At a Glance

1	Fundamentals and Cell Physiology	2
	Fulldamentals and Cell Physiology	
		42
2	Nerve and Muscle, Physical Work	42
3	Autonomic Nervous System (ANS)	78
4	Blood	88
5	Respiration	106
,	кезрицион	
_	AND U. C.	138
6	Acid-Base Homeostasis	
		148
7	Kidneys, Salt, and Water Balance	140
8	Cardiovascular System	186
9	Thermal Balance and Thermoregulation	222
10	Nutrition and Digestion	226
10	Nutrition and Digestion	
		266
11	Hormones and Reproduction	200
		310
12	Central Nervous System and Senses	310
13	Appendix	372
	Further Reading	391
	. at the recurring	
	Index	394
	index	

Color Atlas of Physiology

5th edition, completely revised and expanded

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186 color plates by Ruediger Gay and Astried Rothenburger

Thieme Stuttgart · New York



is available from the publisher

1st German edition 1979 2nd German edition 1983 3rd German edition 1988 4th German edition 1991 5th German edition 2001

1st English edition 1981 2nd English edition 1984 3rd English edition 1986 4th English edition 1991

1st Dutch edition 1981 2nd Dutch edition 2001

1st Italian edition 1981 2nd Italian edition 2001

1st Japanese edition 1982 2nd Japanese edition 1992

1st Spanish edition 1982 2nd Spanish edition 1985 3rd Spanish edition 1994 4th Spanish edition 2001 1st Czech edition 1984 2nd Czech edition 1994

1st French edition 1985 2nd French edition 1992 3rd French edition 2001

1st Turkish edition 1986 2nd Turkish edition 1997

1st Greek edition 1989

1st Chinese edition 1991

1st Polish edition 1994

1st Hungarian edition 1994 2nd Hungarian edition 1996

1st Indonesion edition 2000

This book is an authorized translation of the 5th German edition published and copyrighted 2001 by Georg Thieme Verlag, Stuttgart, Germany.

Title of the German edition: Taschenatlas der Physiologie

Translated by Suzyon O'Neal Wandrey, Berlin, Germany

Illustrated by Atelier Gay + Rothenburger, Sternenfels, Germany

© 1981, 2003 Georg Thieme Verlag Rüdigerstraße 14, D-70469 Stuttgart, Germany http://www.thieme.de Thieme New York, 333 Seventh Avenue, New York, N.Y. 10001, U.S.A. http://www.thieme.com

Cover design: Cyclus, Stuttgart Typesetting by: Druckhaus Götz GmbH, Ludwigsburg, Germany Printed in Germany by: Appl Druck GmbH & Co. KG, Wemding, Germany

ISBN 3-13-545005-8 (GTV)

ISBN 1-58890-061-4 (TNY) 1 2 3 4 5

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Preface to the Fifth Edition

The base of knowledge in many sectors of physiology has grown considerably in magnitude and in depth since the last edition of this book was published. Many advances, especially the rapid progress in sequencing the human genome and its gene products, have brought completely new insight into cell function and communication. This made it necessary to edit and, in some cases, enlarge many parts of the book, especially the chapter on the fundamentals of cell physiology and the sections on neurotransmission, mechanisms of intracellular signal transmission, immune defense. and the processing of sensory stimuli. A list of physiological reference values and important formulas were added to the appendix for quick reference. The extensive index now also serves as a key to abbreviations used in the text.

Some of the comments explaining the connections between pathophysiological principles and clinical dysfunctions had to be slightly truncated and set in smaller print. However, this base of knowledge has also grown considerably for the reasons mentioned above. To make allowances for this, a similarly designed book, the *Color Atlas of Pathophysiology* (S. Silbernagl and F. Lang, Thieme), has now been introduced to supplement the well-established *Color Atlas of Physiology*.

I am very grateful for the many helpful comments from attentive readers (including my son Jakob) and for the welcome feedback from my peers, especially Prof. H. Antoni, Freiburg, Prof. C. von Campenhausen, Mainz, Dr. M. Fischer, Mainz, Prof. K.H. Plattig, Erlangen, and Dr. C. Walther, Marburg, and from my colleagues and staff at the Institute in Würzburg. It was again a great pleasure to work with Rüdiger Gay and Astried Rothenburger, to whom I am deeply indebted for revising practically all the illustrations in the book and for designing a number of new color plates. Their extraordinary enthusiasm and professionalism played a decisive role in the materialization of this new edition. To them I extend my sincere thanks, I would also like to thank Suzyon O'Neal Wandrey for her outstanding translation. I greatly appreciate her capable and careful work. I am also indebted to the publishing staff, especially Marianne Mauch, an extremely competent and motivated editor, and Gert Krüger for invaluable production assistance. I would also like to thank Katharina Völker for her ever observant and conscientious assistance in preparing the index.

I hope that the 5th Edition of the Color Atlas of Physiology will prove to be a valuable tool for helping students better understand physiological correlates, and that it will be a valuable reference for practicing physicians and scientists, to help them recall previously learned information and gain new insights in physiology.

Würzburg, December 2002 Stefan Silbernagl*

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Preface to the First Edition

In the modern world, visual pathways have outdistanced other avenues for informational input. This book takes advantage of the economy of visual representation to indicate the simultaneity and multiplicity of physiological phenomena. Although some subjects lend themselves more readily than others to this treatment, inclusive rather than selective coverage of the key elements of physiology has been attempted.

Clearly, this book of little more than 300 pages, only half of which are textual, cannot be considered as a primary source for the serious student of physiology. Nevertheless, it does contain most of the basic principles and facts taught in a medical school introductory course. Each unit of text and illustration can serve initially as an overview for introduction to the subject and subsequently as a concise review of the material. The contents are as current as the publishing art permits and include both classical information for the beginning students as well as recent details and trends for the advanced student.

A book of this nature is inevitably derivative, but many of the representations are new and, we hope, innovative. A number of people have contributed directly and indirectly to the completion of this volume, but none more than Sarah Iones, who gave much more than editorial assistance. Acknowledgement of helpful criticism and advice is due also to Drs. R. Greger, A. Ratner, J. Weiss, and S. Wood, and Prof. H. Seller. We are grateful to Joy Wieser for her help in checking the proofs. Wolf-Rüdiger and Barbara Gay are especially recognized, not only for their art work, but for their conceptual contributions as well. The publishers, Georg Thieme Verlag and Deutscher Taschenbuch Verlag, contributed valuable assistance based on extensive experience; an author could wish for no better relationship, Finally, special recognition to Dr. Walter Kumpmann for inspiring the project and for his unquestioning confidence in the authors

Basel and Innsbruck, Summer 1979
Agamemnon Despopoulos
Stefan Silbernagl

From the Preface to the Third Edition

The first German edition of this book was already in press when, on November 2nd, 1979, Agamennon Despopoulos and his wife, Sarah Jones-Despopoulos put to sea from Bizerta, Tunisia. Their intention was to cross the Atlantic in their sailing boat. This was the last that was ever heard of them and we have had to abandon all hope of seeing them again.

Without the creative enthusiasm of Agamennon Despopoulos, it is doubtful whether this book would have been possible; without his personal support it has not been easy to continue with the project. Whilst keeping in mind our original aims, I have completely revised the book, incorporating the latest advances in the field of physiology as well as the welcome suggestions provided by readers of the earlier edition, to whom I extend my thanks for their active interest

Würzburg, Fall 1985 Stefan Silbernagl



Dr. Agamemnon Despopoulos

Born 1924 in New York; Professor of Physiology at the University of New Mexico. Albuquerque, USA, until 1971; thereafter scientific adviser to CIBA-GEIGY, Basel.

Table of Contents

1	Fundamentals and Cell Physiology	2
	The Body: an Open System with an Internal Environment ··· 2 Control and Regulation ··· 4 The Cell ··· 8 Transport In, Through, and Between Cells ··· 16 Passive Transport by Means of Diffusion ··· 20 Osmosis, Filtration, and Convection ··· 24 Active Transport ··· 26 Cell Migration ··· 30 Electrical Membrane Potentials and Ion Channels ··· 32 Role of Ca ²⁺ in Cell Regulation ··· 36 Energy Production and Metabolism ··· 38	
2	Nerve and Muscle, Physical Work	42
	Neuron Structure and Function ··· 42 Resting Membrane Potential ··· 44 Action Potential ··· 46 Propagation of Action Potentials in Nerve Fiber ··· 48 Artificial Stimulation of Nerve Cells ··· 50 Synaptic Transmission ··· 50 Motor End-plate ··· 56 Motility and Muscle Types ··· 58 Motor Unit of Skeletal Muscle ··· 58 Contractile Apparatus of Striated Muscle ··· 60 Contraction of Striated Muscle ··· 62 Mechanical Features of Skeletal Muscle ··· 65 Smooth Muscle ··· 70 Energy Supply for Muscle Contraction ··· 72 Physical Work ··· 74 Physical Fitness and Training ··· 76	
3	Autonomic Nervous System (ANS)	78
	Organization of the Autonomic Nervous System ··· 78 Acetylcholine and Cholinergic Transmission ··· 82 Catecholamine, Adrenergic Transmission and Adrenoceptors ··· 84 Adrenal Medulla ··· 86 Non-cholinergic, Non-adrenergic Transmitters ··· 86	

Blood 88 4 Composition and Function of Blood ... 88 Iron Metabolism and Erythropoiesis ... 90 Flow Properties of Blood · · · 92 Plasma, Ion Distribution · · · 92 Immune System · · · 94 Hypersensitivity Reactions (Allergies) ··· 100 Blood Groups · · · 100 Hemostasis · · · 102 Fibrinolysis and Thromboprotection ... 104 Respiration 106 5 Lung Function, Respiration ... 106 Mechanics of Breathing ... 108 Purification of Respiratory Air ... 110 Artificial Respiration · · · 110 Pneumothorax ··· 110 Lung Volumes and their Measurement · · · 112 Dead Space, Residual Volume, and Airway Resistance · · · 114 Lung-Chest Pressure-Volume Curve, Respiratory Work · · · 116 Surface Tension, Surfactant ... 118 Dynamic Lung Function Tests ... 118 Pulmonary Gas Exchange ... 120 Pulmonary Blood Flow, Ventilation-Perfusion Ratio ... 122 CO2 Transport in Blood ... 124 CO₂ Binding in Blood · · · 126 CO₂ in Cerebrospinal Fluid ··· 126 Binding and Transport of O2 in Blood · · · 128 Internal (Tissue) Respiration, Hypoxia · · · 130 Respiratory Control and Stimulation ... 132 Effects of Diving on Respiration · · · 134 Effects of High Altitude on Respiration ... 136 Oxygen Toxicity · · · 136 Acid-Base Homeostasis 138 pH, pH Buffers, Acid-Base Balance · · · 138 Bicarbonate/Carbon Dioxide Buffer · · · 140 Acidosis and Alkalosis · · · 142 Assessment of Acid-Base Status · · · 146 Kidneys, Salt, and Water Balance 148 Kidney Structure and Function · · · 148 Renal Circulation · · · 150 Glomerular Filtration and Clearance · · · 152 Transport Processes at the Nephron · · · 154

Reabsorption of Organic Substances · · · 158

X

Excretion of Organic Substances · · · 160 Reabsorption of Na⁺ and Cl⁻ ··· 162 Reabsorption of Water, Formation of Concentrated Urine · · · 164 Body Fluid Homeostasis · · · 168 Salt and Water Regulation ... 170 Diuresis and Diuretics ... 172 Disturbances of Salt and Water Homeostasis · · · 172 The Kidney and Acid-Base Balance ... 174 Reabsorption and Excretion of Phosphate, Ca²⁺ and Mg²⁺ ··· 178 Potassium Balance · · · 180 Tubuloglomerular Feedback, Renin-Angiotensin System · · · 184 186 Cardiovascular System Overview ··· 186 Blood Vessels and Blood Flow · · · 188 Cardiac Cycle · · · 190 Cardiac Impulse Generation and Conduction · · · 192 Electrocardiogram (ECG) ··· 196 Excitation in Electrolyte Disturbances ... 198 Cardiac Arrhythmias ... 200 Ventricular Pressure-Volume Relationships · · · 202 Cardiac Work and Cardiac Power ... 202 Regulation of Stroke Volume ... 204 Venous Return ··· 204 Arterial Blood Pressure ··· 206 Endothelial Exchange Processes ... 208 Myocardial Oxygen Supply ... 210 Regulation of the Circulation · · · 212 Circulatory Shock · · · 218 Fetal and Neonatal Circulation ... 220 Thermal Balance and Thermoregulation 222 Thermal Balance · · · 222 Thermoregulation ··· 224 226 **Nutrition and Digestion** 10 Nutrition ··· 226 Energy Metabolism and Calorimetry ... 228 Energy Homeostasis and Body Weight ... 230 Gastrointestinal (GI) Tract: Overview, Immune Defense and Blood Flow ... 232 Neural and Hormonal Integration ... 234 Saliva ... 236 Deglutition ··· 238 Vomiting ··· 238 Stomach Structure and Motility ... 240 Gastric Juice · · · 242 Small Intestinal Function ... 244

Bile ... 248 Excretory Liver Function—Bilirubin ... 250 Lipid Digestion · · · 252 Lipid Distribution and Storage ... 254 Digestion and Absorption of Carbohydrates and Protein ... 258 Vitamin Absorption · · · 260 Water and Mineral Absorption ... 262 Large Intestine, Defecation, Feces · · · 264 266 **Hormones and Reproduction** 11 Integrative Systems of the Body ... 266 Hormones ··· 268 Humoral Signals: Control and Effects ... 272 Cellular Transmission of Signals from Extracellular Messengers ... 274 Hypothalamic-Pituitary System ... 280 Carbohydrate Metabolism and Pancreatic Hormones · · · 282 Thyroid Hormones ... 286 Calcium and Phosphate Metabolism ... 290 Biosynthesis of Steroid Hormones ... 294 Adrenal Cortex and Glucocorticoid Synthesis · · · 296 Oogenesis and the Menstrual Cycle ... 298 Hormonal Control of the Menstrual Cycle ... 300 Estrogens ··· 302 Progesterone · · · 302 Prolactin and Oxytocin · · · 303 Hormonal Control of Pregnancy and Birth ... 304 Androgens and Testicular Function · · · 306 Sexual Response, Intercourse and Fertilization ... 308 **Central Nervous System and Senses** 310 12 Central Nervous System · · · 310 Cerebrospinal Fluid ... 310 Stimulus Reception and Processing · · · 312 Sensory Functions of the Skin · · · 314 Proprioception, Stretch Reflex ... 316 Nociception and Pain · · · 318 Polysynaptic Reflexes ··· 320 Synaptic Inhibition · · · 320 Central Conduction of Sensory Input ... 322 Motor System · · · 324 Hypothalamus, Limbic System ... 330 Cerebral Cortex, Electroencephalogram (EEG) ··· 332 Sleep-Wake Cycle, Circadian Rhythms ... 334 Consciousness, Memory, Language ... 336 Glia ... 338 Sense of Taste · · · 338 Sense of Smell ··· 340

Pancreas ... 246

XII

Optical Apparatus of the Eye ... 346 Visual Acuity. Photosensors ··· 348 Adaptation of the Eye to Different Light Intensities ... 352 Retinal Processing of Visual Stimuli ... 354 Color Vision ... 356 Visual Field, Visual Pathway, Central Processing of Visual Stimuli ... 358 Eve Movements, Stereoscopic Vision, Depth Perception ... 360 Physical Principles of Sound—Sound Stimulus and Perception · · · 362 Conduction of Sound, Sound Sensors ... 364 Central Processing of Acoustic Information · · · 368 Voice and Speech ··· 370 **Appendix** 372 Dimensions and Units ... 372 Powers and Logarithms ... 380 Graphic Representation of Data ... 381 The Greek Alphabet ... 384 Reference Values in Physiology ... 384 Important Equations in Physiology ... 388 391 **Further Reading**

Sense of Balance ... 342

13

Index

Eye Structure, Tear Fluid, Aqueous Humor ... 344

394

Fundamentals and Cell Physiology

"... If we break up a living organism by isolating its different parts, it is only for the sake of ease in analysis and by no means in order to conceive them separately. Indeed, when we wish to ascribe to a physiological quality its value and true significance, we must always refer it to the whole and draw our final conclusions only in relation to its effects on the whole."

Claude Bernard (1865)

The Body: an Open System with an Internal Environment

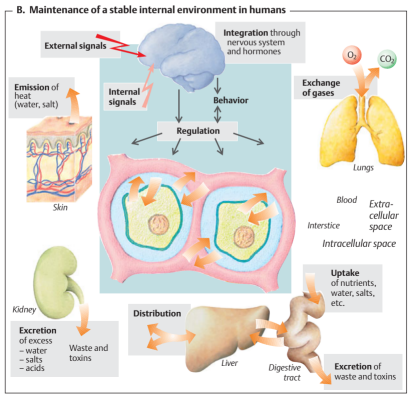
The existence of unicellular organisms is the epitome of life in its simplest form. Even simple protists must meet two basic but essentially conflicting demands in order to survive. A unicellular organism must, on the one hand, isolate itself from the seeming disorder of its inanimate surroundings, yet, as an "open system" (\rightarrow p. 40), it is dependent on its environment for the exchange of heat, oxygen, nutrients, waste materials, and information.

"Isolation" is mainly ensured by the cell membrane, the hydrophobic properties of which prevent the potentially fatal mixing of hydrophilic components in watery solutions inside and outside the cell. Protein molecules within the cell membrane ensure the permeability of the membrane barrier. They may exist in the form of pores (channels) or as more complex transport proteins known as carriers $(\rightarrow p.26 \, \text{ff.})$. Both types are selective for certain substances, and their activity is usually regulated. The cell membrane is relatively well permeable to hydrophobic molecules such as gases. This is useful for the exchange of O2 and CO₂ and for the uptake of lipophilic signal substances, yet exposes the cell to poisonous gases such as carbon monoxide (CO) and lipophilic noxae such as organic solvents. The cell membrane also contains other proteins-namely, receptors and enzymes. Receptors receive signals from the external environment and convev the information to the interior of the cell (signal transduction), and enzymes enable the cell to metabolize extracellular substrates.

Let us imagine the primordial sea as the external environment of the unicellular organism (\rightarrow A). This milieu remains more or less constant, although the organism absorbs nutrients from it and excretes waste into it. In spite of its simple structure, the unicellular or-

ganism is capable of eliciting motor responses to signals from the environment. This is achieved by moving its pseudopodia or flagella, for example, in response to changes in the food concentration.

The evolution from unicellular organisms to multicellular organisms, the transition from specialized cell groups to organs, the emergence of the two sexes, the coexistence of individuals in social groups, and the transition from water to land have tremendously increased the efficiency, survival, radius of action, and independence of living organisms. This process required the simultaneous development of a complex infrastructure within the organism. Nonetheless, the individual cells of the body still need a milieu like that of the primordial sea for life and survival. Today, the extracellular fluid is responsible for providing constant environmental conditions ($\rightarrow B$), but the volume of the fluid is no longer infinite. In fact, it is even smaller than the intracellular volume (\rightarrow p. 168). Because of their metabolic activity, the cells would quickly deplete the oxygen and nutrient stores within the fluids and flood their surroundings with waste products if organs capable of maintaining a stable internal environment had not developed. This is achieved through homeostasis, a process by which physiologic self-regulatory mechanisms (see below) maintain steady states in the body through coordinated physiological activity. Specialized organs ensure the continuous absorption of nutrients, electrolytes and water and the excretion of waste products via the urine and feces. The circulating blood connects the organs to every inch of the body, and the exchange of materials between the blood and the intercellular spaces (interstices) creates a stable environment for the cells. Organs such as the digestive tract and liver absorb nutrients and make them available by processing, metabolizing and distributing



them throughout the body. The lung is responsible for the exchange of gases (O2 intake, CO₂ elimination), the liver and kidney for the excretion of waste and foreign substances, and the skin for the release of heat. The kidney and lungs also play an important role in regulating the internal environment, e.g., water content, osmolality, ion concentrations, pH (kidney, lungs) and O_2 and CO_2 pressure (lungs) ($\rightarrow \mathbf{B}$).

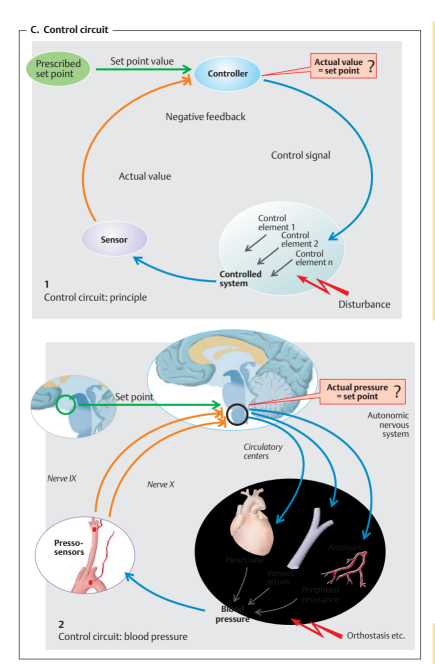
The specialization of cells and organs for specific tasks naturally requires integration, which is achieved by convective transport over long distances (circulation, respiratory tract), humoral transfer of information (hormones), and transmission of electrical signals in the nervous system, to name a few examples. These mechanisms are responsible for supply and disposal and thereby maintain a stable internal environment, even under conditions of extremely high demand and stress. Moreover, they control and regulate functions that ensure survival in the sense of preservation of the species. Important factors in this process include not only the timely development of reproductive organs and the availability of fertilizable gametes at sexual maturity, but also the control of erection, ejaculation, fertilization, and nidation. Others include the coordination of functions in the mother and fetus during pregnancy and regulation of the birth process and the lactation period.

The central nervous system (CNS) processes signals from peripheral sensors (single sensory cells or sensory organs), activates outwardly directed effectors (e.g., muscles), and influences the endocrine glands. The CNS is the focus of attention when studying human or animal behavior. It helps us to locate food and water and protects us from heat or cold. The central nervous system also plays a role in partner selection, concern for offspring even long after their birth, and integration into social systems. The CNS is also involved in the development, expression, and processing of emotions such as desire, listlessness, curiosity, wishfulness, happiness, anger, wrath, and envy and of traits such as creativeness, inquisitiveness, self-awareness, and responsibility. This goes far beyond the scope of physiology which in the narrower sense is the study of the functions of the body-and, hence, of this book.

Although behavioral science, sociology, and psychology are disciplines that border on physiology, true bridges between them and physiology have been established only in exceptional cases.

Control and Regulation

In order to have useful cooperation between the specialized organs of the body, their functions must be adjusted to meet specific needs. In other words, the organs must be subject to control and regulation. Control implies that a controlled variable such as the blood pressure is subject to selective external modification. for example, through alteration of the heart rate (\rightarrow p, 218). Because many other factors also affect the blood pressure and heart rate. the controlled variable can only be kept constant by continuously measuring the current blood pressure, comparing it with the reference signal (set point), and continuously correcting any deviations. If the blood pressure drops-due, for example, to rapidly standing up from a recumbent position—the heart rate will increase until the blood pressure has been reasonably adjusted. Once the blood pressure has risen above a certain limit, the heart rate will decrease again and the blood pressure will normalize. This type of closed-loop control is called a negative feedback control system or a **control circuit** (\rightarrow C1). It consists of a controller with a programmed set-point value (target value) and control elements (effectors) that can adjust the controlled variable to the set point. The system also includes sensors that continuously measure the actual value of the controlled variable of interest and report it (feedback) to the controller, which compares the actual value of the controlled variable with the set-point value and makes the necessary adjustments if disturbance-related discrepancies have occurred. The control system operates either from within the organ itself (autoregulation) or via a superordinate organ such as the central nervous system or hormone glands. Unlike simple control, the elements of a control circuit can work rather imprecisely without causing a deviation from the set point (at least on average). Moreover, control circuits are capable of responding to unexpected dis-



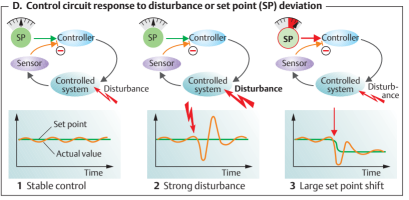
turbances. In the case of blood pressure regulation (\rightarrow C2), for example, the system can respond to events such as orthostasis (\rightarrow p. 204) or sudden blood loss.

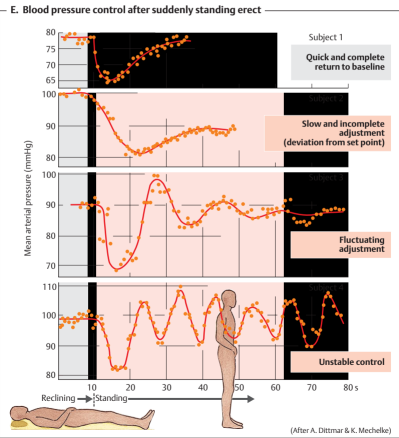
The type of control circuits described above keep the controlled variables constant when disturbance variables cause the controlled variable to deviate from the set point $(\rightarrow D2)$. Within the body, the set point is rarely invariable, but can be "shifted" when requirements of higher priority make such a change necessary. In this case, it is the variation of the set point that creates the discrepancy between the nominal and actual values, thus leading to the activation of regulatory elements $(\rightarrow D3)$. Since the regulatory process is then triggered by variation of the set point (and not by disturbance variables), this is called servocontrol or **servomechanism**. Fever (\rightarrow p. 224) and the adjustment of muscle length by muscle spindles and γ -motor neurons (\rightarrow p. 316) are examples of servocontrol.

In addition to relatively simple variables such as blood pressure, cellular pH, muscle length, body weight and the plasma glucose concentration, the body also regulates complex sequences of events such as fertilization, pregnancy, growth and organ differentiation, as well as sensory stimulus processing and the motor activity of skeletal muscles, e.g., to maintain equilibrium while running. The regulatory process may take parts of a second (e.g., purposeful movement) to several years (e.g., the growth process).

In the control circuits described above, the controlled variables are kept constant on average, with variably large, wave-like deviations. The sudden emergence of a disturbance variable causes larger deviations that quickly normalize in a stable control circuit (\rightarrow E, test subject no. 1). The degree of deviation may be slight in some cases but substantial in others. The latter is true, for example, for the blood glucose concentration, which nearly doubles after meals. This type of regulation obviously functions only to prevent extreme rises and falls (e.g., hyper- or hypoglycemia) or chronic deviation of the controlled variable. More precise maintenance of the controlled variable requires a higher level of regulatory sensitivity (high amplification factor). However, this extends the settling time (\rightarrow **E**, subject no. 3) and can lead to regulatory instability, i.e., a situation where the actual value oscillates back and forth between extremes (*unstable oscillation*, \rightarrow **E**, subject no. 4).

Oscillation of a controlled variable in response to a disturbance variable can be attenuated by either of two mechanisms. First, sensors with differential characteristics (D sensors) ensure that the intensity of the sensor signal increases in proportion with the rate of deviation of the controlled variable from the set point (\rightarrow p. 312 ff.). Second, **feedforward** control ensures that information regarding the expected intensity of disturbance is reported to the controller before the value of the controlled variable has changed at all. Feedforward control can be explained by example of physiologic thermoregulation, a process in which cold receptors on the skin trigger counterregulation before a change in the controlled value (core temperature of the body) has actually occurred (\rightarrow p. 224). The disadvantage of having only D sensors in the control circuit can be demonstrated by example of arterial pressosensors (= pressoreceptors) in acute blood pressure regulation. Very slow but steady changes, as observed in the development of arterial hypertension, then escape regulation. In fact, a rapid drop in the blood pressure of a hypertensive patient will even cause a counterregulatory increase in blood pressure. Therefore, other control systems are needed to ensure proper long-term blood pressure regulation.





The Cell

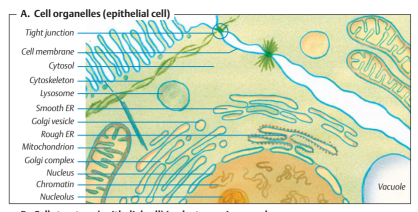
The cell is the smallest functional unit of a living organism. In other words, a cell (and no smaller unit) is able to perform essential vital functions such as metabolism, growth, movement, reproduction, and hereditary transmission (W. Roux) (\rightarrow p. 4). Growth, reproduction, and hereditary transmission can be achieved by *cell division*.

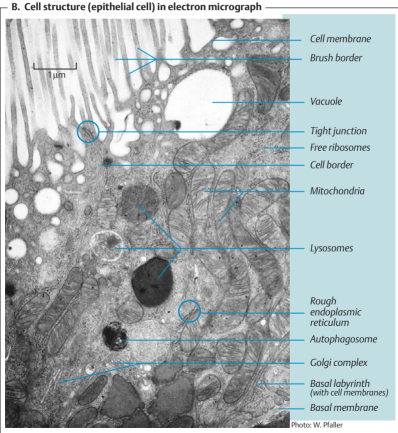
Cell components: All cells consist of a cell membrane, cytosol or cytoplasm (ca. 50 vol.%), and membrane-bound subcellular structures known as $organelles (\rightarrow A, B)$. The organelles of eukaryotic cells are highly specialized. For instance, the genetic material of the cell is concentrated in the cell nucleus, whereas "digestive" enzymes are located in the lysosomes. Oxidative ATP production takes place in the mitochondria.

The cell nucleus contains a liquid known as karyolymph, a nucleolus, and chromatin. Chromatin contains deoxyribonucleic acids (DNA), the carriers of genetic information. Two strands of DNA forming a double helix (up to 7 cm in length) are twisted and folded to form chromosomes 10 µm in length. Humans normally have 46 chromosomes, consisting of 22 autosomal pairs and the chromosomes that determine the sex (XX in females, XY in males). DNA is made up of a strand of three-part molecules called nucleotides, each of which consists of a pentose (deoxyribose) molecule, a phosphate group, and a base. Each sugar molecule of the monotonic sugar-phosphate backbone of the strands (...deoxyribose phosphate-deoxyribose...) is attached to one of four different bases. The sequence of bases represents the genetic code for each of the roughly 100000 different proteins that a cell produces during its lifetime (gene expression). In a DNA double helix, each base in one strand of DNA is bonded to its complementary base in the other strand according to the rule; adenine (A) with thymine (T) and guanine (G) with cytosine (C). The base sequence of one strand of the double helix $(\rightarrow E)$ is always a "mirror image" of the opposite strand. Therefore, one strand can be used as a template for making a new complementary strand, the information content of which is identical to that of the original. In cell division, this process is the means by which duplication of genetic information (replication) is achieved.

Messenger RNA (mRNA) is responsible for code transmission, that is, passage of coding sequences from DNA in the nucleus (base sequence) for protein synthesis in the cytosol (amino acid sequence) (\rightarrow C1), mRNA is formed in the nucleus and differs from DNA in that it consists of only a single strand and that it contains ribose instead of deoxyribose, and uracil (U) instead of thymine. In DNA, each amino acid (e.g., glutamate, \rightarrow E) needed for synthesis of a given protein is coded by a set of three adjacent bases called a codon or triplet (C-T-C in the case of glutamate). In order to transcribe the DNA triplet, mRNA must form a complementary codon (e.g., G-A-G for glutamate). The relatively small transfer RNA (tRNA) molecule is responsible for reading the codon in the ribosomes (\rightarrow C2), tRNA contains a complementary codon called the anticodon for this purpose. The anticodon for glutamate is C-U-C (\rightarrow **E**).

RNA synthesis in the nucleus is controlled by RNA polymerases (types I-III). Their effect on DNA is normally blocked by a repressor protein. Phosphorylation of the polymerase occurs if the repressor is eliminated (de-repression) and the general transcription factors attach to the so-called promoter sequence of the DNA molecule (T-A-T-A in the case of polymerase II). Once activated, it separates the two strands of DNA at a particular site so that the code on one of the strands can be read and transcribed to form mRNA (transcription, → C1a, D). The heterogeneous nuclear RNA (hnRNA) molecules synthesized by the polymerase have a characteristic "cap" at their 5' end and a polyadenine "tail" (A-A-A-...) at the 3' end (\rightarrow **D**). Once synthesized, they are immediately "enveloped" in a protein coat, yielding heterogeneous nuclear ribonucleoprotein (hnRNP) particles. The primary RNA or premRNA of hnRNA contains both coding sequences (exons) and non-coding sequences (introns). The exons code for amino acid sequences of the proteins to be synthesized, whereas the introns are not involved in the coding process. Introns may contain 100 to 10 000 nucleotides; they are removed from the





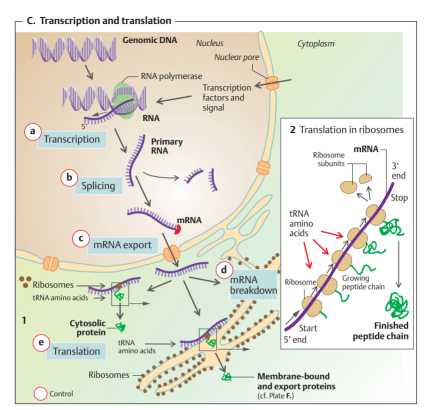
primary mRNA strand by splicing (\rightarrow C1b, D) and then degraded. The introns, themselves, contain the information on the exact splicing site. Splicing is ATP-dependent and requires the interaction of a number of proteins within ribonucleoprotein complex called the spliceosome. Introns usually make up the lion's share of pre-mRNA molecules. For example, they make up 95% of the nucleotide chain of coagulation factor VIII, which contains 25 introns, mRNA can also be modified (e.g., through methylation) during the course of posttranscriptional modification.

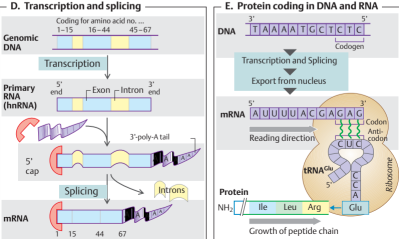
RNA now exits the nucleus through nuclear pores (around 4000 per nucleus) and enters the cytosol (\rightarrow C1c). Nuclear pores are high-molecular-weight protein complexes (125 MDa) located within the nuclear envelope. They allow large molecules such as transcription factors, RNA polymerases or cytoplasmic steroid hormone receptors to pass into the nucleus, nuclear molecules such as mRNA and tRNA to pass out of the nucleus, and other molecules such as ribosomal proteins to travel both ways. The (ATP-dependent) passage of a molecule in either direction cannot occur without the help of a specific signal that guides the molecule into the pore. The abovementioned 5' cap is responsible for the exit of mRNA from the nucleus, and one or two specific sequences of a few (mostly cationic) amino acids are required as the signal for the entry of proteins into the nucleus. These sequences form part of the peptide chain of such nuclear proteins and probably create a peptide loop on the protein's surface. In the case of the cytoplasmic receptor for glucocorticoids (\rightarrow p. 278), the nuclear localization signal is masked by a chaperone protein (heat shock protein 90, hsp90) in the absence of the glucocorticoid, and is released only after the hormone binds, thereby freeing hsp90 from the receptor. The "activated" receptor then reaches the cell nucleus, where it binds to specific DNA sequences and controls specific genes.

The nuclear envelope consists of two membranes (=two phospholipid bilayers) that merge at the nuclear pores. The two membranes consist of different materials. The external membrane is continuous with the membrane of the endoplasmic reticulum (ER), which is described below $(\rightarrow \mathbf{F})$.

The mRNA exported from the nucleus travels to the **ribosomes** (\rightarrow C1), which either float freely in the cytosol or are bound to the cytosolic side of the endoplasmic reticulum, as described below. Each ribosome is made up of dozens of proteins associated with a number of structural RNA molecules called ribosomal RNA (rRNA). The two subunits of the ribosome are first transcribed from numerous rRNA genes in the **nucleolus**, then separately exit the cell nucleus through the nuclear pores. Assembled together to form a ribosome, they now comprise the biochemical "machinery" for protein synthesis (translation) (\rightarrow C2). Synthesis of a peptide chain also requires the presence of specific tRNA molecules (at least one for each of the 21 proteinogenous amino acids). In this case, the target amino acid is bound to the C-C-A end of the tRNA molecule (same in all tRNAs), and the corresponding anticodon that recognizes the mRNA codon is located at the other end ($\rightarrow E$). Each ribosome has two tRNA binding sites: one for the last incorporated amino acid and another for the one beside it (not shown in E). Protein synthesis begins when the start codon is read and ends once the stop codon has been reached. The ribosome then breaks down into its two subunits and releases the mRNA (\rightarrow C2). Ribosomes can add approximately 10-20 amino acids per second. However, since an mRNA strand is usually translated simultaneously by many ribosomes (polyribosomes or polysomes) at different sites, a protein is synthesized much faster than its mRNA. In the bone marrow, for example, a total of around 5×10^{14} hemoglobin copies containing 574 amino acids each are produced per second.

The endoplasmic reticulum (ER. \rightarrow C. F) plays a central role in the synthesis of proteins and lipids; it also serves as an intracellular Ca²⁺ store (\rightarrow p. 17 A). The ER consists of a net-like system of interconnected branched channels and flat cavities bounded by a membrane. The enclosed spaces (cisterns) make up around 10% of the cell volume, and the membrane comprises up to 70% of the membrane mass of a cell. Ribosomes can attach to the cytosolic surface of parts of the ER, forming a rough endo-





plasmic reticulum (RER). These ribosomes synthesize export proteins as well as transmembrane proteins $(\rightarrow \mathbf{G})$ for the plasma membrane, endoplasmic reticulum, Golgi apparatus, lysosomes, etc. The start of protein synthesis (at the amino end) by such ribosomes (still unattached) induces a signal sequence to which a signal recognition particle (SRP) in the cytosol attaches. As a result, (a) synthesis is temporarily halted and (b) the ribosome (mediated by the SRP and a SRP receptor) attaches to a ribosome receptor on the ER membrane. After that, synthesis continues. In export protein synthesis, a translocator protein conveys the peptide chain to the cisternal space once synthesis is completed. Synthesis of membrane proteins is interrupted several times (depending on the number of membrane-spanning domains $(\rightarrow G2)$ by translocator protein closure, and the corresponding (hydrophobic) peptide sequence is pushed into the phospholipid membrane. The smooth endoplasmic reticulum (SER) contains no ribosomes and is the production site of lipids (e.g., for lipoproteins, \rightarrow p. 254 ff.) and other substances. The ER membrane containing the synthesized membrane proteins or export proteins forms vesicles which are transported to the Golgi apparatus.

The Golgi complex or **Golgi apparatus** $(\rightarrow F)$ has sequentially linked functional compartments for further processing of products from the endoplasmic reticulum. It consists of a cis-Golgi network (entry side facing the ER), stacked flattened cisternae (Golgi stacks) and a trans-Golgi network (sorting and distribution). Functions of the Golgi complex:

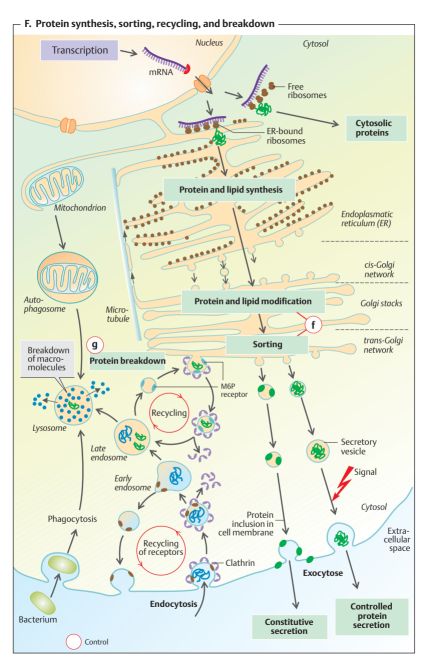
- polysaccharide synthesis;
- protein processing (posttranslational modification), e.g., glycosylation of membrane proteins on certain amino acids (in part in the ER) that are later borne as glycocalyces on the external cell surface (see below) and γ-carboxylation of glutamate residues (\rightarrow p. 102);
- phosphorylation of sugars of glycoproteins (e.g., to mannose-6-phosphate, as described below);
- "packaging" of proteins meant for export into secretory vesicles (secretory granules), the contents of which are exocytosed into the extracellular space; see p. 246, for example.

Hence, the Golgi apparatus represents a central modification, sorting and distribution center for proteins and lipids received from the endoplasmic reticulum.

Regulation of gene expression takes place on the level of transcription (\rightarrow C1a), RNA modification (\rightarrow C1b), mRNA export (\rightarrow C1c), RNA degradation (\rightarrow C1d), translation (\rightarrow C1e), modification and sorting $(\rightarrow F.f)$, and protein degradation (\rightarrow **F.a**).

The **mitochondria** (\rightarrow **A, B**; p. 17 B) are the site of oxidation of carbohydrates and lipids to CO2 and H2O and associated O2 expenditure. The Krebs cycle (citric acid cycle), respiratory chain and related ATP synthesis also occur in mitochondria. Cells intensely active in metabolic and transport activities are rich in mitochondria-e.g., hepatocytes, intestinal cells, and renal epithelial cells. Mitochondria are enclosed in a double membrane consisting of a smooth outer membrane and an inner membrane. The latter is deeply infolded, forming a series of projections (cristae); it also has important transport functions (\rightarrow p. 17 B). Mitochondria probably evolved as a result of symbiosis between aerobic bacteria and anaerobic cells (symbiosis hypothesis). The mitochondrial DNA (mtDNA) of bacterial origin and the double membrane of mitochondria are relicts of their ancient history. Mitochondria also contain ribosomes which synthesize all proteins encoded by mtDNA.

Lysosomes are vesicles $(\rightarrow F)$ that arise from the ER (via the Golgi apparatus) and are involved in the intracellular digestion of macromolecules. These are taken up into the cell either by endocytosis (e.g., uptake of albumin into the renal tubules; \rightarrow p. 158) or by *phagocy*tosis (e.g., uptake of bacteria by macrophages; \rightarrow p. 94 ff.). They may also originate from the degradation of a cell's own organelles (autophagia, e.g., of mitochondria) delivered inside autophagosomes (\rightarrow **B**, **F**). A portion of the endocytosed membrane material recycles (e.g., receptor recycling in receptor-mediated endocytosis; \rightarrow p. 28). Early and late endosomes are intermediate stages in this vesicular transport. Late endosomes and lysosomes contain acidic hydrolases (proteases, nucleases, lipases, glycosidases, phosphatases, etc., that are active only under acidic conditions). The



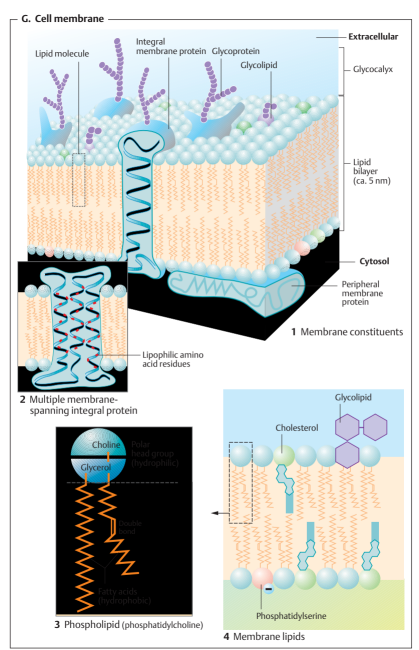
membrane contains an H+-ATPase that creates an acidic (pH5) interior environment within the lysosomes and assorted transport proteins that (a) release the products of digestion (e.g., amino acids) into the cytoplasm and (b) ensure charge compensation during H+ uptake (Clchannels). These enzymes and transport proteins are delivered in primary lysosomes from the Golgi apparatus. Mannose-6-phosphate (M6P) serves as the "label" for this process: it binds to M6 P receptors in the Golgi membrane which, as in the case of receptor-mediated endocytosis (\rightarrow p. 28), cluster in the membrane with the help of a clathrin framework. In the acidic environment of the lysosomes, the enzymes and transport proteins are separated from the receptor, and M6P is dephosphorylated. The M6P receptor returns to the Golgi apparatus (recycling, \rightarrow **F**). The M6 P receptor no longer recognizes the dephosphorylated proteins, which prevents them from returning to the Golgi apparatus.

Peroxisomes are microbodies containing enzymes (imported via a signal sequence) that permit the oxidation of certain organic molecules (R-H₂), such as amino acids and fatty acids: R-H₂ + O₂ \rightarrow R + H₂O₂. The peroxisomes also contain catalase, which transforms 2 H₂O₂ into O₂ + H₂O and oxidizes toxins, such as alcohol and other substances.

Whereas the membrane of organelles is responsible for intracellular compartmentalization, the main job of the **cell membrane** $(\rightarrow G)$ is to separate the cell interior from the extracellular space (\rightarrow p. 2). The cell membrane is a phospholipid bilayer (\rightarrow **G1**) that may be either smooth or deeply infolded, like the brush border or the basal labyrinth (\rightarrow **B**). Depending on the cell type, the cell membrane contains variable amounts of phospholipids, cholesterol, and glycolipids (e.g., cerebrosides). The phospholipids mainly consist of phosphatidylcholine $(\rightarrow G3)$, phosphatidylserine, phosphatidylethanolamine, and sphingomyelin. The hydrophobic components of the membrane face each other, whereas the hydrophilic components face the watery surroundings, that is, the extracellular fluid or cytosol (\rightarrow G4). The lipid composition of the two layers of the membrane differs greatly. Glycolipids are present only in the external layer, as described

below. Cholesterol (present in both layers) reduces both the fluidity of the membrane and its permeability to polar substances. Within the two-dimensionally fluid phospholipid membrane are proteins that make up 25% (myelin membrane) to 75% (inner mitochondrial membrane) of the membrane mass, depending on the membrane type. Many of them span the entire lipid bilayer once $(\rightarrow G1)$ or several times $(\rightarrow G2)$ (transmembrane proteins). thereby serving as ion channels, carrier proteins, hormone receptors, etc. The proteins are anchored by their lipophilic amino acid residues, or attached to already anchored proteins. Some proteins can move about freely within the membrane, whereas others, like the anion exchanger of red cells, are anchored to the cytoskeleton. The cell surface is largely covered by the glycocalyx, which consists of sugar moieties of glycoproteins and glycolipids in the cell membrane (\rightarrow **G1.4**) and of the extracellular matrix. The glycocalyx mediates cellcell interactions (surface recognition, cell docking, etc.). For example, components of the glycocalyx of neutrophils dock onto endothelial membrane proteins, called selectins $(\rightarrow p.94)$.

The **cytoskeleton** allows the cell to maintain and change its shape (during cell division, etc.), make selective movements (migration, cilia), and conduct intracellular transport activities (vesicle, mitosis). It contains *actin filaments* as well as *microtubules* and *intermediate filaments* (e.g., vimentin and desmin filaments, neurofilaments, keratin filaments) that extend from the centrosome.



Transport In, Through and Between Cells

The lipophilic cell membrane protects the cell interior from the extracellular fluid, which has a completely different composition $(\rightarrow p, 2)$. This is imperative for the creation and maintenance of a cell's internal environment by means of metabolic energy expenditure. Channels (pores), carriers, ion pumps ($\rightarrow p.26ff$.) and the process of cytosis $(\rightarrow p.28)$ allow transmembrane transport of selected substances. This includes the import and export of metabolic substrates and metabolites and the selective transport of ions used to create or modify the *cell potential* (\rightarrow p. 32), which plays an essential role in excitability of nerve and muscle cells. In addition, the effects of substances that readily penetrate the cell membrane in most cases (e.g., water and CO₂) can be mitigated by selectively transporting certain other substances. This allows the cell to compensate for undesirable changes in the cell volume or pH of the cell interior.

Intracellular Transport

The cell interior is divided into different compartments by the organelle membranes. In some cases, very broad intracellular spaces must be crossed during transport. For this purpose, a variety of specific intracellular transport mechanisms exist, for example:

- Nuclear pores in the nuclear envelope provide the channels for RNA export out of the nucleus and protein import into it (→ p. 11 C);
- ♦ Protein transport from the rough endoplasmic reticulum to the Golgi complex $(\rightarrow p. 13 F)$;
- ◆ Axonal transport in the nerve fibers, in which distances of up to 1 meter can be crossed (→p.42). These transport processes mainly take place along the filaments of the cytoskeleton. Example: while expending ATP, the microtubules set dynein-bound vesicles in motion in the one direction, and kinesinbound vesicles in the other (→p.13 F).

Intracellular Transmembrane Transport

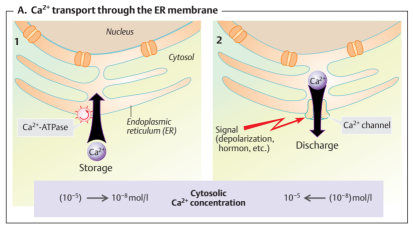
Main sites:

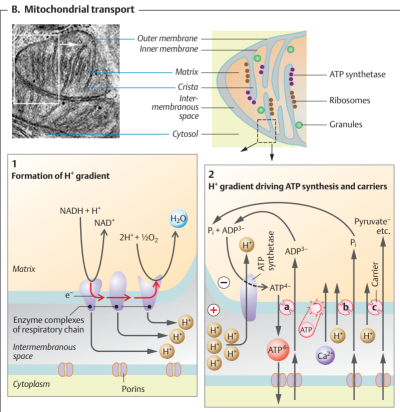
 Lysosomes: Uptake of H⁺ ions from the cytosol and release of metabolites such as amino acids into the cytosol (→ p. 12);

- ◆ Endoplasmic reticulum (ER): In addition to a translocator protein (\rightarrow p. 10), the ER has two other proteins that transport Ca²⁺ (\rightarrow A). Ca²⁺ can be pumped from the cytosol into the ER by a Ca²⁺-ATPase called SERCA (sarcoplasmic endoplasmic reticulum Ca²⁺-transporting ATPase). The resulting Ca²⁺ stores can be released into the cytosol via a Ca²⁺ channel (ryanodine receptor, RyR) in response to a triggering signal (\rightarrow p. 36).
- Mitochondria: The outer membrane contains large pores called porins that render it permeable to small molecules (< 5 kDa), and the inner membrane has high concentrations of specific carriers and enzymes $(\rightarrow B)$. Enzyme complexes of the respiratory chain transfer electrons (e-) from high to low energy levels, thereby pumping H⁺ ions from the matrix space into the intermembrane space $(\rightarrow B1)$, resulting in the formation of an H^+ ion gradient directed into the matrix. This not only drives ATP synthetase (ATP production; \rightarrow **B2**), but also promotes the inflow of pyruvate - and anorganic phosphate, P_i^- (symport; \rightarrow **B2b,c** and p. 28). Ca2+ ions that regulate Ca2+-sensitive mitochondrial enzymes in muscle tissue can be pumped into the matrix space with ATP expenditure (\rightarrow **B2**), thereby allowing the mitochondria to form a sort of Ca²⁺ buffer space for protection against dangerously high concentrations of Ca2+ in the cytosol. The insidenegative membrane potential (caused by H+ release) drives the uptake of ADP3- in exchange for ATP⁴⁻ (potential-driven transport; \rightarrow **B2a** and p. 22).

Transport between Adjacent Cells

In the body, transport between adjacent cells occurs either via diffusion through the extracellular space (e.g., paracrine hormone effects) or through channel-like connecting structures (connexons) located within a so-called gap junction or nexus (\rightarrow C). A connexon is a hemichannel formed by six connexin molecules (\rightarrow C2). One connexon docks with another connexon on an adjacent cell, thereby forming a common channel through which substances with molecular masses of up to around 1 kDa can pass. Since this applies not only for ions such as Ca²⁺, but also for a number of organic substances such as ATP, these types of cells are





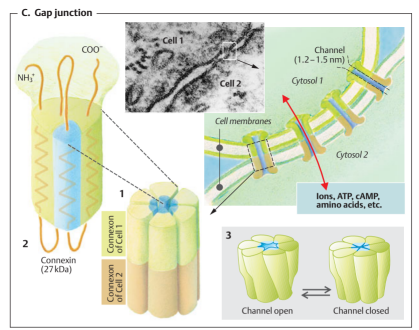
Transport through Cell Layers

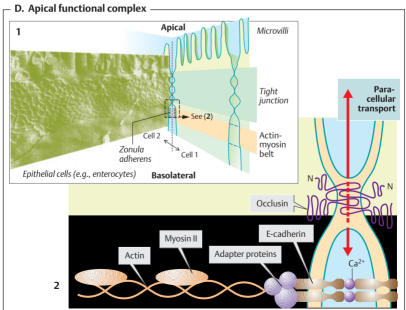
Multicellular organisms have cell layers that are responsible for separating the "interior" from the "exterior" of the organism and its larger compartments. The epithelia of skin and gastrointestinal, urogenital and respiratory tracts, the endothelia of blood vessels, and neuroglia are examples of this type of extensive barrier. They separate the immediate extracellular space from other spaces that are greatly different in composition, e.g., those filled with air (skin, bronchial epithelia), gastrointestinal contents, urine or bile (tubules, urinary bladder, gallbladder), aqueous humor of the eve. blood (endothelia) and cerebrospinal fluid (blood-cerebrospinal fluid barrier), and from the extracellular space of the CNS (blood-brain barrier). Nonetheless. certain substances must be able to pass through these cell layers. This requires selective transcellular transport with import into the cell followed by export from the cell. Unlike cells with a completely uniform plasma membrane (e.g., blood cells), epi- and endothelial cells are polar cells, as defined by

their structure (\rightarrow p. 9A and B) and transport function. Hence, the *apical membrane* (facing exterior) of an epithelial cell has a different set of transport proteins from the *basolateral membrane* (facing the blood). Tight junctions (described below) at which the outer phospholipid layer of the membrane folds over, prevent lateral mixing of the two membranes (\rightarrow D2).

Whereas the apical and basolateral membranes permit transcellular transport, paracellular transport takes place between cells. Certain epithelia (e.g., in the small intestinal and proximal renal tubules) are relatively permeable to small molecules (leaky), whereas others are less leaky (e.g., distal nephron, colon). The degree of permeability depends on the strength of the tight junctions (zonulae occludentes) holding the cells together $(\rightarrow \mathbf{D})$. The paracellular pathway and the extent of its permeability (sometimes cation-specific) are essential functional elements of the various epithelia. Macromolecules can cross the barrier formed by the endothelium of the vessel wall by transcytosis (\rightarrow p. 28), yet paracellular transport also plays an essential role, especially in the fenestrated endothelium. Anionic macromolecules like albumin, which must remain in the bloodstream because of its colloid osmotic action (\rightarrow p. 208), are held back by the wall charges at the intercellular spaces and, in some cases, at the fenestra.

Long-distance transport between the various organs of the body and between the body and the outside world is also necessary. *Convection* is the most important transport mechanism involved in long-distance transport $(\rightarrow p, 24)$.





Diffusion is movement of a substance owing to the random thermal motion (brownian movement) of its molecules or ions (→A1) in all directions throughout a solvent. Net diffusion or selective transport can occur only when the solute concentration at the starting point is higher than at the target site. (Note: unidirectional fluxes also occur in absence of a concentration gradient—i.e., at equilibrium—but net diffusion is zero because there is equal flux in both directions.) The driving force of diffusion is, therefore, a concentration gradient. Hence, diffusion equalizes concentration differences and requires a driving force: passive transport (= downhill transport).

Example: When a layer of O_2 gas is placed on water, the O_2 quickly diffuses into the water along the initially high gas pressure gradient (\rightarrow **A2**). As a result, the partial pressure of O_2 (Po_2) rises, and O_2 can diffuse further downward into the next O_2 -poor layer of water (\rightarrow **A1**). (*Note:* with gases, partial pressure is used in lieu of concentration.) However, the steepness of the Po_2 profile or gradient (dPo_2/dx) decreases (exponentially) in each subsequent layer situated at distance x from the O_2 source (\rightarrow **A3**). Therefore, diffusion is only feasible for transport across short distances within the body. Diffusion in liquids is slower than in gases.

The diffusion rate, $J_{\rm diff}$ (mol·s⁻¹), is the amount of substance that diffuses per unit of time. It is proportional to the area available for diffusion (A) and the absolute temperature (T) and is inversely proportional to the viscosity (η) of the solvent and the radius (r) of the diffused particles.

According to the Stokes–Einstein equation, the coefficient of diffusion (D) is derived from T, η , and r as

$$D = \frac{R \cdot T}{N_A \cdot 6\pi \cdot r \cdot \eta} [m^2 \cdot s^{-1}], \qquad [1.1]$$

where R is the general gas constant $(8.3144\,\mathrm{J}\cdot\mathrm{K}^{-1}\cdot\mathrm{mol}^{-1})$ and N_A Avogadro's constant $(6.022\cdot10^{23}\,\mathrm{mol}^{-1})$. In Fick's first law of diffusion (Adolf Fick, 1855), the diffusion rate is expressed as

$$J_{diff} = A \cdot D \cdot \left(\frac{dC}{dx}\right) [mol \cdot s^{-1}] \eqno{[1.2]}$$

where C is the molar concentration and x is the distance traveled during diffusion. Since the driving "force"—i.e., the concentration gradient (dC/dx)—decreases with distance, as was explained above, the *time* required for diffusion increases exponentially with the distance traveled $(t \sim x^2)$. If, for example, a molecule travels the first μ m in 0.5 ms, it will require 5 s to travel 100 μ m and a whopping 14 h for 1 cm.

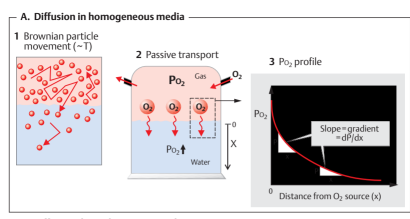
Returning to the previous example $(\rightarrow$ **A2**), if the above-water partial pressure of free O_2 diffusion $(\rightarrow$ **A2**) is kept constant, the Po_2 in the water and overlying gas layer will eventually equalize and net diffusion will cease (diffusion equilibrium). This process takes place within the body, for example, when O_2 diffuses from the alveoli of the lungs into the bloodstream and when CO_2 diffuses in the opposite direction $(\rightarrow$ p. 120).

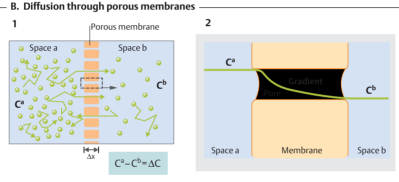
Let us imagine two spaces, a and b $(\rightarrow B1)$ containing different concentrations $(C_a > C_b)$ of an uncharged solute. The membrane separating the solutions has pores Δx in length and with total cross-sectional area of A. Since the pores are permeable to the molecules of the dissolved substance, the molecules will diffuse from a to b, with $C^a - C^b = \Delta C$ representing the concentration gradient. If we consider only the spaces a and b (while ignoring the gradients dC/dx in the pore, as shown in B2, for the sake of simplicity), Fick's first law of diffusion (Eq. 1.2) can be modified as follows:

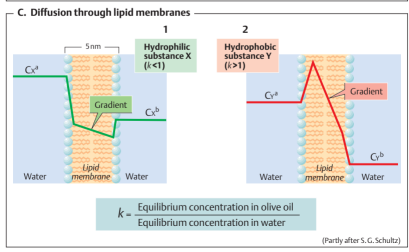
$$\textbf{J}_{\text{diff}} = \textbf{A} \cdot \textbf{D} \cdot \frac{\Delta \textbf{C}}{\Delta \textbf{x}} \left[mol \cdot s^{-1} \right]. \tag{1.3}$$

In other words, the rate of diffusion increases as A, D, and ΔC increase, and decreases as the thickness of the membrane (Δx) decreases.

When diffusion occurs through the **lipid membrane** of a cell, one must consider that hydrophilic substances in the membrane are sparingly soluble (compare intramembrane gradient in C1 to C2) and, accordingly, have a hard time penetrating the membrane by means of "simple" diffusion. The oil-and-water partition coefficient (k) is a measure of the lipid solubility of a substance $(\rightarrow C)$.







The higher the *k* value, the more quickly the substance will diffuse through a pure phospholipid bilayer membrane. Substitution into Eq. 1.3 gives

$$J_{\text{diff}} = k \cdot A \cdot D \cdot \frac{\Delta C}{\Delta x} [\text{mol} \cdot \text{s}^{-1}];$$
 [1.4]

Whereas the molecular radius $r (\rightarrow \text{Eq. 1.1})$ still largely determines the magnitude of D when k remains constant (cf. diethylmalonamide with ethylurea in **D**), k can vary by many powers of ten when r remains constant (cf. urea with ethanol in **D**) and can therefore have a decisive effect on the permeability of the membrane.

Since the value of the variables k, D, and Δx within the body generally cannot be determined, they are usually summarized as the permeability coefficient P, where

$$P = k \cdot \frac{D}{\Delta x} [m \cdot s^{-1}].$$
 [1.5]

If the diffusion rate, J_{diff} [mol·s⁻¹], is related to area A, Eq. 1.4 is transformed to yield

$$\frac{\mathbf{J}_{\text{diff}}}{\mathbf{A}} = \mathbf{P} \cdot \Delta \mathbf{C} \left[mol \cdot m^{-2} \cdot s^{-1} \right]. \tag{1.6}$$

The quantity of substance (net) diffused per unit area and time is therefore proportional to ΔC and $P (\rightarrow E$, blue line with slope P).

When considering the **diffusion of gases**, ΔC in Eq. 1.4 is replaced by $\alpha \cdot \Delta P$ (solubility coefficient times partial pressure difference; \rightarrow p. 126) and J_{diff} [mol·s⁻¹] by \dot{V}_{diff} [m³·s⁻¹], $k \cdot \alpha \cdot D$ is then summarized as diffusion conductance, or *Krogh's diffusion coefficient* K [m²·s⁻¹·Pa⁻¹]. Substitution into Fick's first diffusion equation yields

$$\frac{\dot{V}_{\text{diff}}}{A} = K \cdot \frac{\Delta P}{\Delta x} [m \cdot s^{-1}]. \qquad [1.7]$$

Since A and Δx of alveolar gas exchange (\rightarrow p. 120) cannot be determined in living organisms, $K \cdot F/\Delta x$ for O_2 is often expressed as the O_2 diffusion capacity of the lung, D_L :

$$\dot{V}_{O_{2 \, diff}} = D_L \cdot \Delta P_{O_2} \, [m^3 \cdot s^{-1}]. \eqno(1.8)$$

Nonionic diffusion occurs when the uncharged form of a weak base (e.g., ammonia = NH_3) or acid (e.g., formic acid, HCOOH) passes through a membrane more readily than the charged form (\rightarrow **F**). In this case, the membrane would be more permeable to NH_3 than to NH_4 ⁺

 $(\rightarrow$ p. 176 ff.). Since the pH of a solution determines whether these substances will be charged or not (pK value; \rightarrow p. 378), the diffusion of weak acids and bases is clearly dependent on the pH.

The previous equations have not made allowances for the diffusion of electrically charged particles (ions). In their case, the electrical potential difference at cell membranes must also be taken into account. The electrical potential difference can be a driving force of diffusion (electrodiffusion). In that case, positively charged ions (cations) will then migrate to the negatively charged side of the membrane, and negatively charged ions (anions) will migrate to the positively charged side. The prerequisite for this type of transport is, of course, that the membrane contain ion channels (\rightarrow p. 32 ff.) that make it permeable to the transported ions. Inversely, every ion diffusing along a concentration gradient carries a charge and thus creates an electric diffusion potential $(\rightarrow p.32 \text{ ff.}).$

As a result of the electrical charge of an ion, the permeability coefficient of the ion x (= P_x) can be transformed into the **electrical conductance** of the membrane for this ion, \mathbf{g}_x (\rightarrow \mathbf{p} . 32):

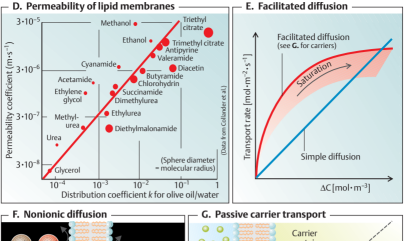
$$g_x = \cdot P_x \cdot z_x^2 \cdot F^2 R^{-1} \cdot T^{-1} \cdot \overline{c}_x [S \cdot m^{-2}]$$
 [1.9]

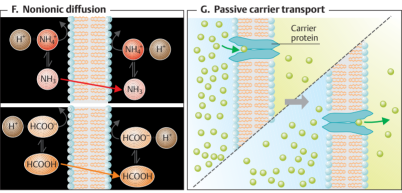
where R and T have their usual meaning (explained above) and z_x equals the charge of the ion, F equals the Faraday constant (9,65 \cdot 10⁴ A \cdot s \cdot mol⁻¹), and \bar{c}_x equals the mean ionic activity in the membrane. Furthermore,

$$\bar{c} = \frac{c_1 - c_2}{\ln c_1 - \ln c_2}.$$
 [1.10]

where index 1 = one side and index 2 = the other side of the membrane. Unlike P, g is concentration-dependent. If, for example, the extracellular K^+ concentration rises from 4 to 8 mmol/kg H_2O (cytosolic concentration remains constant at 160 mmol/kg H_2O), \bar{c} will rise, and g will increase by 20%.

Since most of the biologically important substances are so polar or lipophobic (*small k value*) that simple diffusion of the substances through the membrane would proceed much too slowly, other membrane transport proteins called **carriers** or transporters exist in addition to ion channels. Carriers bind the target molecule (e.g., glucose) on one side of the membrane and detach from it on the other side





(after a conformational change) (\rightarrow **G**). As in simple diffusion, a concentration gradient is necessary for such carrier-mediated transport (passive transport), e.g., with GLUT uniporters for glucose (\rightarrow p. 158). On the other hand, this type of "facilitated diffusion" is subject to *satu*-

ration and is specific for structurally similar substances that may *competitively inhibit* one another. The carriers in both passive and active transport have the latter features in common $(\rightarrow p, 26)$.

Osmosis. Filtration and Convection

Water flow or volume flow (Iv) across a membrane, in living organisms is achieved through osmosis (diffusion of water) or filtration. They can occur only if the membrane is water-permeable. This allows osmotic and hydrostatic pressure differences ($\Delta \pi$ and ΔP) across the membrane to drive the fluids through it.

Osmotic flow equals the hydraulic conductivity (K_f) times the osmotic pressure difference $(\Delta \pi)$ $(\rightarrow A)$:

$$\textbf{J}_{\text{V}} = \textbf{K}_{\text{f}} \cdot \Delta \pi \tag{1.11}$$
 The osmotic pressure difference (\$\Delta \pi\$) can be

calculated using van't Hoff's law, as modified by Staverman:

$$\Delta \pi = \sigma \cdot R \cdot T \cdot \Delta C_{osm}$$
, [1.12]
where σ is the reflection coefficient of the parcles (see below). R is the universal gas con-

where σ is the reflection coefficient of the particles (see below). R is the universal gas constant (\rightarrow p. 20), T is the absolute temperature, and ΔC_{osm} [osm · kgH₂O⁻¹] is the difference between the lower and higher particle concentrations, $C_{osm}^a - C_{osm}^b$ ($\rightarrow A$). Since ΔC_{osm} , the driving force for osmosis, is a negative value, Jv is also negative (Eq. 1.11). The water therefore flows against the concentration gradient of the solute particles. In other words, the higher concentration, C_{osm}, attracts the water. When the concentration of water is considered in osmosis, the H_2O concentration in A,a, $C_{H_2O}^a$, is greater than that in **A,b**, $C_{H_2O}^b$. $C_{H_2O}^a - C_{H_2O}^b$ is therefore the driving force for H₂O diffusion $(\rightarrow A)$. Osmosis also cannot occur unless the reflection coefficient is greater than zero $(\sigma > 0)$, that is, unless the membrane is less permeable to the solutes than to water.

Aquaporins (AQP) are water channels that permit the passage of water in many cell membranes. A chief cell in the renal collecting duct contains a total of ca. 107 water channels, comprising AOP2 (regulated) in the luminal membrane, and AOP3 and 4 (permanent?) in the basolateral membrane. The permeability of the epithelium of the renal collecting duct to water (\rightarrow A, right panel) is controlled by the insertion and removal of AQP2, which is stored in the membrane of intracellular vesicles. In the presence of the antidiuretic hormone ADH (V2 receptors, cAMP; \rightarrow p. 274), water channels are inserted in the luminal membrane within minutes, thereby increasing the water permeability of the membrane to around 1.5×10^{-17} L s⁻¹ per channel.

In filtration $(\rightarrow B)$,

therefore calculated as

 $I_V = K_f \cdot \Delta P - \Delta \pi$ [1.13] Filtration occurs through capillary walls, which allow the passage of small ions and molecules ($\sigma = 0$; see below), but not of plasma proteins $(\rightarrow \mathbf{B}, \text{ molecule } \mathbf{x})$. Their concentration difference leads to an oncotic pressure

difference $(\Delta \pi)$ that opposes ΔP . Therefore, filtration can occur only if $\Delta P > \Delta \pi$ ($\rightarrow B$, p. 152. p. 208).

Solvent drag occurs when solute particles are carried along with the water flow of osmosis or filtration. The amount of solvent drag for solute X (J_X) depends mainly on osmotic flow (J_V) and the mean solute activity $\overline{a}_x (\rightarrow p.376)$ at the site of penetration, but also on the degree of particle reflection from the membrane, which is described using the reflection **coefficient** (σ). Solvent drag for solute X (J_X) is

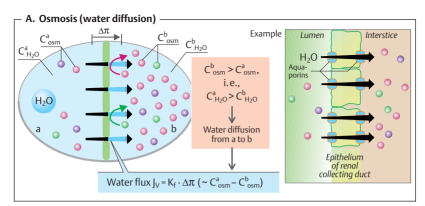
 $I_x = I_V (1 - \sigma) \overline{a}_x [\text{mol} \cdot \text{s}^{-1}]$ [1.14] Larger molecules such as proteins are entirely reflected, and $\sigma = 1 (\rightarrow \mathbf{B}, \text{ molecule X})$. Reflection of smaller molecules is lower, and σ < 1. When urea passes through the wall of the proximal renal tubule, for example, σ = 0.68. The value $(1-\sigma)$ is also called the sieving coefficient (\rightarrow p. 154).

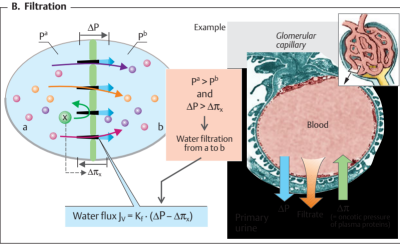
Plasma protein binding occurs when smallmolecular substances in plasma bind to proteins $(\rightarrow \mathbf{C})$. This hinders the free penetration of the substances through the endothelium or the glomerular filter (\rightarrow p. 154 ff.). At a glomerular filtration fraction of 20%, 20% of a freely filterable substance is filtered out. If, however, 9/10 of the substance is bound to plasma proteins, only 2% will be filtered during each renal pass.

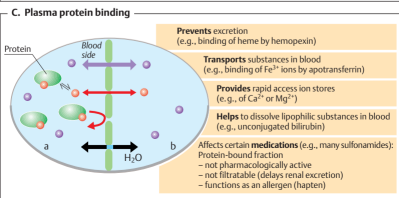
Convection functions to transport solutes over long distances-e.g., in the circulation or urinary tract. The solute is then carried along like a piece of driftwood. The quantity of solute transported over time (Jconv) is the product of volume flow J_V (in $m^3 \cdot s^{-1}$) and the solute concentration C (mol · m⁻³):

 $J_{conv} = J_{v} \cdot C [mol \cdot s^{-1}].$ [1.15]

The flow of gases in the respiratory tract, the transmission of heat in the blood and the release of heat in the form of warmed air occurs through convection (\rightarrow p. 222).







Active transport occurs in many parts of the body when solutes are transported against their concentration gradient (uphill transport) and/or, in the case of ions, against an electrical potential (\rightarrow p. 22). All in all, active transport occurs against the electrochemical gradient or potential of the solute. Since passive transport mechanisms represent "downhill" transport $(\rightarrow p.20 \text{ ff.})$, they are not appropriate for this task. Active transport requires the expenditure of energy. A large portion of chemical energy provided by foodstuffs is utilized for active transport once it has been made readily available in the form of ATP (\rightarrow p. 41). The energy created by ATP hydrolysis is used to drive the transmembrane transport of numerous ions. metabolites, and waste products. According to the laws of thermodynamics, the energy expended in these reactions produces order in cells and organelles-a prerequisite for survival and normal function of cells and, therefore, for the whole organism (\rightarrow p. 38 ff.).

In **primary active transport**, the energy produced by hydrolysis of ATP goes *directly* into ion transport through an ion pump. This type of ion pump is called an **ATPase**. They establish the electrochemical gradients rather slowly, e.g., at a rate of around $1 \, \mu \text{mol} \cdot \text{s}^{-1} \cdot \text{m}^{-2}$ of membrane surface area in the case of Na*-K*-ATPase. The gradient can be exploited to achieve *rapid ionic currents* in the opposite direction after the permeability of ion channels has been increased (\rightarrow p. 32 ff.). Na* can, for example, be driven into a nerve cell at a rate of up to $1000 \, \mu \text{mol} \cdot \text{s}^{-1} \cdot \text{m}^{-2}$ during an action potential.

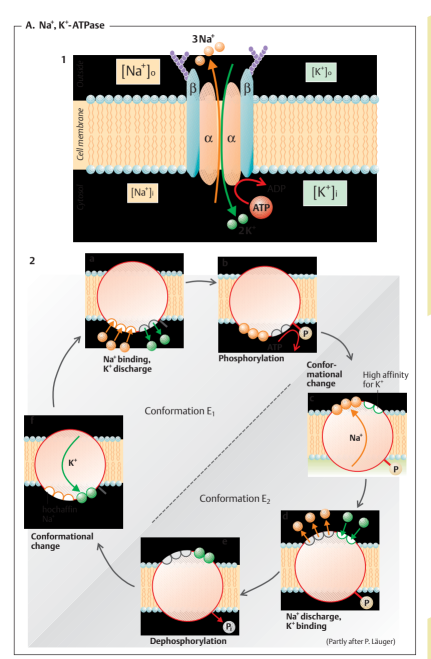
ATPases occur ubiquitously in cell membranes (Na⁺-K⁺-ATPase) and in the endoplasmic reticulum and plasma membrane (Ca²⁺-ATPase), renal collecting duct and stomach glands (H⁺,K⁺ -ATPase), and in lysosomes (H⁺-ATPase). They transport Na⁺, K⁺, Ca²⁺ and H⁺, respectively, by primarily active mechanisms. All except H⁺-ATPase consist of 2 α-subunits and 2 β-subunits (P-type ATPases). The α-subunits are phosphorylated and form the ion transport channel (\rightarrow **A1**).

Na⁺-K⁺-ATPase is responsible for maintenance of intracellular Na⁺ and K⁺ homeostasis

and, thus, for maintenance of the cell membrane potential. During each transport cycle $(\rightarrow A1, A2)$, 3 Na⁺ and 2 K⁺ are "pumped" out of and into the cell, respectively, while 1 ATP molecule is used to phosphorylate the carrier protein $(\rightarrow A2b)$. Phosphorylation changes the conformation of the protein and subsequently alters the affinities of the Na⁺ and K+ binding sites. The conformational change is the actual ion transport step since it moves the binding sites to the opposite side of the membrane $(\rightarrow A2b-d)$. Dephosphorylation restores the pump to its original state $(\rightarrow A2e-f)$. The Na⁺/K⁺ pumping rate increases when the cytosolic Na+ concentration risesdue, for instance, to increased Na+ influx, or when the extracellular K+ rises. Therefore, Na+.K+-activatable ATPase is the full name of the pump. Na-+K+-ATPase is inhibited by ouabain and cardiac glycosides.

Secondary active transport occurs when uphill transport of a compound (e.g., glucose) via a carrier protein (e.g., sodium glucose transporter type 2, SGLT2) is coupled with the passive (downhill) transport of an ion (in this example Na⁺; \rightarrow **B1**). In this case, the electrochemical Na+ gradient into the cell (created by Na+-K+-ATPase at another site on the cell membrane; \rightarrow **A**) provides the driving force needed for secondary active uptake of glucose into the cell. Coupling of the transport of two compounds across a membrane is called cotransport, which may be in the form of symport or antiport. Symport occurs when the two compounds (i.e., compound and driving ion) are transported across the membrane in the same direction (\rightarrow B1-3). Antiport (countertransport) occurs when they are transported in opposite directions. Antiport occurs, for example, when an electrochemical Na+ gradient drives H⁺ in the opposite direction by secondary active transport (\rightarrow **B4**). The resulting H⁺ gradient can then be exploited for tertiary active symport of molecules such as peptides (\rightarrow **B5**).

Electroneutral transport occurs when the net electrical charge remains balanced during transport, e.g., during Na⁺/H⁺ antiport (\rightarrow B4) and Na⁺-Cl⁻ symport (\rightarrow B2). Small charge separation occurs in **electrogenic (rheogenic) transport**, e.g., in Na⁺-glucose⁰ symport (\rightarrow B1), Na⁺-amino acid⁰ symport (\rightarrow B3),



2 Na⁺-amino acid⁻ symport, or H⁺-peptide⁰ symport (\rightarrow **B5**). The chemical Na⁺ gradient provides the sole driving force for electroneutral transport (e.g., Na+/H+ antiport). whereas the negative membrane potential (→ p. 32 ff.) provides an additional driving force for electrogenication-coupled cotransport into the cell. When secondary active transport (e.g., of glucose) is coupled with the influx of not one but two Na+ ions (e.g., SGLT1 symporter), the driving force is doubled. The aid of ATPases is necessary, however, if the required "uphill" concentration ratio is several decimal powers large, e.g., 106 in the extreme case of H⁺ ions across the luminal membrane of parietal cells in the stomach, ATPase-mediated transport can also be electrogenic or electroneutral, e.g., Na+-K+-ATPase (3 Na+/2 K+: cf. p. 46) or H+/K+-ATPase (1 H+/1 K+), respectively. Characteristics of active transport:

- ◆ It can be saturated, i.e., it has a limited *maximum capacity* (J_{max}).
- ◆ It is more or less *specific*, i.e., a carrier molecule will transport only certain chemically similar substances which inhibit the transport of each other (*competitive inhibition*).
- lack Variable quantities of the similar substances are transported at a given concentration, i.e., each has a different *affinity* (~1/K_M) to the transport system.
- Active transport is inhibited when the *energy supply* to the cell is disrupted.

All of these characteristics except the last apply to passive carriers, that is, to uniporter-mediated (facilitated) diffusion (\rightarrow p. 22).

The **transport rate** of saturable transport (J_{sat}) is usually calculated according to Michaelis–Menten kinetics:

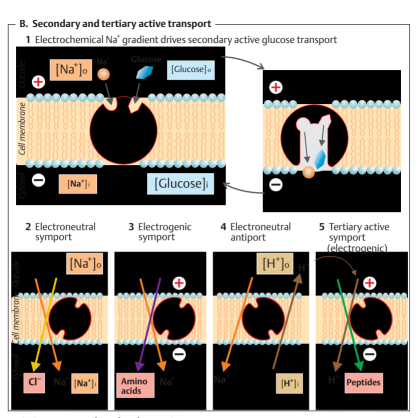
$$J_{\text{sat}} = J_{\text{max}} \cdot \frac{C}{K_{\text{M}} + C}$$
 [1.16]

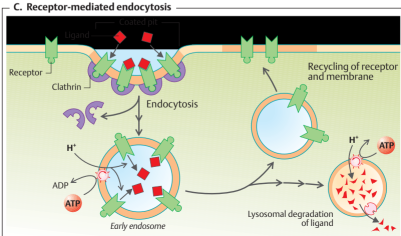
where C is the concentration of the substrate in question, J_{max} is its maximum transport rate, and K_M is the substrate concentration that produces one-half J_{max} (\rightarrow p. 383).

Cytosis is a completely different type of active transport involving the formation of membrane-bound vesicles with a diameter of 50–400 nm. **Vesicles** are either pinched off from the plasma membrane (*exocytosis*) or incorporated into it by invagination (*endocyto-*

sis) in conjunction with the expenditure of ATP. In cytosis, the uptake and release of macromolecules such as proteins, lipoproteins, polynucleotides, and polysaccharides into and out of a cell occurs by specific mechanisms similar to those involved in intracellular transport $(\rightarrow p.12 \, \text{ff.})$.

Endocytosis (\rightarrow p. 13) can be broken down into different types, including pinocytosis, receptor-mediated endocytosis, and phagocytosis. Pinocytosis is characterized by the continuous unspecific uptake of extracellular fluid and molecules dissolved in it through relatively small vesicles. Receptor-mediated en**docytosis** $(\rightarrow C)$ involves the selective uptake of specific macromolecules with the aid of receptors. This usually begins at small depressions (pits) on the plasma membrane surface. Since the insides of the pits are often densely covered with the protein clathrin, they are called clathrin-coated pits. The receptors involved are integral cell membrane proteins such as those for low-density lipoprotein (LPL; e.g., in hepatocytes) or intrinsic factor-bound cobalamin (e.g., in ileal epithelial cells). Thousands of the same receptor type or of different receptors can converge at coated pits $(\rightarrow C)$, vielding a tremendous increase in the efficacy of ligand uptake. The endocytosed vesicles are initially coated with clathrin, which is later released. The vesicles then transform into early endosomes, and most of the associated receptors circulate back to the cell membrane ($\rightarrow C$ and p. 13). The endocytosed ligand is either exocytosed on the opposite side of the cell (transcytosis, see below), or is digested by lysosomes (\rightarrow C and p. 13). Phagocytosis involves the endocytosis of particulate matter, such as microorganisms or cell debris, by phagocytes $(\rightarrow p.94 \, \text{ff.})$ in conjunction with lysosomes. Small digestion products, such as amino acids. sugars and nucleotides, are transported out of the lysosomes into the cytosol, where they can be used for cellular metabolism or secreted into the extracellular fluid. When certain hormones such as insulin (→ p.282) bind to receptors on the surface of target cells, hormonereceptor complexes can also enter the coated pits and are endocytosed (internalized) and digested by lysosomes. This reduces the density of receptors available for hormone bind-





ing. In other words, an increased hormone supply down-regulates the receptor density.

Exocytosis (\rightarrow p. 13) is a method for selective export of macromolecules out of the cell (e.g., pancreatic enzymes; \rightarrow p. 246 ff.) and for release of many hormones (e.g., posterior pituitary hormone; → p. 280) or neurotransmitters (\rightarrow p. 50 ff.). These substances are kept "packed" and readily available in (clathrin-coated) secretory vesicles, waiting to be released when a certain signal is received (increase in cytosolic Ca2+). The "packing material" (vesicle membrane) is later re-endocytosed and recycled. Exocytotic membrane fusion also helps to insert vesicle-bound proteins into the plasma membrane (\rightarrow p. 13). The liquid contents of the vesicle then are automatically emptied in a process called constitutive exocytosis $(\rightarrow D)$.

In constitutive exocytosis, the protein complex coatomer (coat assembly protomer) takes on the role of clathrin (see above). Within the Golgi membrane, GNRP (quanine nucleotide-releasing protein) phosphorylates the GDP of the ADP-ribosylation factor (ARF) to GTP \rightarrow **D1**), resulting in the dispatch of vesicles from the trans-Golgi network. ARF-GTP complexes then anchor on the membrane and bind coatomer (→ **D2**), thereby producina **coatomer-coated vesicles** (\rightarrow **D3**). The membranes of the vesicles contain v-SNAREs (vesicle synaptosome-associated protein receptors), which recognize t-SNAREs (target-SNAREs) in the target membrane (the plasma membrane, in this case). This results in cleavage of ARF-GTP, dissociation of ARF-GDP and coatomer molecules and, ultimately, to membrane fusion and exocytosis (→ D4, D5) to the extracellular space (ECS).

Transcytosis is the uptake of macromolecules such as proteins and hormones by endocytosis on one side of the cell, and their release on the opposite side. This is useful for *transcellular transport of the macromolecules* across cell layers such as endothelia.

Cell Migration

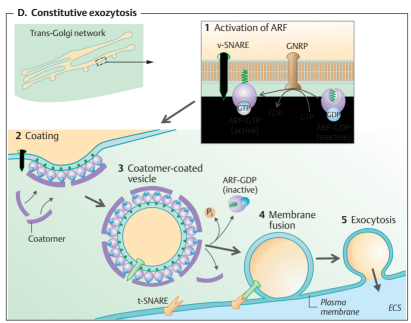
Most cells in the body are theoretically able to move from one place to another or *migrate* $(\rightarrow \mathbf{E})$, but only a few cell species actually do so. The *sperm* are probably the only cells with a special propulsion mechanism. By waving their whip-like tail, the sperm can travel at

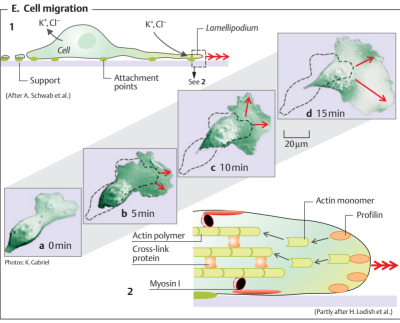
speeds of up to around $2000\,\mu\text{m/min}$. Other cells also migrate, but at much slower rates. Fibroblasts, for example, move at a rate of around 1.2 $\mu\text{m/min}$. When an injury occurs, fibroblasts migrate to the wound and aid in the formation of scar tissue. Cell migration also plays a role in embryonal development. Chemotactically attracted neutrophil granulocytes and macrophages can even migrate through vessel walls to attack invading bacteria (\rightarrow p. 94ff.). Cells of some tumors can also migrate to various tissues of the body or metastasize, thereby spreading their harmful effects.

Cells migrate by "crawling" on a stable surface (\rightarrow E1). The following activities occur during cell migration:

- Back end of the cell: (a) Depolymerization of actin and tubulin in the cytoskeleton; (b) endocytosis of parts of the cell membrane, which are then propelled forward as endocytotic vesicles to the front of the cell, and (c) release of ions and fluids from the cell.
- ◆ Front end of the cell (lamellipodia): (a) Polymerization of actin monomers is achieved with the aid of profilin (→ E2). The monomers are propelled forward with the help of plasma membrane-based myosin I (fueled by ATP); (b) reinsertion of the vesicles in the cell membrane; (c) uptake of ions and fluids from the environment.

Parts of the cell membrane that are not involved in cytosis are conveyed from front to back, as on a track chain. Since the cell membrane is attached to the stable surface (primarily fibronectin of the extracellular matrix in the case of fibroblasts), the cell moves forward relative to the surface. This is achieved with the aid of specific receptors, such as fibronectin receptors in the case of fibroblasts.





Electrical Membrane Potentials and Ion Channels

An electrical potential difference occurs due to the net movement of charge during ion transport. A diffusion potential develops for instance, when ions (e.g., K+) diffuse (down a chemical gradient; \rightarrow p. 20ff.) out of a cell, making the cell interior negative relative to the outside. The rising diffusion potential then drives the ions back into the cell (potentialdriven ion transport; → p. 22). Outward K⁺ diffusion persists until equilibrium is reached. At equilibrium, the two opposing forces become equal and opposite. In other words, the sum of the two or the electrochemical gradient (and thereby the electrochemical potential) equals zero, and there is no further net movement of ions (equilibrium concentration) at a certain voltage (equilibrium potential).

The **equilibrium potential** (E_X) for any species of ion X distributed inside (i) and outside (o) a cell can be calculated using the Nernst equation:

$$E_X = \frac{R \cdot T}{F \cdot z_x} \cdot \ln \frac{[X]_o}{[X]_i} [V]$$
 [1.17]

where R is the universal gas constant (= 8.314 $J \cdot K^{-1} \cdot mol^{-1}$), T is the absolute temperature (310 °K in the body), F is the Faraday constant or charge per mol (= 9.65 × 10⁴ A · s · mol^{-1}), z is the valence of the ion in question (+1 for K⁺, +2 for Ca²⁺, -1 for Cl⁻, etc.), In is the natural logarithm, and [X] is the effective concentration = activity of the ion X (\rightarrow p. 376). R·T/F = 0.0267 V⁻¹ at body temperature (310 °K). It is sometimes helpful to convert $\ln([X]_o/[X]_i)$ into $-\ln([X]_i/[X]_o)$, V into mV and In into log before calculating the equilibrium potential (\rightarrow p. 380). After insertion into Eq. 1.17, the Nernst equation then becomes

$$E_X = -61 \cdot \frac{1}{z_X} \cdot log \frac{[X]_i}{[X]_o} [mV] \qquad [1.18]$$

If the ion of species X is K*, and $[K^*]_i = 140$, and $[K^*]_o = 4.5 \text{ mmol/kg H}_2O$, the equilibrium potential $E_K = -61 \cdot \log 31 \text{ mV or } -91 \text{ mV}$. If the cell membrane is permeable only to K^* , the **membrane potential (E_m)** will eventually reach a value of -91 mV, and $E_m \text{ will equal } E_K (\rightarrow \textbf{A1})$.

At equilibrium potential, the chemical gradient will drive just as many ions of species X in the one direction as the electrical potential does in the opposite direction. The **electrochemical potential** (E_m-E_k) or so-called electrochemical driving "force", will equal zero, and the sum of ionic inflow and outflow or the **net flux** (I_k) will also equal zero.

Membrane conductance (g_x), a concentration-dependent variable, is generally used to describe the permeability of a cell membrane to a given ion instead of the permeability coefficient P (see Eq. 1.5 on p.22 for conversion). Since it is relative to membrane surface area, g_x is expressed in siemens (S = 1/Ω) per $m^2(\rightarrow p.22, Eq. 1.9)$. Ohm's law defines the net ion current (I_x) per unit of membrane surface area as

$$\mathbf{I}_{X} = \mathbf{g}_{X} \cdot (\mathbf{E}_{m} - \mathbf{E}_{X}) [\mathbf{A} \cdot \mathbf{m}^{-2}]$$
 [1.19]

 I_x will therefore differ from zero when the prevailing membrane potential, E_m , does not equal the equilibrium potential, E_x . This occurs, for example, after strong transient activation of Na^+ -K $^+$ -ATPase (electrogenic; \rightarrow p. 26): hyperpolarization of the membrane (\rightarrow A2), or when the cell membrane conducts more than one ion species, e.g., K^+ as well as CI^- and Na^+ : depolarization (\rightarrow A3). If the membrane is permeable to different ion species, the total conductance of the membrane (g_m) equals the sum of all parallel conductances ($g_1 + g_2 + g_3 + ...$). The **fractional conductance** for the ion species X (f_x) can be calculated as

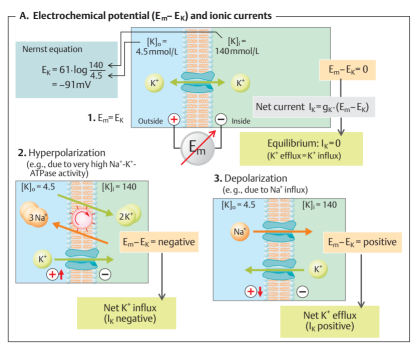
$$f_X = g_X/g_m \qquad [1.20]$$

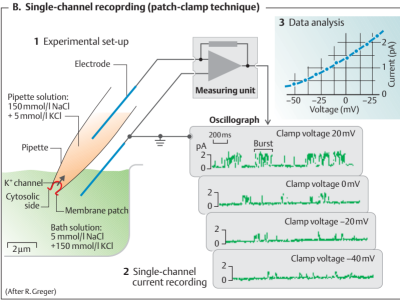
The membrane potential, E_m , can be determined if the fractional conductances and equilibrium potentials of the conducted ions are known (see Eq. 1.18). Assuming K^+ , Na^+ , and Cl^- are the ions in question,

$$E_m = (E_K \cdot f_K) + (E_{Na} \cdot f_{Na}) + (E_{Cl} \cdot f_{Cl}) \qquad \text{[1.21]}$$

Realistic values in resting nerve cells are: $f_K = 0.90$, $f_{Na} = 0.03$, $f_{Cl} = 0.07$; $E_K = -90$ mV, $E_{Na} = +70$ mV, $E_{Cl} = -83$ mV. Inserting these values into equation 1.21 results in an E_m of -85 mV. Thus, the driving forces (= electrochemical potentials = $E_m - E_x$), equal +5 mV for K^+ , -145 mV for Na^+ , and -2 mV for Cl^- . The driv-

▶





ing force for K^+ efflux is therefore low, though g_K is high. Despite a high driving force for Na^+ , Na^+ influx is low because the g_{Na} and f_{Na} of resting cells are relatively small. Nonetheless, the sodium current, I_{Na} , can rise tremendously when large numbers of Na^+ channels open during an action potential (\rightarrow p. 46).

Electrodiffusion. The potential produced by the transport of one ion species can also drive other cations or anions through the cell membrane (\rightarrow p. 22), provided it is permeable to them. The K⁺diffusion potential leads to the efflux of Cl-, for example, which continues until $E_{Cl} = E_{m}$. According to Equation 1.18, this means that the cytosolic Cl- concentration is reduced to 1/25 th of the extracellular concentration (passive distribution of Cl- between cytosol and extracellular fluid). In the above example. there was a small electrochemical Cl-potential driving Cl^- out of the cell $(E_m - E_{Cl} = -2 \text{ mV})$. This means that the cytosolic Cl⁻ concentration is higher than in passive Cl^- distribution (E_{Cl} = E_m). Therefore, Cl⁻ ions must have been actively taken up by the cell, e.g., by a Na+- Clsymport carrier (\rightarrow p. 29 B): active distribution of Cl-.

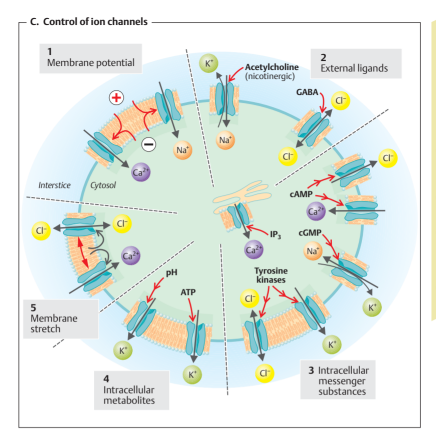
To achieve ion transport, membranes have a variable number of channels (pores) specific for different ion species (Na+, Ca2+, K+, Cl-, etc.). The conductance of the cell membrane is therefore determined by the type and number of ion channels that are momentarily open. Patch-clamp techniques permit the direct measurement of ionic currents through single ion channels (\rightarrow **B**). Patch-clamp studies have shown that the conductance of a membrane does not depend on the change of the pore diameter of its ion channels, but on their average frequency of opening. The ion permeability of a membrane is therefore related to the open-probability of the channels in question. Ion channels open in frequent bursts (\rightarrow **B2**). Several ten thousands of ions pass through the channel during each individual burst, which lasts for only a few milliseconds.

During a **patch–clamp recording**, the opening $(0.3–3 \, \mu m$ in diameter) of a glass electrode is placed over a cell membrane in such a way that the opening covers only a small part of the membrane (*patch*) containing only one or a small number of ion channels. The whole cell

can either be left intact, or a membrane patch can excised for isolated study (\rightarrow **B1**). In singlechannel recording, the membrane potential is kept at a preset value (voltage clamp). This permits the measurement of ionic current in a single channel. The measurements are plotted $(\rightarrow B3)$ as current (I) over voltage (V). The slope of the I/V curve corresponds to the conductance of the channel for the respective ion species (see Eq. 1.18). The zero-current potential is defined as the voltage at which the I/V curve intercepts the x-axis of the curve (I = 0). The ion species producing current I can be deduced from the zero-current potential. In example B, the zero-current potential equals - 90 mV. Under the conditions of this experiment, an electrochemical gradient exists only for Na+ and K^+ , but not for $Cl^-(\rightarrow \mathbf{B})$. At these gradients, $E_K = -90 \text{ mV}$ and $E_{Na} = +90 \text{ mV}$. As E_K equals the zero-current potential, the channel is exclusively permeable to K⁺ and does not allow other ions like Na⁺ to pass. The channel type can also be determined by adding specific channel blockers to the system.

Control of ion channels $(\rightarrow C)$. Channel open-probability is controlled by five main factors:

- ♦ *Membrane potential*, especially in Na^+ , Ca^{2+} and K^+ channels in nerve and muscle fibers (\rightarrow C1; pp. 46 and 50).
- ◆ External ligands that bind with the channel (→ C2). This includes acetylcholine on the postsynaptic membrane of nicotinic synapses (cation channels), glutamate (cation channels), and glycine or GABA (Cl⁻ channels).
- ♦ Intracellular messenger substances $(\rightarrow C3)$ such as:
- cAMP (e.g., in Ca²⁺ channels in myocardial cells and Cl⁻ channels in epithelial cells);
- cGMP (plays a role in muscarinergic effects of acetylcholine and in excitation of the retinal rods);
- IP3 (opening of Ca²⁺ channels of intracellular Ca²⁺ stores);
- Small G-proteins (Ca²⁺ channels of the cell membrane);
- Tyrosine kinases (Cl⁻ and K⁺ channels during apoptosis);
- Ca²⁺ (affects K⁺ channels and degree of activation of rapid Na⁺ channels; → p. 46).



- Intracellular metabolites (→ C4) such as ATP (e.g., in K⁺ channels in the heart and B cells in pancreatic islets) or H+ ions (e.g., in K⁺ channels in renal epithelial cells);
- ♦ *Membrane stretch* (\rightarrow **C5**), the direct or indirect (?) effects of which play a role in Ca²⁺ channels of smooth muscle fibers and generally in normal K⁺ and Cl⁻ channels in swelling cells.

Role of Ca²⁺ in Cell Regulation

The cytosolic Ca²⁺ concentration, [Ca²⁺]_i, (ca. 0.1 to 0.01 µmol/L) is several decimal powers lower than the extracellular Ca²⁺ concentration [Ca²⁺]_o (ca. 1.3 mmol/L). This is because Ca²⁺ is continuously pumped from the cytosol into intracellular Ca²⁺ stores such as the endoplasmic and sarcoplasmic reticulum (\rightarrow p. 17 A), vesicles, mitochondria and nuclei (?) or is *transported out of the cell*. Both processes occur by primary active transport (Ca²⁺-ATPases) and, in the case of efflux, by additional secondary active transport through Ca²⁺/3 Na⁺ antiporters (\rightarrow A1).

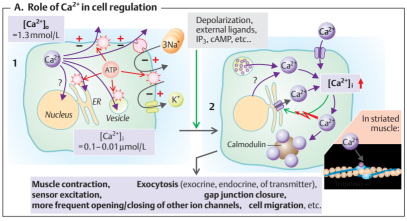
To increase the cytosolic Ca²⁺ concentration, Ca²⁺ channels conduct Ca²⁺ from intracellular stores and the extracellular space into the cytosol (→A2). The frequency of Ca²⁺ channel opening in the cell membrane is increased by *Depolarization* of the cell membrane (nerve and muscle cells);

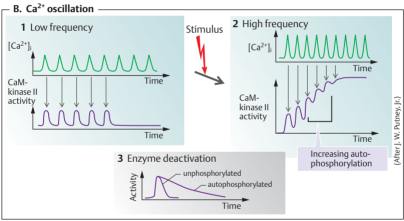
- ♦ *Ligands* (e.g., via G_0 proteins; \rightarrow p. 274);
- ◆ Intracellular messengers (e.g., IP₃ and cAMP; → p. 274ff.);
- ♦ Stretching or heating of the cell membrane. The Ca^{2+} channels of the endoplasmic and sarcoplasmic reticulum open more frequently in response to signals such as a rise in $[Ca^{2+}]_i$ (influx of external Ca^{2+} works as the "spark" or trigger) or inositol *tris*-phosphate (IP₃; → A2 and p. 276).

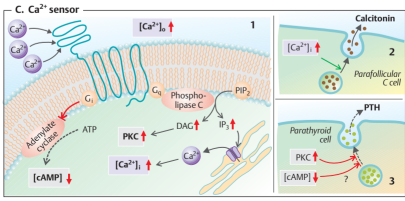
A rise in [Ca2+]; is a signal for many important cell functions $(\rightarrow A)$, including myocyte contraction, exocytosis of neurotransmitters in presynaptic nerve endings, endocrine and exocrine hormone secretion, the excitation of certain sensory cells, the closure of gap junctions in various cells (\rightarrow p. 19 C), the opening of other types of ion channels, and the migration of leukocytes and tumor cells (\rightarrow p. 30) as well as thrombocyte activation and sperm mobilization. Some of these activities are mediated by calmodulin. A calmodulin molecule can bind up to 4 Ca²⁺ ions when the [Ca²⁺]_i rises $(\rightarrow A2)$. The Ca²⁺-calmodulin complexes activate a number of different enzymes, including calmodulin-dependent protein kinase II (CaMkinase II) and myosin light chain kinase (MLCK), which is involved in smooth muscle contraction (\rightarrow p. 70).

[Ca2+]i oscillation is characterized by multiple brief and regular [Ca2+]; increases (Ca2+ spikes) in response to certain stimuli or hormones $(\rightarrow B)$. The frequency, not amplitude, of [Ca²⁺]_i oscillation is the quantitative signal for cell response. When low-frequency [Ca2+]i oscillation occurs, CaM-kinase II, for example, is activated and phosphorylates only its target proteins, but is quickly and completely deactivated (→ B1, B3), High-frequency [Ca2+]; oscillation results in an increasing degree of autophosphorylation and progressively delays the deactivation of the enzyme (\rightarrow **B3**). As a result, the activity of the enzyme decays more and more slowly between [Ca2+]i signals, and each additional [Ca2+]i signal leads to a summation of enzyme activity (\rightarrow **B2**). As with action potentials (\rightarrow p. 46), this frequency-borne, digital all-or-none type of signal transmission provides a much clearer message than the [Ca²⁺]; amplitude, which is influenced by a number of factors.

Ca2+ sensors. The extracellular Ca2+ concentration [Ca²⁺]_o plays an important role in blood coagulation and bone formation as well as in nerve and muscle excitation. [Ca2+]0 is tightly controlled by hormones such as PTH, calcitriol and calcitonin (\rightarrow p. 290), and represents the feedback signal in this control circuit $(\rightarrow p.290)$. The involved Ca²⁺sensors are membrane proteins that detect high [Ca2+]o levels on the cell surface and dispatch IP3 and DAG (diacylglycerol) as intracellular second messengers with the aid of a G_q protein (\rightarrow C1 and p. 274ff.). IP₃ triggers an increase in the [Ca²⁺]_i of parafollicular C cells of the thyroid gland. This induces the exocytosis of calcitonin, a substance that reduces $[Ca^{2+}]_0 (\rightarrow C2)$. In parathyroid cells, on the other hand, a high [Ca2+]0 reduces the secretion of PTH, a hormone that increases the [Ca2+]o. This activity is mediated by DAG and PKC (protein kinase C) and, perhaps, by a (G_i protein-mediated; \rightarrow p. 274) reduction in the cAMP concentration $(\rightarrow C3)$. Ca2+ sensors are also located on osteoclasts as well as on renal and intestinal epithelial cells.







Energy Production and Metabolism

Energy is the ability of a system to perform work: both are expressed in joules (I). A potential difference (potential gradient) is the socalled driving "force" that mobilizes the matter involved in the work. Water falling from height X (in meters) onto a power generator, for example, represents the potential gradient in mechanical work. In electrical and chemical work, potential gradients are provided respectively by voltage (V) and a change in free enthalpy ΔG (J·mol⁻¹). The amount of work performed can be determined by multiplying the potential difference (intensity factor) by the corresponding capacity factor. In the case of the water fall, the work equals the height the water falls (m) times the force of the falling water (in N). In the other examples, the amount work performed equals the voltage (V) times the amount of charge (C). Chemical work performed = ΔG times the amount of substance (mol).

Living organisms cannot survive without an adequate supply of energy. Plants utilize solar energy to convert atmospheric CO2 into oxygen and various organic compounds. These, in turn, are used to fill the energy needs of humans and animals. This illustrates how energy can be converted from one form into another. If we consider such a transformation taking place in a closed system (exchange of energy, but not of matter, with the environment), energy can neither appear nor disappear spontaneously. In other words, when energy is converted in a closed system, the total energy content remains constant. This is described in the first law of thermodynamics. which states that the change of internal energy (= change of energy content, ΔU) of a system (e.g. of a chemical reaction) equals the sum of the work absorbed (+W) or performed (-W) by a system and the heat lost (-Q) or gained (+Q) by the system. This is described as:

 ΔU = heat gained (Q) – work performed (W) [J] and [1.22] ΔU = work absorbed (W) – heat lost (Q) [J]. [1.23] y definition, the signs indicate the direction

(By definition, the signs indicate the direction of flow with respect to the system under consideration.)

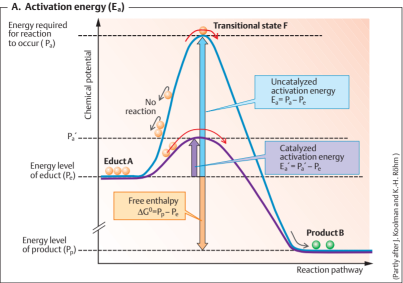
Heat is transferred in all chemical reactions. The amount of heat produced upon conversion of a given substance into product X is the same, *regardless of the reaction pathway* or whether the system is closed or open, as in a biological system. For caloric values, see p. 228.

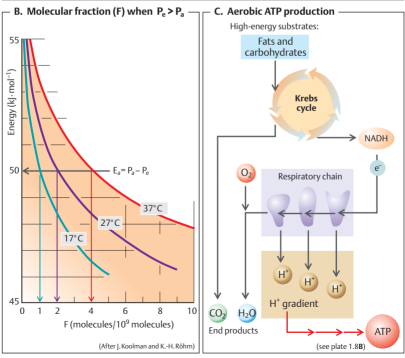
Enthalpy change (\DeltaH) is the heat gained or lost by a system at constant pressure and is related to work, pressure, and volume ($\Delta H = \Delta U$ + p · ΔV). Heat is lost and ΔH is negative in exothermic reactions, while heat is gained and AH is positive in endothermic reactions. The second law of thermodynamics states that the total disorder (randomness) or entropy (S) of a closed system increases in any spontaneous process, i.e., entropy change (ΔS) > 0. This must be taken into consideration when attempting to determine how much of ΔH is freely available. This free energy or free enthalpy (ΔG) can be used, for example, to drive a chemical reaction. The heat produced in the process is the product of absolute temperature and entropy change $(T \cdot \Delta S)$.

Free enthalpy (\DeltaG) can be calculated using the Gibbs-Helmholtz equation:

 $\Delta G = \Delta H - T \cdot \Delta S$. [1.24] ΔG and ΔH are approximately equal when ΔS approaches zero. The maximum chemical work of glucose in the body can therefore be determined based on heat transfer. ΔH . measured during the combustion of glucose in a calorimeter (see p.228 for caloric values). Equation 1.24 also defines the conditions under which chemical reactions can occur. Ex**ergonic** reactions ($\Delta G < 0$) are characterized by the release of energy and can proceed spontaneously, whereas **endergonic** reactions (ΔG > 0) require the absorption of energy and are not spontaneous. An endothermic reaction $(\Delta H > 0)$ can also be exergonic $(\Delta G < 0)$ when the entropy change ΔS is so large that ΔH - $T \cdot \Delta S$ becomes negative. This occurs, for example, in the endothermic dissolution of crystalline NaCl in water.

Free enthalpy, ΔG , is a concentration-dependent variable that can be calculated from the change in **standard free enthalpy** (ΔG^0) and the prevailing concentrations of the substances in question. ΔG^0 is calculated assuming for all reaction partners that concentration = 1 mol/L, pH = 7.0, T = 298 K, and p = 1013 hPa.





Given the reaction

where A is the educt and B and C are the products, ΔG^0 is converted to ΔG as follows:

$$\Delta \mathbf{G} = \Delta \mathbf{G}^{0} + \mathbf{R} \cdot \mathbf{T} \cdot \ln \frac{[\mathbf{B}] + [\mathbf{C}]}{[\mathbf{A}]}$$
 [1.26]

or, at a temperature of 37 °C,

$$\Delta G = \Delta G^0 + 8.31 \cdot 310 \cdot 2.3 \cdot \log \frac{[B] + [C]}{[A]} [J \cdot mol^{-1}]$$
[1.27]

Assuming the ΔG^0 of a reaction is $+20 \text{ kJ} \cdot \text{mol}^{-1}$ (endergonic reaction), ΔG will be exergonic (<0) if [B] · [C] is 10^4 times smaller than A:

$$\Delta G = 20000 + 5925 \cdot \log 10^{-4} = -3.7 \text{ kJ} \cdot \text{mol}^{-1}.$$

In this case, A is converted to B and C and reaction 1.25 proceeds to the right.

If $[B] \cdot [C]/[A] = 4.2 \times 10^{-4}$, ΔG will equal zero and the reaction will come to equilibrium (no net reaction). This numeric ratio is called the **equilibrium constant** (K_{eq}) of the reaction. K_{eq} can be converted to ΔG^0 and vice versa using Equation 1.26:

$$0 = \Delta G^0 + R \cdot T \cdot lnK_{eq} \text{ or}$$

$$\Delta G^0 = -R \cdot T \cdot lnK_{eq} \text{ and}$$
 [1.29]

$$K_{eq} = e^{-\Delta G^{\circ}/(R \cdot T)}.$$
 [1.30]

Conversely, when $[B] \cdot [C]/[A] > 4.2 \times 10^{-4}$, ΔG will be > 0, the net reaction will proceed backwards, and A will arise from B and C.

 ΔG is therefore a measure of the direction of a reaction and of its distance from equilibrium. Considering the concentration-dependency of ΔG and assuming the reaction took place in an open system (see below) where reaction products are removed continuously, e.g., in subsequent metabolic reactions, it follows that ΔG would be a large negative value, and that the reaction would persist without reaching equilibrium.

The magnitude of ΔG^0 , which represents the difference between the energy levels (chemical potentials) of the product P_p and educt P_e (\rightarrow A), does not tell us anything about the **rate of the reaction**. A reaction may be very slow, even if $\Delta G^0 < 0$, because the reaction rate also depends on the energy level (P_a) needed *transiently* to create the necessary transitional state. P_a is higher than P_e (\rightarrow A). The additional amount of energy required to reach this level is

called the **activation energy (E_a)**: $E_a = P_a - P_e$. It is usually so large ($\approx 50 \, \text{kJ} \cdot \text{mol}^{-1}$) that only a tiny fraction ($F \approx 10^{-9}$) of the educt molecules are able to provide it (\rightarrow **A**, **B**). The energy levels of these individual educt molecules are incidentally higher than P_e , which represents the mean value for all educt molecules. The size of **fraction F** is temperature-dependent (\rightarrow **B**). A 10 °C decrease or rise in temperature lowers or raises F (and usually the reaction rate) by a factor of 2 to 4, i.e. the **Q**₁₀ value of the reaction is 2 to 4.

Considering the high Ea values of many noncatalyzed reactions, the development of enzymes as biological catalysts was a very imstep in evolution. Enzymes enormously accelerate reaction rates by lowering the activation energy $E_a (\rightarrow \mathbf{A})$. According to the Arrhenius equation, the rate constant k (s⁻¹) of a unimolecular reaction is proportional to $e^{-E_a/(R \cdot T)}$. For example, if a given enzyme reduces the Ea of a unimolecular reaction from 126 to 63 kJ · mol-1, the rate constant at 310 °K (37 °C) will rise by e^{-63000/(8.31 · 310)}/e^{-126000/(8.31 ·} ³¹⁰⁾, i.e., by a factor of 4 10¹⁰. The enzyme would therefore reduce the time required to metabolize 50% of the starting materials (t1/2) from, say, 10 years to 7 msec! The forward rate of a reaction $(mol \cdot L^{-1} \cdot s^{-1})$ is related to the product of the rate constant (s-1) and the starting substrate concentration (mol \cdot L⁻¹).

The second law of thermodynamics also implies that a continuous loss of free energy occurs as the total disorder or entropy (S) of a closed system increases. A living organism represents an open system which, by definition, can absorb energy-rich nutrients and discharge end products of metabolism. While the entropy of a closed system (organism + environment) increases in the process, an open system (organism) can either maintain its entropy level or reduce it using free enthalpy. This occurs, for example, when ion gradients or hydraulic pressure differences are created within the body. A closed system therefore has a maximum entropy, is in a true state of chemical equilibrium, and can perform work only once. An open system such as the body can continuously perform work while producing only a minimum of entropy. A true state of equilibrium is achieved in only a very few processes within the body, e.g., in the reaction $CO_2 + H_2O - HCO_3^- + H^+$. In most cases (e.g. metabolic pathways, ion gradients), only a *steady state* is reached. Such metabolic pathways are usually *irreversible* due, for example, to excretion of the end products. The thought of reversing the "reaction" germ cell \rightarrow adult illustrates just how impossible this is.

At **steady state**, the *rate* of a reaction is more important than its equilibrium. The regulation of body functions is achieved by controlling reaction rates. Some reactions are so slow that it is impossible to achieve a sufficient reaction rate with enzymes or by reducing the concentration of the reaction products. These are therefore endergonic reactions that require the input of outside energy. This can involve "activation" of the educt by attachment of a high-energy phosphate group to raise the Pe.

ATP (adenosine triphosphate) is the universal carrier and transformer of free enthalpy within the body. ATP is a nucleotide that derives its chemical energy from energy-rich nutrients (\rightarrow C). Most ATP is produced by **oxi**dation of energy-rich biological molecules such as glucose. In this case, oxidation means the removal of electrons from an electron-rich (reduced) donor which, in this case, is a carbohydrate. CO₂ and H₂O are the end products of the reaction. In the body, oxidation (or electron transfer) occurs in several stages, and a portion of the liberated energy can be simultaneously used for ATP synthesis. This is therefore a **coupled reaction** (\rightarrow **C** and p. 17 B). The standard free enthalpy ΔG^0 of ATP hydrolysis,

ATP ADP + P_i [1.31] is $-30.5 \text{ kJ} \cdot \text{mol}^{-1}$. According to Eq. 1.27, the ΔG of reaction 1.31 should increase when the ratio ([ADP] \cdot [P_i)]/[ATP] falls below the equilibrium constant K_{eq} of ATP hydrolysis. The fact that a high cellular ATP concentration does indeed yield a ΔG of approximately -46 to $-54 \text{ kJ} \cdot \text{mol}^{-1}$ shows that this also applies in practice.

Some substances have a much higher ΔG^0 of hydrolysis than ATP, e.g., *creatine phosphate* ($-43 \text{ kJ} \cdot \text{mol}^{-1}$). These compounds react with ADP and P_i to form ATP. On the other hand, the energy of ATP can be used to synthesize other compounds such as UTP, GTP and glucose-6-phosphate. The energy content of these sub-

stances is lower than that of ATP, but still relatively high.

The free energy liberated upon hydrolysis of ATP is used to drive hundreds of reactions within the body, including the active transmembrane transport of various substances, protein synthesis, and muscle contraction. According to the laws of thermodynamics, the expenditure of energy in all of these reactions leads to increased order in living cells and, thus, in the organism as a whole. Life is therefore characterized by the continuous reduction of entropy associated with a corresponding increase in entropy in the immediate environment and, ultimately, in the universe.

42

Nerve and Muscle. Physical Work

Neuron Structure and Function

An excitable cell reacts to stimuli by altering its membrane characteristics (\rightarrow p. 32). There are two types of excitable cells: nerve cells, which transmit and modify impulses within the nervous system, and muscle cells, which contract either in response to nerve stimuli or autonomously (\rightarrow p. 59).

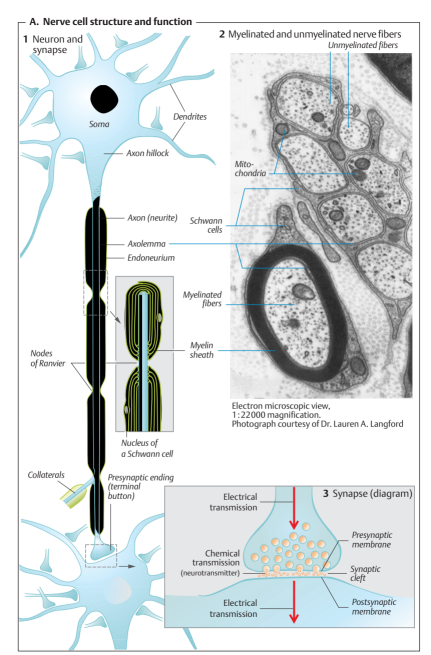
The human nervous system consists of more than 1010 nerve cells or neurons. The neuron is the structural and functional unit of the nervous system. A typical neuron (motor neuron, \rightarrow A1) consists of the soma or cell body and two types of processes: the axon and dendrites. Apart from the usual intracellular organelles (\rightarrow p. 8 ff.), such as a nucleus and mitochondria (\rightarrow A2), the neuron contains neurofibrils and neurotubules. The neuron receives afferent signals (excitatory and inhibitory) from a few to sometimes several thousands of other neurons via its dendrites (usually arborescent) and sums the signals along the cell membrane of the soma (summation). The axon arises from the axon hillock of the soma and is responsible for the transmission of efferent neural signals to nearby or distant effectors (muscle and glandular cells) and adjacent neurons. Axons often have branches (collaterals) that further divide and terminate in swellings called synaptic knobs or terminal buttons. If the summed value of potentials at the axon hillock exceeds a certain threshold, an action poten**tial** (\rightarrow p. 46) is generated and sent down the axon, where it reaches the next synapse via the **terminal buttons** (\rightarrow **A1,3**) described below.

Vesicles containing materials such as proteins, lipids, sugars, and transmitter substances are conveyed from the Golgi complex of the soma (\rightarrow p. 13 F) to the terminal buttons and the tips of the dendrites by rapid axonal transport (40 cm/day). This type of anterograde transport along the neurotubules is promoted by kinesin, a myosin-like protein, and the energy required for it is supplied by ATP (→p. 16). Endogenous and exogenous substances such as nerve growth factor (NGF), herpes virus, poliomyelitis virus, and tetanus toxin are conveyed by retrograde transport from the peripheral regions to the soma at a

rate of ca. 25 cm/day. Slow axon transport (ca. 1 mm/day) plays a role in the regeneration of severed neurites.

Along the axon, the plasma membrane of the soma continues as the axolemma (\rightarrow A1,2). The axolemma is surrounded by oligodendrocvtes (\rightarrow p. 338) in the central nervous system (CNS), and by **Schwann cells** in the peripheral nervous system (\rightarrow A1,2). A nerve fiber consists of an axon plus its sheath. In some neurons. Schwann cells form multiple concentric double phospholipid layers around an axon, comprising the myelin sheath $(\rightarrow A1,2)$ that insulates the axon from ion currents. The sheath is interrupted every 1.5 mm or so at the **nodes** of Ranvier (→ A1). The conduction velocity of myelinated nerve fibers is much higher than that of unmyelinated nerve fibers and increases with the diameter of the nerve fiber $(\rightarrow p.49 C)$.

A **synapse** $(\rightarrow A3)$ is the site where the axon of a neuron communicates with effectors or other neurons (see also p. 50 ff.). With very few exceptions, synaptic transmissions in mammals are mediated by chemicals, not by electrical signals. In response to an electrical signal in the axon, vesicles (\rightarrow p. 1.6) on the **presynaptic** membrane release transmitter substances (neurotransmitters) by exocytosis ($\rightarrow p.30$). The transmitter diffuses across the synaptic cleft (10-40 nm) to the postsynaptic membrane, where it binds to receptors effecting new electrical changes (\rightarrow A3). Depending on the type of neurotransmitter and postsynaptic receptor involved, the transmitter will either have an excitatory effect (e.g., acetylcholine in skeletal muscle) or inhibitory effect (e.g., glycine in the CNS) on the postsynaptic membrane. Since the postsynaptic membrane normally does not release neurotransmitters (with only few exceptions), nerve impulses can pass the synapse in one direction only. The synapse therefore acts like a valve that ensures the orderly transmission of signals. Synapses are also the sites at which neuronal signal transmissions can be modified by other (excitatory or inhibitory) neurons.



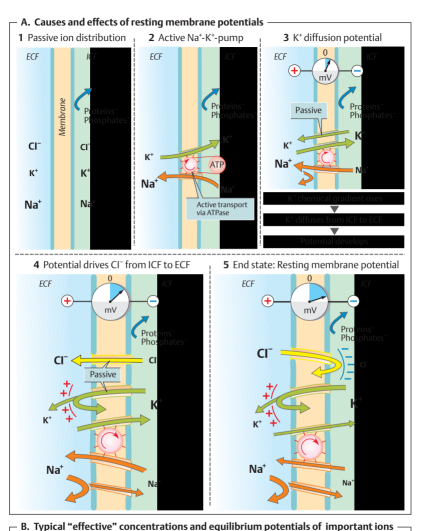
An electrical potential difference, or membrane potential (E_m), can be recorded across the plasma membrane of living cells. The potential of unstimulated muscle and nerve cells, or resting potential, amounts to -50 to - 100 mV (cell interior is negative). A resting potential is caused by a slightly unbalanced distribution of ions between the intracellular fluid (ICF) and extracellular fluid (ECF) ($\rightarrow B$). The following factors are involved in establishing the membrane potential (see also p. 32 ff.). Maintenance of an unequal distribution of ions: The Na⁺-K⁺-ATPase (\rightarrow p. 26) continuously "pumps" Na+ out of the cell and K+ into it (→ A2). As a result, the intracellular K⁺ concentration is around 35 times higher and the intracellular Na⁺ concentration is roughly 20 times lower than the extracellular concentration $(\rightarrow B)$. As in any active transport, this process requires energy, which is supplied by ATP. Lack of energy or inhibition of the Na+-K+-ATPase results in flattening of the ion gradient and breakdown of the membrane potential.

Because anionic proteins and phosphates present in high concentrations in the cytosol are virtually unable to leave the cell, purely passive mechanisms (Gibbs-Donnan distribution) could, to a slight extent, contribute to the unequal distribution of diffusable ions (\rightarrow A1). For reasons of electroneutrality, [K*+Na*]_{ECF} = md [Cl*]_{ECF} < [Cl*]_{ECF}. However, this has practically no effect on the development of resting potentials.

- ♦ Low resting Na⁺ and Ca²⁺ conductance, g_{Na} , g_{Ca} : The membrane of a resting cell is only very slightly permeable to Na⁺ and Ca²⁺, and the resting g_{Na} comprises only a small percentage of the total conductance (→ p. 32 ff.). Hence, the Na⁺ concentration difference (→ A3–A5) cannot be eliminated by immediate passive diffusion of Na⁺ back into the cell.
- ♦ **High K**+**conductance**, **g**_K: It is relatively easy for K⁺ ions to diffuse across the cell membrane ($g_K \approx 90\%$ of total conductance; → p. 32ff.). Because of the steep concentration gradient (→ point 1), K⁺ ions diffuse from the ICF to the ECF (→ **A3**). Because of their positive charge, the diffusion of even small amounts of K⁺ ions leads to an electrical potential (*diffusion potential*) across the membrane. This (inside negative for the same propertial) across the membrane.

- tive) diffusion potential drives K^+ back into the cell and rises until large enough to almost completely compensate for the K^+ concentration gradient driving the K^+ -ions out of the cell $(\rightarrow$ **A4**). As a result, the membrane potential, E_{m_r} is approximately equal to the K^+ equilibrium potential E_K $(\rightarrow$ p. 32).
- ◆ Cl⁻ **distribution:** Since the cell membrane is also conductive to Cl⁻ (g_{Cl} greater in muscle cells than in nerve cells), the membrane potential (electrical driving "force") expels Cl⁻ ions from the cell (→ A4) until the Cl⁻ concentration gradient (chemical driving "force") drives them back into the cell at the same rate. The intracellular Cl⁻ concentration, [Cl⁻]_i, then continues to rise until the Cl⁻ equilibrium potential equals E_m (→ A5). [Cl⁻]_i can be calculated using the *Nernst equation* (→ p. 32, Eq. 1.18). Such a "passive" distribution of Cl⁻ between the intra- and extracellular spaces exists only as long as there is no active Cl⁻ uptake into the cell (→ p. 34).
- ♦ Why is E_m less negative than E_K? Although the conductances of Na+ and Ca2+ are very low in resting cells, a few Na+ and Ca2+ ions constantly enter the cell (\rightarrow A4, 5). This occurs because the equilibrium potential for both types of ions extends far into the positive range, resulting in a high outside-to-inside electrical and chemical driving "force" for these ions $(\rightarrow B; \rightarrow p. 32f.)$. This cation influx depolarizes the cell, thereby driving K⁺ ions out of the cell (1 K⁺ for each positive charge that enters). If Na+-K+-ATPase did not restore these gradients continuously (Ca2+ indirectly via the 3 Na+/Ca2+ exchanger; \rightarrow p. 36), the intracellular Na⁺ and Ca2+ concentrations would increase continuously, whereas [K+]i would decrease, and EK and Em would become less negative.

All living cells have a (resting) membrane potential, but only excitable cells such as nerve and muscle cells are able to greatly change the ion conductance of their membrane in response to a stimulus, as in an *action potential* $(\rightarrow p.46)$.



in skeletal muscle (at 37°C) "Effective" concentration (mmol/kg H₂O) Equilibrium potential Interstice (ECF) Cell (ICF) K⁺ 4.5 160 - 95 mV Na⁺ 144 7 + 80 mV Ca²⁺ 0.0001-0.00001 +125 to +310 mV 1.3 (After Conway) - 24 mV H⁺ 4.10^{-5} (pH 7.4) 10⁻⁴ (pH 7.0) - 80 mV CI-114 - 27 mV HCO₃ 28

Action Potential

An action potential is a signal passed on through an axon or along a muscle fiber that influences other neurons or induces muscle contraction. Excitation of a neuron occurs if the membrane potential, Em, on the axon hillock of a motor neuron, for example (\rightarrow p. 42), or on the motor end-plate of a muscle fiber changes from its resting value (\rightarrow p. 44) to a less negative value (slow depolarization, $\rightarrow A1$). This depolarization may be caused by neurotransmitter-induced opening of postsynaptic cation channels (\rightarrow p. 50) or by the (electrotonic) transmission of stimuli from the surroundings (\rightarrow p. 48). If the E_m of a stimulated cell comes close to a critical voltage or threshold potential (→ A1), "rapid" voltage-gated Na⁺ channels are activated (\rightarrow B4 and B1 \Rightarrow B2). This results in increased Na⁺ conductance, g_{Na} $(\rightarrow p.32)$, and the entry of Na⁺ into the cell $(\rightarrow A2)$. If the threshold potential is not reached, this process remains a local (subthreshold) response.

Once the threshold potential is reached, the cell responds with a fast all-or-none depolarization called an **action potential**, AP $(\rightarrow A1)$. The AP follows a pattern typical of the specific cell type, irregardless of the magnitude of the stimulus that generated it. Large numbers of Na+ channels are activated, and the influxing Na+ accelerates depolarization which, in turn, increases g_{Na} and so on (positive feedback). As a result, the E_m rapidly collapses (0.1 ms in nerve cells: fast depolarization phase or upsweep) and temporarily reaches positive levels (overshooting, +20 to +30 mV). The g_{Na} drops before overshooting occurs $(\rightarrow A2)$ because the Na+ channels are inactivated within 0.1 ms $(\rightarrow B2 \Rightarrow B3)$. The potential therefore reverses, and restoration of the resting potential, the repolarization phase of the action potential, begins. Depolarization increases (relatively slowly) the open-probability of voltage-gated K⁺ channels. This increases the potassium conductance, g_K, thereby accelerating repolarization.

In many cases, potassium conductance, g_K is still increased after the original resting potential has been restored (\rightarrow **A2**), and E_m temporarily approaches E_K (\rightarrow pp. 44 and 32 ff.),

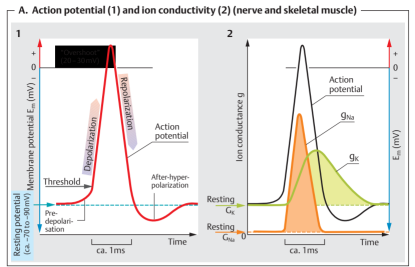
resulting in a hyperpolarizing **afterpotential** (\rightarrow **A1**). Increased Na⁺-K⁺-ATPase pumping rates (electrogenic; \rightarrow p. 28) can contribute to this afterpotential.

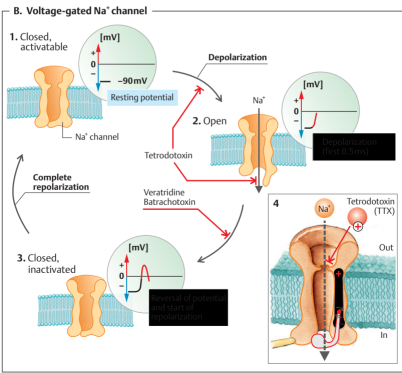
Very long trains of action potentials can be generated (up to 1000/s in some nerves) since the quantity of ions penetrating the membrane is very small (only ca. $1/100\,000$ th the number of intracellular ions). Moreover, the Na*-K*-ATPase (\rightarrow p. 26) ensures the continuous restoration of original ion concentrations (\rightarrow p. 46).

During an action potential, the cell remains unresponsive to further stimuli; this is called the **refractory period**. In the *absolute refractory period*, no other action potential can be triggered, even by extremely strong stimuli, since Na^+ channels in depolarized membranes cannot be activated (\rightarrow **B3**). This is followed by a *relative refractory period* during which only action potentials of smaller amplitudes and rates of rise can be generated, even by strong stimuli. The refractory period ends once the membrane potential returns to its resting value (\rightarrow e.g. p. 59 A).

The extent to which Na+ channels can be activated and, thus, the strength of the Na+ current, I_{Na}, depends on the pre-excitatory resting potential, not the duration of depolarization. The activation of the Na+ channels reaches a maximum at resting potentials of ca. - 100 mV and is around 40% lower at -60 mV. In mammals. Na+ channels can no longer be activated at potentials of -50 mV and less negative values (\rightarrow **B3**). This is the reason for the absolute and relative refractory periods (see above) and the non-excitability of cells after the administration of continuously depolarizing substances such as suxamethonium (\rightarrow p. 56). An increased extracellular Ca2+ concentration makes it more difficult to stimulate the cell because the threshold potential becomes less negative. On the other hand, excitability increases (lower threshold) in hypocalcemic states, as in muscle spasms in tetany $(\to p.290)$.

The special features of action potentials in cardiac and smooth muscle fibers are described on pages 192, 70 and 59 A.





Propagation of Action Potentials in Nerve Fiber

Electrical current flows through a *cable* when voltage is applied to it. The metal wire inside the cable is well insulated and has very low-level resistance, reducing current loss to a minimum. As a result, it can conduct electricity over long distances. *Nerve fibers*, especially unmyelinated ones (\rightarrow p. 42), have a much greater internal longitudinal resistance (R_i) and are not well insulated from their surroundings. Therefore, the cable-like, **electrotonic transmission** of neural impulses dwindles very rapidly, so the conducted impulses must be continuously "refreshed" by generating new **action potentials** (\rightarrow p. 46).

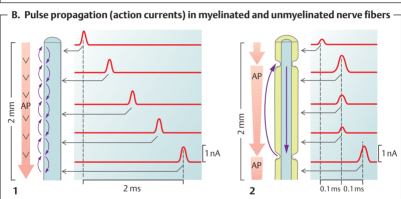
Propagation of action potentials: The start of an action potential is accompanied by a brief influx of Na^+ into the nerve fiber (\rightarrow **A1a**). The cell membrane that previously was inside negative now becomes positive (+20 to +30 mV), thus creating a longitudinal potential difference with respect to the adjacent, still unstimulated nerve segments (internal -70 to -90 mV; \rightarrow p. 44). This is followed by a passive *electrotonic withdrawal of charge* from the adjacent segment of the nerve fiber, causing its *depolarization*. If it exceeds threshold, another action potential is created in the adjacent segment and the action potential in the previous segment dissipates (\rightarrow **A1b**).

Because the membrane acts as a *capacitor*. the withdrawal of charge represents a capacitating (depolarizing) flow of charge that becomes smaller and rises less steeply as the spatial distance increases. Because of the relatively high R_i of nerve fiber, the outward loops of current cross the membrane relatively close to the site of excitation, and the longitudinal current decreases as it proceeds towards the periphery. At the same time, depolarization increases the driving force (= $E_m - E_K$; \rightarrow p. 32) for K⁺ outflow. K⁺ fluxing out of the cell therefore accelerates repolarization. Hence, distal action potentials are restricted to distances from which the capacitative current suffices to depolarize the membrane quickly and strongly enough. Otherwise, the Na+ channels will be deactivated before the threshold potential is reached (\rightarrow p. 46).

Action potentials normally run forward (orthodromic) because each segment of nerve fiber becomes refractory when an action potential passes (\rightarrow **A1b** and p. 46). If, however, the impulses are conducted backwards (antidomic) due, for example, to electrical stimulation of nerve fibers from an external source (\rightarrow p. 50), they will terminate at the next synapse (valve-like function, \rightarrow p. 42).

Although the continuous generation of action potentials in the immediately adjacent fiber segment guarantees a refreshed signal. this process is rather time-consuming (\rightarrow **B1**). The **conduction velocity**, θ , in unmyelinated (type C) nerve fibers $(\rightarrow \mathbf{C})$ is only around 1 m/s. Myelinated (types A and B) nerve fibers $(\rightarrow \mathbf{C})$ conduct much faster (up to 80 m/s = 180 mph in humans). In the internode regions, a myelin sheath (\rightarrow p.42) insulates the nerve fibers from the surroundings; thus, longitudinal currents strong enough to generate action potentials can travel further down the axon (ca. 1.5 mm) (\rightarrow **A2**). This results in more rapid conduction because the action potentials are generated only at the unmyelinated nodes of Ranvier, where there is a high density of Na+ channels. This results in rapid, jump-like passage of the action potential from node to node (saltatory propagation). The saltatory length is limited since the longitudinal current (1 to 2 nA) grows weaker with increasing distance $(\rightarrow B2)$. Before it drops below the threshold level, the signal must therefore be refreshed by a new action potential, with a time loss of 0.1 ms.

Since the internal resistance, R_i , of the nerve fiber limits the spread of depolarization, as described above, the **axon diameter** (2r) also affects the conduction velocity, $\theta (\rightarrow C)$. R_i is proportional to the cross-sectional area of the nerve fiber (πr^2) , i.e., $R_i \sim 1/r^2$. Thick fibers therefore require fewer new APs per unit of length, which is beneficial for θ . Increases in fiber diameter are accompanied by an increase in both fiber circumference $(2\pi r)$ and membrane capacity, $K(K \sim r)$. Although θ decreases, the beneficial effect of the smaller R_i predominates because of the quadratic relationship.



— C. Classif	ication of nerve fibers (in humans) —	n according to fiber type vd and Hunt types I – IV) cle efferent, afferents in muscle and tendon organs (Ib) Diameter (μm) cle efferent, afferents in muscle and tendon organs (Ib)								
C. Classification of fictive fibers (in figure 1)										
Fiber type	Function according to fiber type (Lloyd and Hunt types I – IV)									
Αα	Skeletal muscle efferent, afferents in muscle spindles (lb) and tendon organs (lb)	11–16	60 – 80							
Αβ	Mechanoafferents of skin (II)	6 –11	30 – 60							
Αγ	Muscle spindle efferents									
Αδ	Skin afferents (temperature and "fast" pain) (III)	1-6	2 – 30							
В	Sympathetic preganglionic; visceral afferents	3	3 – 15							
С	Skin afferents ("slow" pain); sympathetic postganglionic afferents (IV)	0.5–1.5 (unmyelinated)	0.25 – 1.5							
			(After Erlanger and Gasser)							

Artificial Stimulation of Nerve Cells

When an electrical stimulus is applied to a nerve cell from an external source, current flows from the positive stimulating electrode (anode) into the neuron, and exits at the negative electrode (cathode). The nerve fiber below the cathode is depolarized and an action potential is generated there if the threshold potential is reached.

The **conduction velocity** of a nerve can be measured by placing two electrodes on the skin along the course of the nerve at a known distance from each other, then stimulating the nerve (containing multiple neurons) and recording the time it takes the summated action potential to travel the known distance. The conduction velocity in humans is normally 40 to 70 m · s⁻¹. Values below 40 m · s⁻¹ are considered to be pathological.

Accidental electrification. Exposure of the body to high-voltage electricity, especially low-frequency alternating current (e.g., in an electrical outlet) and low contact resistance (bare feet, bathtub accidents), primarily affects the conduction of impulses in the heart and can cause ventricular fibrillation (\rightarrow p. 200).

Direct current usually acts as a stimulus only when switched on or off: High-frequency alternating current (> 15 kHz), on the other hand, cannot cause depolarization but heats the body tissues. **Diathermy** works on this principle.

Synaptic Transmission

Synapses connect nerve cells to other nerve cells (also applies for certain muscle cells) as well as to sensory and effector cells (muscle and glandular cells).

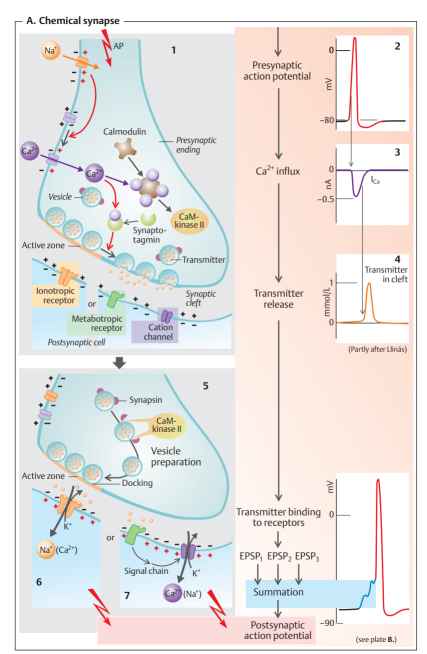
Electrical synapses are direct, ion-conducting cell–cell junctions through channels (*connexons*) in the region of *gap junctions* (→ p. 16 f.). They are responsible for the conduction of impulses between neighboring smooth or cardiac muscle fibers (and sometimes between neurons in the retina and in the CNS) and ensure also communication between neighboring epithelial or glial cells.

Chemical synapses utilize (neuro)transmitters for the transmission of information and

provide not only simple 1:1 connections, but also serve as switching elements for the nervous system. They can facilitate or inhibit the neuronal transmission of information or process them with other neuronal input. At the chemical synapse, the arrival of an action potential (AP) in the axon (\rightarrow **A1,2** and p.48) triggers the release of the transmitter from the presynaptic axon terminals. The transmitter then diffuses across the narrow synaptic cleft (ca. 30 nm) to bind postsynaptically to receptors in the subsynaptic membrane of a neuron or of a glandular or muscle cell. Depending on the type of transmitter and receptor involved, the effect on the postsynaptic membrane may either be excitatory or inhibitory, as is described below.

Transmitters are released by regulated exocytosis of so-called synaptic vesicles $(\rightarrow A1)$. Each vesicle contains a certain quantum of neurotransmitters. In the case of the motor end-plate (\rightarrow p. 56), around 7000 molecules of acetylcholine (ACh) are released. Some of the vesicles are already docked on the membrane (active zone), ready to exocytose their contents. An incoming action potential functions as the signal for transmitter release (\rightarrow A1,2). The higher the action potential frequency in the axon the more vesicles release their contents. An action potential increases the open probability of voltage-gated Ca2+ channels in the presynaptic membrane (sometimes oscillating), thereby leading to an increase in the cytosolic Ca^{2+} concentration, $[Ca^{2+}]_i (\rightarrow A1, 3)$ and p. 36). Extracellular Mg2+ inhibits this process. Ca^{2+} binds to synaptotagmin ($\rightarrow A1$), which triggers the interaction of syntaxin and SNAP-25 on the presynaptic membrane with synaptobrevin on the vesicle membrane, thereby triggering exocytosis of already docked vesicles (approximately 100 per AP) $(\rightarrow A1, 4)$. On the other hand, Ca^{2+} activates calcium-calmodulin-dependent protein kinase-II (CaM-kinase-II: \rightarrow A5, and p. 36), which activates the enzyme synapsin at the presynaptic terminal. As a result, vesicles dock anew on the active zone.

Synaptic facilitation (= potentiation). If an action potential should arrive at the presynaptic terminal immediately after another AP (AP frequency > approx. 30 Hz), the cytosolic Ca²⁺



concentration will not yet drop to the resting value, and residual Ca^{2+} will accumulate. As a result, the more recent rise in $[Ca^{2+}]_i$ builds on the former one. $[Ca^{2+}]_i$ rises to a higher level after the second stimulus than after the first, and also releases more transmitters. Hence, the first stimulus *facilitates* the response to the second stimulus. Muscle strength increases at high stimulus frequencies for similar reasons $(\rightarrow p, 67 \text{ A})$.

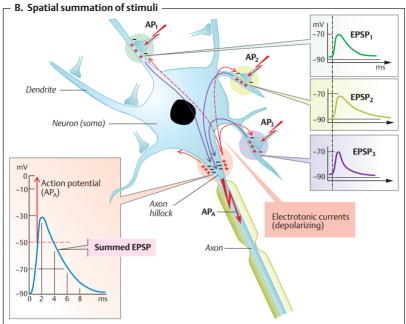
Among the many substances that act as excitatory transmitters are acetylcholine (ACh) and glutamate (Glu). They are often released together with co-transmitters which modulate the transmission of a stimulus (e.g., ACh together with substance P, VIP or galanin; Glu with substance P or enkephalin). If the transmitter's receptor is an ion channel itself (ionotropic receptor or ligand-gated ion channel; \rightarrow A6 and F), e.g., at the N-cholinergic synapse (\rightarrow p. 82), the channels open more often and allow a larger number of cations to enter (Na+, sometimes Ca2+) and leave the cell (K+). Other, so-called metabotropic receptors influence the channel via G proteins that control channels themselves or by means of "second messengers" (\rightarrow A7 and F). Because of the high electrochemical Na⁺ gradient (\rightarrow p.32), the number of incoming Na+ ions is much larger than the number of exiting K⁺ ions, Ca²⁺ can also enter the cell, e.g., at the glutamate-NMDA receptor $(\rightarrow F)$. The net influx of cations leads to depolarization: excitatory postsynaptic **potential (EPSP)** (maximum of ca. 20 mV; \rightarrow **B**). The EPSP begins approx. 0.5 ms after the arrival of an action potential at the presynaptic terminal. This synaptic delay (latency) is caused by the relatively slow release and diffusion of the transmitter.

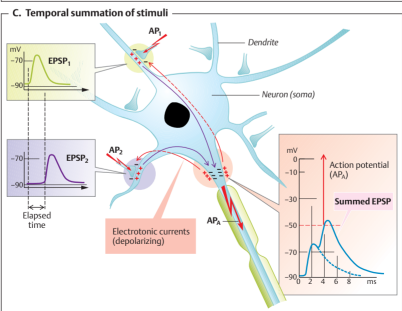
A single EPSP normally is not able to generate a postsynaptic (axonal) action potential (AP_A), but requires the triggering of a large number of local depolarizations in the dendrites. Their depolarizations are transmitted electrotonically across the soma (\rightarrow p. 48) and summed on the axon hillock (**spatial summation**; \rightarrow **B**). Should the individual stimuli arrived each other), the prior depolarization will not have dissipated before the next one arrives, and summation will make it easier to reach

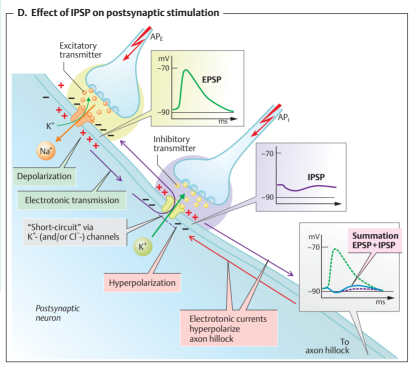
threshold. This type of **temporal summation** therefore increases the excitability of the post-synaptic neuron $(\rightarrow \mathbb{C})$.

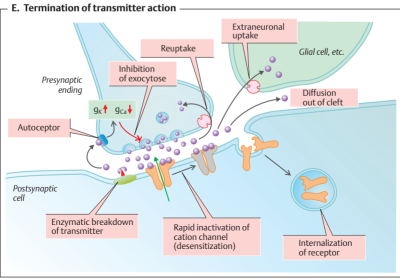
Inhibitory transmitters include substances as glycine, GABA (y-aminobutyric acid), and acetylcholine (at M2 and M3 receptors: \rightarrow p. 82). They increase the conductance, g. of the subsynaptic membrane only to K+ (e.g., the metabotropic GABA_R receptor.) or Cl⁻ (e.g., the ionotropic glycine and GABA $_{\Lambda}$ receptors: \rightarrow **F**). The membrane usually becomes hyperpolarized in the process (ca. 4 mV max.). Increases in g_K occur when E_m approaches E_K $(\rightarrow p.44)$. However, the main effect of this inhibitory postsynaptic potential IPSP $(\rightarrow \mathbf{D})$ is not hyperpolarization-which works counter to EPSP-related depolarization (the IPSP is sometimes even slightly depolarizing). Instead, the IPSP-related increase in membrane conductance short circuits the electrotonic currents of the EPSP (high g_K or g_{Cl} levels). Since both E_K and E_{CI} are close to the resting potential $(\rightarrow p.44)$, stabilization occurs, that is, the EPSP is cancelled out by the high K+ and Cl- shortcircuit currents. As a result, EPSP-related depolarization is reduced and stimulation of postsynaptic neurons is inhibited ($\rightarrow \mathbf{D}$).

Termination of synaptic transmission $(\rightarrow E)$ can occur due to inactivation of the cation channels due to a conformational change in the channel similar to the one that occurs during an action potential (\rightarrow p. 46). This very rapid process called desensitization also functions in the presence of a transmitter. Other terminating pathways include the rapid enzymatic decay of the transmitter (e.g., acetylcholine) while still in the synaptic cleft, the re-uptake of the transmitter (e.g., noradrenaline) into the presynaptic terminal or uptake into extraneuronal cells (e.g., in glial cells of the CNS), endocytotic internalization of the receptor (\rightarrow p. 28), and binding of the transmitter to a receptor on the presynaptic membrane (autoceptor). In the latter case, a rise in g_K and a drop in g_{Ca} can occur, thus inhibiting transmitter release, e.g., of GABA via GABAB receptors or of noradrenaline via α_2 -adrenoceptors ($\rightarrow \mathbf{F}$ and p. 86).









Transmitter	Receptor	Recep	tor	Effect						
	subtypes	types		lon	conc	lucta	nce	Second n	Second messenger	
				Na ⁺	K ⁺	Ca ²⁺	Cl-	cAMP	IP ₃ /DAG	
Acetylcholine	Nicotinic Muscarinic: M1, M2, M3			†	† †	1		<u>¥</u>		
ADH (= vasopressin)	V1 V2							^	^	
CCK (= cholecystokinin)	CCK_{A-B}								^	
Dopamine	D1, D5 D2				1	ţ		<u>↓</u>		
GABA (= γ-aminobutyric acid)	GABA _A , GABA _C GABA _B				1	1	1	<u>¥</u>		
Glutamate (aspartate)	AMPA Kainat NMDA m-GLU			† †	† †	t		业	^	
Glycine	_	•					1			
Histamine	H ₁ H ₂							^	^	
Neurotensin	-							<u>¥</u>	^	
Norepinephrine, epinephrine	$\begin{array}{l}\alpha_{1\text{(A-D)}}\\\alpha_{2\text{(A-C)}}\\\beta_{1-3}\end{array}$				†	+		₩ ^	^	
Neuropeptide Y (NPY)	Y1-2				1	1		<u>¥</u>		
Opioid peptides	μ, δ, κ				1	1		<u>¥</u>		
Oxytocin	-								^	
Purines	P_{1} : A_{1} A_{2a} P_{2X} P_{2Y}			†	†	1		<u>▼</u>	^	
Serotonin (5-hydroxytryptamine)	5-HT ₁ 5-HT ₂ 5-HT ₃ 5-HT ₄₋₇	•		†	†			<u>↓</u>	^	
Somatostatin (= SIH)	SRIF				1	+		<u>¥</u>		
Tachykinin	NK1-3								^	
Amino acids Catecholamines					•)	Inhibits o	or promote	
Peptides Others				botropic receptor otein-mediated t)				↑(CAMP) ATP	PIP ₂ IP	

(Modified from F. E. Bloom)

Motor End-plate

The transmission of stimuli from a motor axon to a skeletal muscle fiber occurs at the motor end-plate, MEP (\rightarrow A), a type of chemical synapse (\rightarrow p. 50 ff.). The transmitter involved is acetylcholine (ACh, \rightarrow cf. p. 82), which binds to the N(nicotinergic)-cholinoceptors of the subsynaptic muscle membrane (\rightarrow A3). N-cholinoceptors are ionotropic, that is, they also function as *ion channels* (\rightarrow A4). The N-cholinoceptor of the MEP (type N_M) has 5 subunits (2α , 1β , 1γ , 1δ), each of which contains 4 membranespanning α -helices (\rightarrow p. 14).

The channel opens briefly (\rightarrow **B1**) (for approx. 1 ms) when an ACh molecule binds to the two α -subunits of an N-cholinoceptor (\rightarrow **A4**). Unlike voltage-gated Na⁺-channels, the openprobability p_o of the N_M-cholinoceptor is not increased by depolarization, but is determined by the *ACh concentration in the synaptic cleft* (\rightarrow p. 50 ff.).

The channel is specific to cations such as Na⁺, K⁺, and Ca²⁺. Opening of the channel at a resting potential of ca. -90 mV leads mainly to an influx of Na⁺ ions (and a much lower outflow of K^+ ; \rightarrow pp. 32 ff. and 44). Depolarization of the subsynaptic membrane therefore occurs: endplate potential (EPP). Single-channel currents of 2.7 pA (\rightarrow B1) are summated to yield a miniature end-plate current of a few nA when spontaneous exocytosis occurs and a vesicle releases a quantum of ACh activating thousands of N_M -cholinoceptors (\rightarrow **B2**). Still, this is not enough for generation of a postsynaptic action potential unless an action potential transmitted by the motor neuron triggers exocytosis of around a hundred vesicles. This opens around 200,000 channels at the same time, yielding a neurally induced end-plate current (I_{EP}) of ca. 400 nA (\rightarrow **B3). End-plate current, I**_{EP}, is therefore dependent on:

• the number of open channels, which is equal to the total number of channels (n) times the open-probability (p_o) , where p_o is determined by the concentration of ACh in the synaptic cleft (up to 1 mmol/L);

• the single-channel conductance γ (ca. 30 pS);

♦ and, to a slight extent, the membrane potential, E_m, since the electrical driving

"force" (= E_m – $E_{Na,K}$; \rightarrow p. 32 ff.) becomes smaller when E_m is less negative.

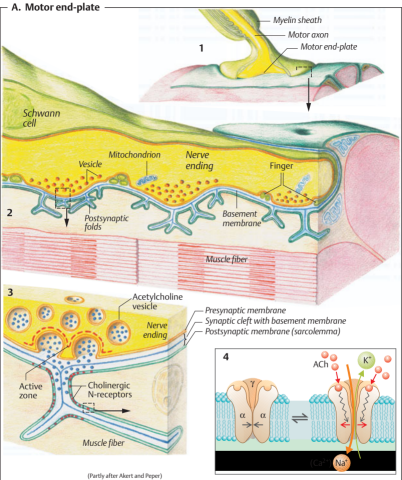
 $E_{Na,K}$ is the common equilibrium potential for Na^+ and K^+ and amounts to approx. 0 mV. It is also called the **reversal potential** because the direction of I_{EP} (= I_{Na} + I_K), which enters the cell when E_m is negative (Na^+ influx > K^+ outflow), reverses when E_m is positive (K^+ outflow > Na^+ influx). As a result,

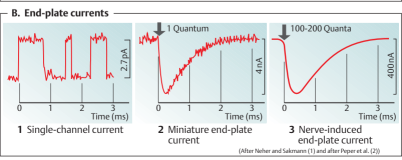
$$I_{EP} = n \cdot p_o \cdot \gamma \cdot (E_m - E_{Na, K}) [A]$$
 [2.1]

Because **neurally induced EPPs** in skeletal muscle are much larger (depolarization by ca. 70 mV) than neuronal EPSPs (only a few mV; \rightarrow p. 50 ff.), single motor axon action potentials are above threshold. The EPP is transmitted electrotonically to the adjacent sarcolemma, where muscle action potentials are generated by means of voltage-gated Na $^+$ channels, resulting in muscle contraction.

Termination of synaptic transmission in MEPs occurs (1) by rapid degradation of ACh in the synaptic cleft by *acetylcholinesterase* localized at the subsynaptic basal membrane, and (2) by diffusion of ACh out of the synaptic cleft $(\rightarrow p.82)$.

A motor end-plate can be blocked by certain poisons and drugs, resulting in muscular weakness and, in some cases, paralysis. Botulinum neurotoxin, for example, inhibits the discharge of neurotransmitters from the vesicles, and α -bungarotoxin in cobra venom blocks the opening of ion channels. Curare-like substances such as (+)-tubocurarine are used as muscle relaxants in surgical operations. They displace ACh from its binding site (competitive inhibition) but do not have a depolarizing effect of their own. Their inhibitory effect can be reversed by cholinesterase inhibitors such as neostigmine (decurarinization). These agents increase the concentration of ACh in the synaptic cleft, thereby displacing curare. Entry of anticholinesterase agents into intact synapses leads to an increase in the ACh concentration and, thus, to paralysis due to permanent depolarization. ACh-like substances such as suxamethonium have a similar depolarizing effect, but decay more slowly than ACh. In this case, paralysis occurs because permanent depolarization also permanently inactivates Na+ channels near the motor end-plate on the sarcolemma (\rightarrow p. 46).





Motility and Muscle Types

Active motility (ability to move) is due to either the **interaction of** energy-consuming **motor proteins** (fueled by ATPase) such as myosin, kinesin and dynein with other proteins such as actin or the **polymerization** and depolymerization of actin and tubulin. Cell division (cytokinesis), cell migration (\rightarrow p. 30), intracellular vesicular transport and cytosis (\rightarrow p. 12f.), sperm motility (\rightarrow p. 306f.), axonal transport (\rightarrow p. 42), electromotility of hair cells (\rightarrow p. 366), and ciliary motility (\rightarrow p. 110) are examples of cell and organelle motility.

The muscles consist of cells (fibers) that contract when stimulated. **Skeletal muscle** is responsible for locomotion, positional change, and the convection of respiratory gases. **Cardiac muscle** (\rightarrow p. 190 ff.) is responsible for pumping the blood, and **smooth muscle** (\rightarrow p. 70) serves as the motor of internal organs and blood vessels. The different muscle types are distinguished by several functional characteristics (\rightarrow **A**).

Motor Unit of Skeletal Muscle

Unlike some types of smooth muscle (single-unit type; \rightarrow p. 70) and cardiac muscle fibers, which pass electric stimuli to each other through gap junctions or nexus (\rightarrow A; p. 16f.), skeletal muscle fibers are not stimulated by adjacent muscle fibers, but by *motor neurons*. In fact, muscle paralysis occurs if the nerve is severed.

One motor neuron together with all muscle fibers innervated by it is called a **motor unit** (MU). Muscle fibers belonging to a single motor unit can be distributed over large portions $(1\,\text{cm}^2)$ of the muscle cross-sectional area. To supply its muscle fibers, a motor neuron splits into collaterals with terminal branches (\rightarrow p. 42). A given motor neuron may supply only 25 muscle fibers (mimetic muscle) or well over 1000 (temporal muscle).

Two **types of skeletal muscle fibers** can be distinguished: **S** – slow-twitch fibers (type 1) and **F** – fast-twitch fibers (type 2), including two subtypes, **FR** (2 A) and **FF** (2 B). Since each motor unit contains only one type of fiber, this classification also applies to the motor unit.

Slow-twitch fibers are the least fatigable and are therefore equipped for sustained performance. They have high densities of capillaries and mitochondria and high concentrations of fat droplets (high-energy substrate reserves) and the red pigment myoglobin (short-term O₂ storage). They are also rich in oxidative enzymes (→ p. 72). Fast-twitch fibers are mainly responsible for brief and rapid contractions. They are quickly fatigued (FF > FR) and are rich in glycogen (FF > FR) but contain little myoglobin (FF≪FR).

The **fiber type distribution** of a muscle depends on the muscle type. Motor units of the *S type* predominate in "red" muscles such as the soleus muscle, which helps to maintain the body in an upright position, whereas the *F type* predominates in "white" muscles such as the gastrocnemius muscle, which is involved in running activity. Each fiber type can also be *converted* to the other type. If, for example, the prolonged activation of fast-twitch fibers leads to a chronic increase in the cytosolic Ca²⁺ concentration, fast-twitch fibers will be converted to slow-twitch fibers and vice versa.

Graded muscle activity is possible because a variable number of motor units can be recruited as needed. The more motor units a muscle has, the more finely graded its contractions. Contractions are much finer in the external eve muscles, for example, which have around 2000 motor units, than in the lumbrical muscles, which have only around 100 motor units. The larger the number of motor units recruited, the stronger the contraction. The number and type of motor units recruited depends on the type of movement involved (fine or coarse movement, intermittent or persistent contraction, reflex activity, voluntary or involuntary movement, etc.). In addition, the strength of each motor unit can be increased by increasing the frequency of neuronal impulses, as in the tetanization of skeletal muscle $(\rightarrow p.67 A)$.

Contractile Apparatus of Striated Muscle

The **muscle cell** is a fiber (\rightarrow **A2**) approximately 10 to 100 µm in diameter. Skeletal muscles fibers can be as long as 15 cm. Meat "fibers" visible with the naked eye are actually bundles of muscle fibers that are around 100 to 1000 µm in diameter (\rightarrow **A1**). Each striated muscle fiber is invested by a cell membrane called the *sarcolemma*, which surrounds the *sarcoplasm* (cytoplasm), several cell nuclei, mitochondria (*sarcosomes*), substances involved in supplying O₂ and energy (\rightarrow p. 72), and several hundreds of *myofibrils*.

So-called Z lines or, from a three-dimensional aspect, *Z plates* (plate-like proteins; \rightarrow B) subdivide each **mvofibril** (\rightarrow A3) into approx. 2 µm long, striated compartments called **sarcomeres** $(\rightarrow B)$. When observed by (two-dimensional) microscopy, one can identify alternating light and dark bands and lines (hence the name "striated muscle") created by the thick myosin II filaments and thin actin filaments (\rightarrow B; for myosin I, see p. 30). Roughly 2000 actin filaments are bound medially to the Z plate. Thus, half of the filament projects into two adjacent sarcomeres (\rightarrow **B**). The region of the sarcomere proximal to the Z plate contains only actin filaments, which form a so-called Iband (\rightarrow **B**). The region where the actin and myosin filaments overlap is called the Aband. The Hzone solely contains myosin filaments (ca. 1000 per sarcomere), which thicken towards the middle of the sarcomere to form the M line (M plate). The (actin) filaments are anchored to the sarcolemma by the protein dystrophin.

Each myosin filament consists of a bundle of ca. 300 **myosin-II molecules** (\rightarrow **B**). Each molecule has two globular heads connected by flexible necks (head and neck = subfragment S1; formed after proteolysis) to the filamentous tail of the molecule (two intertwined α -helices = subfragment S2) (\rightarrow **C**). Each of the heads has a motor domain with a nucleotide binding pocket (for ATP or ADP + P₁) and anottin binding site. Two light protein chains are located on each neck of this heavy molecule (220 kDa): one is regulatory (20 kDa), the other essential (17 kDa). Conformational

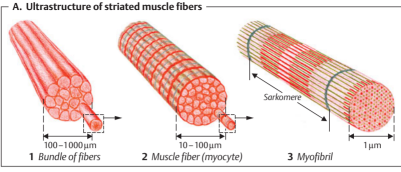
changes in the head–neck segment allow the myosin head to "tilt" when interacting with actin (*sliding filaments*; \rightarrow p. 62).

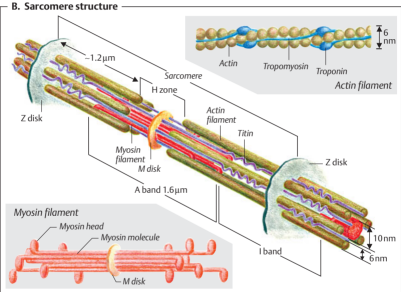
Actin is a globular protein molecule (G-actin). Four hundered such molecules join to form F-actin, a beaded polymer chain. Two of the twisted protein filaments combine to form an **actin filament** (\rightarrow **B**), which is positioned by the equally long protein *nebulin*.

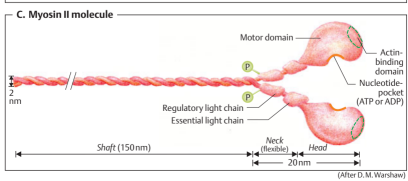
Tropomyosin molecules joined end-to-end (40 nm each) lie adjacent to the actin filament, and a **troponin** (TN) molecule is attached every 40 nm or so $(\rightarrow B)$. Each troponin molecule consists of three subunits: TN-C, which has two regulatory bindings sites for Ca^{2+} at the amino end, TN-I, which prevents the filaments from sliding when at rest $(\rightarrow p. 62)$, and TN-T, which interacts with TN-C, TN-I, and actin.

The sarcomere also has another system of filaments $(\rightarrow B)$ formed by the filamentous protein **titin** (connectin). Titin is more than 1000 nm in length and has some 30 000 amino acids $(M_r > 3000 \, kDa)$. It is the longest known polypeptide chain and comprises 10% of the total muscle mass. Titin is anchored at its carboxyl end to the M plate and, at the amino end, to the Z plate $(\rightarrow p.66$ for functional description).

The sarcolemma forms a Tsystem with several transverse tubules (tube-like invaginations) that run perpendicular to the myofibrils $(\rightarrow p.63 A)$. The endoplasmic reticulum $(\rightarrow p. 10 \text{ ff.})$ of muscle fibers has a characteristic shape and is called the sarcoplasmic reti**culum** (SR; \rightarrow p. 63 A). It forms closed chambers without connections between the intraand extracellular spaces. Most of the chambers run lengthwise to the myofibrils, and are called longitudinal $(\rightarrow p.63 \text{ A})$. The sarcoplasmic reticulum is more prominently developed in skeletal muscle than in the myocardium and serves as a Ca²⁺ storage space. Each T system separates the adjacent longitudinal tubules, forming triads (\rightarrow p. 63 A, B).







Contraction of Striated Muscle

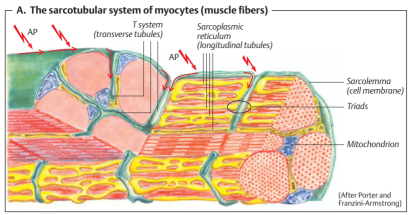
Stimulation of muscle fibers. The release of *acetylcholine* at the motor end-plate of skeletal muscle leads to an *end-plate current* that spreads electrotonically and activates voltagegated Na^+ *channels* in the sarcolemma (\rightarrow p. 56). This leads to the firing of **action potentials** (AP) that travel at a rate of 2 m/s along the sarcolemma of the entire muscle fiber, and penetrate rapidly into the depths of the fiber along the T system (\rightarrow A).

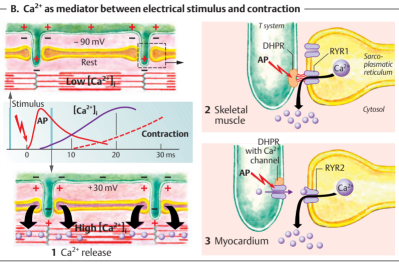
The conversion of this excitation into a contraction is called electromechanical coupling $(\rightarrow B)$. In the skeletal muscle, this process begins with the action potential exciting voltage-sensitive dihydropyridine receptors (DHPR) of the sarcolemma in the region of the triads. The DHPR are arranged in rows, and directly opposite them in the adjacent membrane of the sarcoplasmic reticulum (SR) are rows of Ca2+ channels called ryanodine receptors (type 1 in skeletal muscle: RYR1). Every other RYR1 is associated with a DHPR (\rightarrow **B2**). RYR1 open when they directly "sense" by mechanical means an AP-related conformational change in the DHPR. In the myocardium, on the other hand, each DHPR is part of a voltagegated Ca2+ channel of the sarcolemma that opens in response to an action potential. Small quantities of extracellular Ca2+ enter the cell through this channel, leading to the opening of myocardial RYR2 (so-called trigger effect of Ca^{2+} or Ca^{2+} spark; \rightarrow **B3**). Ca^{2+} ions stored in the SR now flow through the opened RYR1 or RYR2 into the cytosol, increasing the cytosolic Ca²⁺ concentration [Ca²⁺]_i from a resting value of ca. $0.01 \, \text{umol/L}$ to over $1 \, \text{umol/L}$ (\rightarrow **B1**). In skeletal muscle, DHPR stimulation at a single site is enough to trigger the coordinated opening of an entire group of RYR1, thereby increasing the reliability of impulse transmission. The increased cytosolic Ca2+ concentration saturates the Ca2+ binding sites on troponin-C. thereby canceling the troponin-mediated inhibitory effect of tropomyosin on filament sliding $(\rightarrow \mathbf{D})$. It is still unclear whether this type of disinhibition involves actin-myosin binding or the detachment of ADP and Pi, as described below.

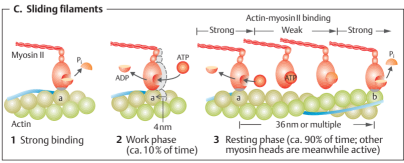
ATP (\rightarrow p. 72) is essential for filament sliding and, hence, for muscle contraction. Due to their ATPase activity, the myosin heads $(\rightarrow p.60)$ act as the motors (motor proteins) of this process. The myosin-II and actin filaments of a sarcomere (\rightarrow p. 60) are arranged in such a way that they can slide past each other. The myosin heads connect with the actin filaments at a particular angle, forming so-called crossbridges (\rightarrow C1). Due to a conformational change in the region of the nucleotide binding site of myosin-II (\rightarrow p. 61 C), the spatial extent of which is increased by concerted movement of the neck region, the myosin head tilts down, drawing the thin filament a length of roughly $4 \text{ nm} (\rightarrow C2)$. The second myosin head may also move an adjacent actin filament. The head then detaches and "tenses" in preparation for the next "oarstroke" when it binds to actin anew (\rightarrow C3).

Kinesin, another motor protein (\rightarrow pp. 42 u. 58), independently advances on the microtubule by incremental movement of its two heads (8 nm increments), as in tug-of-war. In this case, fifty percent of the cycle time is "work time" (duty ratio = 0.5). Between two consecutive interactions with actin in skeletal muscle, on the other hand, myosin-II "jumps" 36 nm (or multiples of 36, e.g. 396 nm or more in rapid contractions) to reach the next (or the 11th) suitably located actin binding site (\rightarrow C3, jump from a to b). Meanwhile, the other myosin heads working on this particular actin filament must make at least another 10 to 100 oarstrokes of around 4 nm each. The duty ratio of a myosin-II head is therefore 0.1 to 0.01. This division of labor by the myosin heads ensures that a certain percentage of the heads will always be ready to generate rapid contractions.

When filament sliding occurs, the Z plates approach each other and the overlap region of thick and thin filaments becomes larger, but the length of the filaments remains unchanged. This results in shortening of the I band and H zone (\rightarrow p. 60). When the ends of the thick filaments ultimately bump against the Z plate, maximum muscle shortening occurs, and the ends of the thin filaments overlap (\rightarrow p. 67 C). Shortening of the sarcomere therefore occurs at both ends of the myosin bundle, but in opposite directions.







binding pocket. The resulting M-ATP complex lies at an approx. 90° angle to the rest of the myosin filament (\rightarrow **D4**). In this state, myosin has only a weak affinity for actin binding. Due to the influence of the increased cytosolic Ca2+ concentration on the troponin - tropomyosin complex, actin (A) activates myosin's ATPase. resulting in hydrolysis of ATP (ATP \rightarrow ADP + P_i) and the formation of an A-M-ADP-Pi complex $(\rightarrow D1)$. Detachment of P_i (inorganic phosphate) from the complex results in a conformational change of myosin that increases the actin-myosin association constant by four powers of ten (binding affinity now strong). The myosin heads consequently tilt to a 40° angle (\rightarrow D2a), causing the actin and myosin filaments to slide past each other. The release of ADP from myosin ultimately brings the myosin heads to their final position, a 45° angle $(\rightarrow D2b)$. The remaining A-M complex (rigor complex) is stable and can again be transformed into a weak bond when the myosin heads bind ATP anew ("softening effect" of ATP). The high flexibility of the muscle at rest is important for processes such as cardiac filling or the relaxing of the extensor muscles during rapid bending movement. If a new ATP is bound to myosin, the subsequent weakening of the actin-myosin bond allows the realignment of the myosin head from 45° to 90° $(\rightarrow D3, 4)$, the position preferred by the M-ATP

Contraction cycle (\rightarrow **C** and **D**). Each of the

two myosin heads (M) of a myosin-II molecule

bind one ATP molecule in their nucleotide

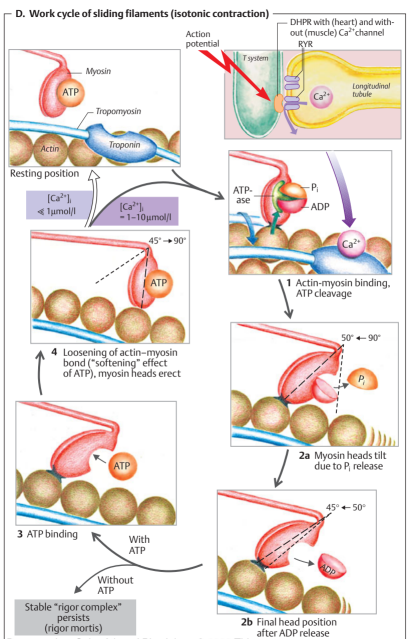
The Ca^{2+} ions released from the sarcoplasmic reticulum (SR) are continuously pumped back to the SR due to active transport by Ca^{2+} -ATPase (\rightarrow pp. 17 A and 26), also called SERCA (\rightarrow p. 16). Thus, if the RYR-mediated release of Ca^{2+} from the SR is interrupted, the cytosolic Ca^{2+} concentration rapidly drops below 10^{-6} mol/L and filament sliding ceases (resting position; \rightarrow D, upper left corner).

complex. If the cytosolic Ca²⁺ concentration remains > 10⁻⁶ mol/L, the **D1** to **D4** cycle will begin anew. This depends mainly on whether subsequent action potentials arrive. Only a portion of the myosin heads that pull actin filaments are "on duty" (low duty ratio; see p. 62) to ensure the smoothness of contractions.

Parvalbumin, a protein that occurs in the cytosol of fast-twitch muscle fibers (\rightarrow type F; p.58), accelerates muscle relaxation after short contractions by binding cytosolic Ca²⁺ in exchange for Mg²⁺. Parvalbumin's binding affinity for Ca²⁺ is higher than that of troponin, but lower than that of SR's Ca²⁺-ATPase. It therefore functions as a "slow" Ca²⁺ buffer.

The course of the filament sliding cycle as described above mainly applies to **isotonic contractions**, that is, to contractions where muscle shortening occurs. During strictly **isometric contractions** where muscular tension increases but the muscle length remains unchanged, the sliding process tenses elastic components of a muscle, e.g. titin (\rightarrow p. 66), and then soon comes to a halt. Afterwards, the A-M-ATP complex (\rightarrow D3) probably transforms directly into A-M-ADP-P_i (\rightarrow D1).

The muscle fibers of a dead body do not produce any ATP. This means that, after death, Ca²⁺ is no longer pumped back into the SR, and the ATP reserves needed to break down stable A-M complexes are soon depleted. This results in stiffening of the dead body or **rigor mortis**, which passes only after the actin and myosin molecules in the muscle fibers decompose.



Mechanical Features of Skeletal Muscle

Action potentials generated in muscle fibers increase the cytosolic Ca2+ concentration [Ca²⁺]_i, thereby triggering a contraction (skeletal muscle; \rightarrow p. 36 B; myocardium; \rightarrow p. 194). In skeletal muscles, gradation of contraction force is achieved by variable recruitment of motor units (\rightarrow p.58) and by changing the action notential frequency. A single stimulus always leads to maximum Ca2+ release and, thus, to a maximum single twitch of skeletal muscle fiber if above threshold (allor-none response). Nonetheless, a single stimulus does not induce maximum shortening of muscle fiber because it is too brief to keep the sliding filaments in motion long enough for the end position to be reached. Muscle shortening continues only if a second stimulus arrives before the muscle has completely relaxed after the first stimulus. This type of stimulus repetition leads to incremental mechanical summation or super**position** of the individual contractions $(\rightarrow A)$. Should the frequency of stimulation become so high that the muscle can no longer relax at all between stimuli, sustained maximum contraction of the motor units or tetanus will occur (\rightarrow A). This occurs, for example, at 20 Hz in slow-twitch muscles and at 60-100 Hz in fast-twitch muscles (\rightarrow p.58). The muscle force during tetanus can be as much as four times larger than that of single twitches. The Ca2+ concentration, which decreases to some extent between superpositioned stimuli, remains high in tetanus.

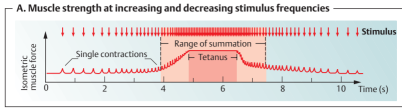
Rigor (→ p.2.13) as well as contracture, another state characterized by persistent muscle shortening, must be distinguished from tetanus. Contracture is not caused by action potentials, but by persistent local depolarization due, for example, to increased extracellular K^+ concentrations (K^+ contracture) or drug-induced intracellular Ca^{2+} release, e.g., in response to caffeine. The contraction of so-called tonus fibers (specific fibers in the external eye muscles and in muscle spindles; → p. 318) is also a form of contracture. Tonus fibers do not respond to stimuli according to the all-or-none law, but contract in proportion

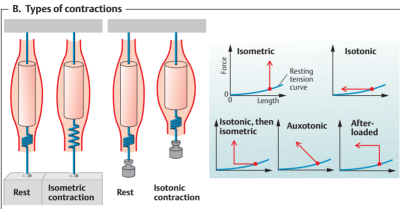
with the magnitude of depolarization. The magnitude of contraction of tonus fibers is regulated by *variation of the cytosolic Ca²⁺ concentration* (not by action potentials!)

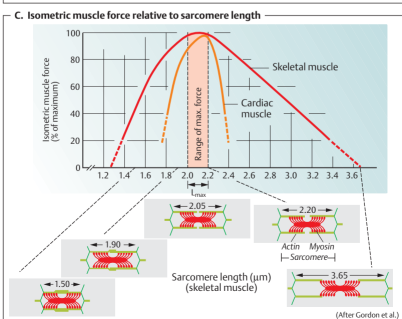
In contrast, the general **muscle tone** (*reflex tone*), or the tension of skeletal muscle at rest, is attributable to the arrival of normal action potentials at the individual motor units. The individual contractions cannot be detected because the motor units are alternately (asynchronously) stimulated. When apparently at rest, muscles such as the postural muscles are in this involuntary state of tension. Resting muscle tone is regulated by reflexes $(\rightarrow p.318\,\mathrm{ff.})$ and increases as the state of attentiveness increases.

Types of contractions $(\rightarrow B)$. There are different types of muscle contractions. In isometric contractions, muscle force ("tension") varies while the length of the muscle remains constant. (In cardiac muscle, this also represents isovolumetric contraction, because the muscle length determines the atrial or ventricular volume.) In isotonic contractions. the length of the muscle changes while muscle force remains constant. (In cardiac muscle, this also represents isobaric contraction, because the muscle force determines the atrial or ventricular pressure.) In auxotonic contractions, muscle length and force both vary simultaneously. An isotonic or auxotonic contraction that builds on an isometric one is called an afterloaded contraction.

Muscle extensibility. A resting muscle containing ATP can be stretched like a rubber band. The force required to start the stretching action (\rightarrow **D**, **E**; extension force at rest) is very small, but increases exponentially when the muscle is under high elastic strain (see resting tension curve, $\rightarrow \mathbf{D}$). A muscle's resistance to stretch, which keeps the sliding filaments in the sarcomeres from separating, is influenced to a small extent by the fascia (fibrous tissue). The main factor, however, is the giant filamentous elastic molecule called titin (or connectin: $1000 \,\mathrm{nm}$ long, $M_r = 3$ to $3.7 \,\mathrm{MDa}$) which is incorporated in the sarcomere (6 titin molecules per myosin filament). In the Aband region of the sarcomere (\rightarrow p. 61 B), titin lies adjacent to a myosin filament and helps to keep it in the center of the sarcomere. Titin molecules in the







I band region are flexible and function as "elastic bands" that counteract passive stretching of a muscle and influence its shortening velocity.

The **extensibility of titin molecules**, which can stretch to up to around ten times their normal length in skeletal muscle and somewhat less in cardiac muscle, is mainly due to frequent repetition of the *PEVK motif* (proline-glutamate-valine-lysine). In very strong muscle extension, which represents the steepest part of the resting extensibility curve (\rightarrow **D**), globular chain elements called immunoglobulin C2 domains also unfold. The quicker the muscle stretches, the more sudden and crude this type of "shock absorber" action will be.

The length (L) and force (F) or "tension" of a muscle are closely related ($\rightarrow C$, E). The total force of a muscle is the sum of its active force and its extension force at rest, as was explained above. Since the active force is determined by the magnitude of all potential actinmyosin interactions, it varies in accordance with the initial sarcomere length $(\rightarrow C, D)$. Skeletal muscle can develop maximum active (isometric) force (F_0) from its resting length (L_{max} ; sarcomere length ca. 2 to 2.2 μ m; \rightarrow **C**). When the sarcomeres shorten ($L < L_{max}$), part of the thin filaments overlap, allowing only forces smaller than F_0 to develop ($\rightarrow \mathbf{C}$). When L is 70% of L_{max} (sarcomere length: 1.65 μ m), the thick filaments make contact with the Z disks, and F becomes even smaller. In addition, a greatly pre-extended muscle ($L > L_{max}$) can develop only restricted force, because the number of potentially available actin-myosin bridges is reduced $(\rightarrow \mathbf{C})$. When extended to 130% or more of the Lmax, the extension force at rest becomes a major part of the total muscle force $(\rightarrow E)$.

The length–force curve corresponds to the **cardiac pressure–volume diagram** in which ventricular filling volume corresponds to muscle length, and ventricular pressure corresponds to muscle force; \rightarrow p. 202. Changes in the cytosolic Ca²⁺ concentration can modify the pressure–volume relationship by causing a change in *contractility* (\rightarrow p. 203 B2).

Other important functional **differences between cardiac muscle and skeletal muscle** are listed below (see also p. 59 A):

Since skeletal muscle is more extensible than the cardiac muscle, the passive extension

force of cardiac muscle at rest is greater than that of skeletal muscle (\rightarrow **E1, 2**).

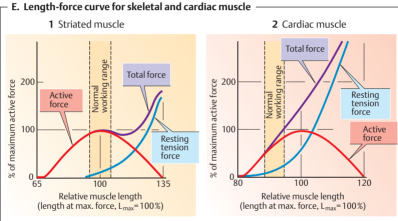
Skeletal muscle normally functions in the plateau region of its length–force curve, whereas cardiac muscle tends to operate in the ascending limb (below L_{max}) of its length–force curve without a plateau (\rightarrow C, E1, 2). Hence, the ventricle responds to increased diastolic filling loads by increasing its force development (Frank–Starling mechanism; \rightarrow p. 204). In cardiac muscle, extension also affects troponin's sensitivity to Ca^{2+} , resulting in a steeper curve (\rightarrow E2).

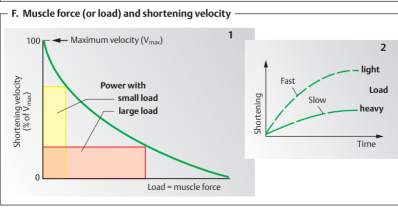
Action potentials in cardiac muscle are of much longer duration than those in skeletal muscle (\rightarrow p.59 A) because g_K temporarily decreases and g_{Ca} increases for 200 to 500 ms after rapid inactivation of Na⁺ channels. This allows the slow influx of Ca²⁺, causing the action potential to reach a *plateau*. As a result, the refractory period does not end until a contraction has almost subsided (\rightarrow p.59 A). Therefore, *tetanus cannot be evoked in cardiac muscle*

Unlike skeletal muscle, cardiac muscle has no motor units. Instead, the stimulus spreads across all myocardial fibers of the atria and subsequently of the ventricles generating an all-or-none contraction of both atria and, thereafter, both ventricles.

In cardiac muscle but not in skeletal muscle, the duration of an action potential can change the force of contraction, which is controlled by the variable influx of Ca²⁺ into the cell.

The greater the force (load), the lower the **velocity** of an (isotonic) contraction (see velocity–force diagram, **F1**). Maximal force and a small amount of heat will develop if shortening does not occur. The maximal velocity (biceps: ca. 7 m/s) and a lot of heat will develop in muscle without a stress load. Light loads can therefore be picked up more quickly than heavy loads (\rightarrow **F2**). The total amount of energy consumed for work and heat is greater in isotonic contractions than in isometric ones. **Muscle power** is the product of force and the shortening velocity: $N \cdot m \cdot s^{-1} = W (\rightarrow$ **F1**, colored areas).





Smooth Muscle

Smooth muscle (SmM) consists of multiple layers of spindle-shaped cells. It is involved in the function of many organs (stomach, intestine, gall bladder, urinary bladder, uterus, bronchi, eves, etc.) and the blood vessels. where it plays an important role in circulatory control. SmM contains a special type of Factin-tropomyosin and myosin II filaments $(\rightarrow p.60)$, but lacks troponin and myofibrils. Furthermore, it has no distinct tubular system and no sarcomeres (nonstriated). It is therefore called smooth muscle because of this lack of striation (see p. 59 A for further differences in the muscle types). SmM filaments form a loose contractile apparatus arranged approximately longitudinally within the cell and attached to discoid plagues (see B for model). which also provide a mechanical means for cell-cell binding of SmM. Smooth muscle can shorten much more than striated muscle.

The **membrane potential** of the SmM cells of many organs (e.g., the intestine) is not constant, but fluctuates rhythmically at a low frequency (3 to 15 min⁻¹) and amplitude (10 to 20 mV), producing slow waves. These waves trigger a burst of action potentials (spikes) when they exceed a certain threshold potential. The longer the slow wave remains above the threshold potential, the greater the number and frequency of the action potentials it produces. A relatively sluggish contraction occurs around 150 ms after a spike (\rightarrow p. 59 A, left panel). Tetanus occurs at relatively low spike frequencies (\rightarrow p. 66). Hence, SmM is constantly in a state of a more or less strong contraction (tonus or tone). The action potential of SmM cells of some organs has a plateau similar to that of the cardiac action potential $(\rightarrow p.59 A. middle panel)$.

There are two types of smooth muscles $(\rightarrow A)$. The cells of **single-unit SmM** are electrically coupled with each other by *gap junctions* $(\rightarrow pp. 18 \text{ and } 50)$. Stimuli are passed along from cell to cell in organs such as the stomach, intestine, gallbladder, urinary bladder, ureter, uterus, and some types of blood vessels. Stimuli are generated autonomously from within the SmM, partly by pacemaker cells). In other words, the stimulus is innervation-inde-

pendent and, in many cases, spontaneous (*myogenic tonus*). The second type, **multi-unit SmM**, contracts primarily due to stimuli from the autonomic nervous system (*neurogenic tonus*). This occurs in structures such as the arterioles, spermatic ducts, iris, ciliary body, and the muscles at the roots of the hair. Since these SmM cells generally are not connected by gap junctions, stimulation remains localized, as in the motor units of the skeletal muscle.

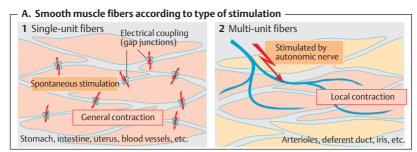
Smooth muscle tonus is regulated by the degree of depolarization (e.g., through stretch or pacemaker cells) as well as by transmitter substances (e.g., acetylcholine or noradrenaline) and numerous hormones (e.g., estrogens, progesterone and oxytocin in the uterus and histamine, angiotensin II, adjuretin, serotonin and bradykinin in vascular muscle). An increase in tonus will occur if any of these factors directly or indirectly increases the cytosolic Ca2+ concentration to more than 10-6 mol/L. The Ca2+ influx comes mainly from extracellular sources, but a small portion comes from intracellular stores (\rightarrow B1). Ca²⁺ ions bind to calmodulin (CM) (\rightarrow B2), and Ca²⁺-CM promotes contraction in the following manner.

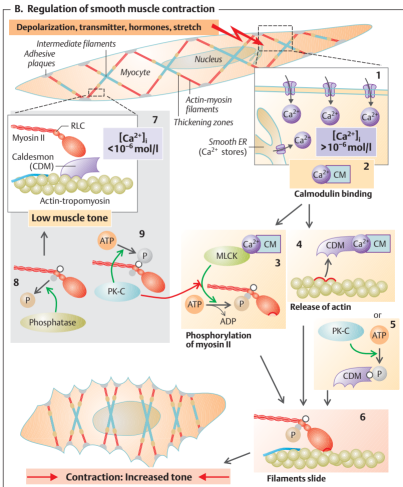
Regulation at myosin II (\rightarrow B3): The Ca²⁺-CM complex activates myosin light chain kinase (MLCK), which phosphorylates myosin's regulatory light chain (RLC) in a certain position, thereby enabling the myosin head to interact with actin (\rightarrow B6).

Regulation at the actin level (\rightarrow B4). The Ca²⁺-CM complex also binds with caldesmon (CDM), which then detaches from the actintropomyosin complex, thus making it available for filament sliding (\rightarrow B6). Phosphorylation of CDM by protein kinase C (PK-C) also seems to be able to induce filament sliding (\rightarrow B5).

Factors that lead to a **reduction of tonus** are: reduction of the cytosolic Ca²⁺ concentration to less than 10^{-6} mol/L (\rightarrow B7), phosphatase activity (\rightarrow B8), and PK-C if it phosphorylates another position on the RLC (\rightarrow B9).

When length-force curves are recorded for smooth muscle, the curve shows that muscle force decreases continuously while muscle length remains constant. This property of a muscle is called **plasticity**.





Adenosine triphosphate (ATP) is a direct source of chemical energy for muscle contraction (\rightarrow A, pp. 40 and 64). However, a muscle cell contains only a limited amount of ATP-only enough to take a sprinter some 10 to 20 m or so. Hence, spent ATP is continuously regenerated to keep the intracellular ATP concentration constant, even when large quantities of it are needed. The three routes of ATP regeneration are (\rightarrow B):

- 1. Dephosphorylation of creatine phosphate
- 2. Anaerobic glycolysis
- 3. Aerobic oxidation of glucose and fatty acids. Routes 2 and 3 are relatively slow, so **creatine phosphate** (**CrP**) must provide the chemical energy needed for *rapid ATP regeneration*. ADP derived from metabolized ATP is immediately transformed to ATP and creatine (Cr) by mitochondrial creatine kinase (→ **B1** and p. 40). The CrP reserve of the muscle is sufficient for short-term high-performance bursts of 10−20 s (e.g., for a 100-m sprint).

Anaerobic glycolysis occurs later than CrP dephosphorylation (after a maximum of ca. 30 s). In anaerobic glycolysis, muscle glycogen is converted via glucose-6-phosphate to lactic acid (→ lactate + H⁺), yielding 3 ATP molecules for each glucose residue (→ B2). During light exercise, lactate⁻ is broken down in the heart and liver whereby H⁺ ions are used up. Aerobic oxidation of glucose and fatty acids takes place approx. 1 min after this less productive anaerobic form of ATP regeneration. If aerobic oxidation does not produce a sufficient supply of ATP during strenuous exercise, anaerobic glycolysis must also be continued.

In this case, however, glucose must be imported from the liver where it is formed by glycogenolysis and gluconeogenesis (see also p. 282f.). Imported glucose yields only two ATP for each molecule of glucose, because one ATP is required for 6-phosphorylation of glucose.

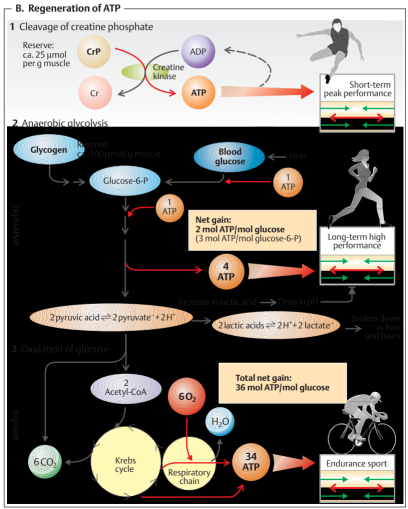
Aerobic regeneration of ATP from glucose (2+34 ATP per glucose residue) or fatty acids is required for *sustained exercise* (\rightarrow **B3**). The cardiac output (= heart rate \times stroke volume) and total ventilation must therefore be increased to meet the increased metabolic requirements of the muscle; the heart rate then becomes

constant (\rightarrow p. 75 B). The several minutes that pass before this *steady state* is achieved are bridged by anaerobic energy production, *increased O*₂ *extraction* from the blood and depletion of short-term O₂ reserves in the muscle (*myoglobin*). The interim between the two phases is often perceived as the "low point" of physical performance.

The O_2 affinity of **myoglobin** is higher than that of hemoglobin, but lower than that of respiratory chain enzymes. Thus, myoglobin is normally saturated with O_2 and can pass on its oxygen to the mitochondria during brief arterial oxygen supply deficits.

The endurance limit, which is some 370 W ($\approx 0.5 \text{ HP}$) in top athletes, is mainly dependent on the speed at which O₂ is supplied and on how fast aerobic oxidation takes place. When the endurance limit is exceeded, steady state cannot occur, the heart rate then rises continuously (\rightarrow p. 75 B). The muscles can temporarily compensate for the energy deficit (see above), but the H+-consuming lactate metabolism cannot keep pace with the persistently high level of anaerobic ATP regeneration. An excess of lactate and H+ ions, i.e. lactacidosis, therefore develops. If an individual exceeds his or her endurance limit by around 60%, which is about equivalent to maximum O2 consumption $(\rightarrow p.74)$, the plasma lactate concentration will increase sharply, reaching the so-called anaerobic threshold at 4 mmol/L. No significant increase in performance can be expected after that point. The systemic drop in pH results in increasing inhibition of the chemical reactions needed for muscle contraction. This ultimately leads to an ATP deficit, rapid muscle fatique and, finally, a stoppage of muscle work.

CrP metabolism and anaerobic glycolysis enable the body to achieve three times the performance possible with aerobic ATP regeneration, albeit for only about 40 s. However, these processes result in an O_2 deficit that must be compensated for in the post-exercise recovery phase (O_2 debt). The body "pays off" this debt by regenerating its energy reserves and breaking down the excess lactate in the liver and heart. The O_2 debt after strenuous exercise is much larger (up to $20\,L$) than the O_2 deficit for several reasons.



Physical Work

There are three types of muscle work:

- Positive dynamic work, which requires to muscles involved to alternately contract and relax (e.g., going uphill).
- Negative dynamic work, which requires the muscles involved to alternately extend while braking (braking work) and contract without a load (e.g., going downhill).
- ◆ Static postural work, which requires continuous contraction (e.g., standing upright). Many activities involve a combination of two or three types of muscle work. Outwardly directed mechanical work is produced in dynamic muscle activity, but not in purely postural work. In the latter case, force × distance = 0. However, chemical energy is still consumed and completely transformed into a form of heat called maintenance heat (= muscle force times the duration of postural work).

In strenuous exercise, the muscles require up to 500 times more O_2 than when at rest. At the same time, the muscle must rid itself of metabolic products such as H^+ , CO_2 , and lactate (\rightarrow p.72). Muscle work therefore requires drastic cardiovascular and respiratory changes.

In untrained subjects (UT), the cardiac output (CO: \rightarrow p. 186) rises from 5-6 L/min at rest to a maximum of 15-20 L/min during exercise $(\rightarrow p. 77 C)$. Work-related activation of the sympathetic nervous system increases the heart rate up to ca. 2.5 fold and the stroke volume up to ca. 1.2 fold (UT). In light to moderate exercise, the heart rate soon levels out at a new constant level, and no fatigue occurs. Very strenuous exercise, on the other hand, must soon be interrupted because the heart cannot achieve the required long-term performance $(\rightarrow B)$. The increased CO provides more blood for the muscles $(\rightarrow A)$ and the skin (heat loss; \rightarrow p. 222.). The blood flow in the kidney and intestine, on the other hand, is reduced by the sympathetic tone below the resting value $(\rightarrow A)$. The systolic **blood pressure** $(\rightarrow p.206)$ rises while the diastolic pressure remains constant, yielding only a moderate increase in the mean pressure.

The smaller the muscle mass involved in the work, the higher the increase in blood pressure. Hence, the blood pressure increase in arm activity (cutting hedges) is higher than that in leg activity (cycling). In patients with coronary artery disease or cerebrovascular sclerosis, arm activity is therefore more dangerous than leg activity due to the risk of myocardial infarction or brain hemorrhage.

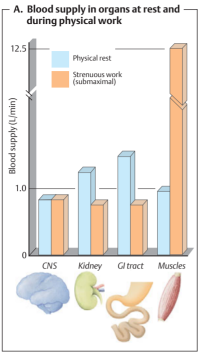
Muscular blood flow. At the maximum work level, the blood flow in 1 kg of active muscle rises to as much as $2.5\,\mathrm{L/min}$ (\rightarrow p. 213 A), equivalent to 10% of the maximum cardiac output. Hence, no more than 10 kg of muscle (<1/3 the total muscle mass) can be fully active at any one time. Vasodilatation, which is required for the higher blood flow, is mainly achieved through *local chemical influences* (\rightarrow p. 212). In purely postural work, the increase in blood flow is prevented in part by the fact that the continuously contracted muscle squeezes its own vessels. The muscle then *fatigues* faster than in rhythmic dynamic work.

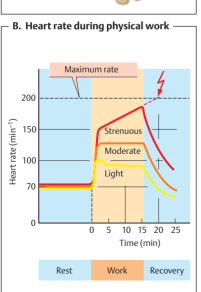
During physical exercise (\rightarrow C1), the ventilation (\dot{V}_E) increases from a resting value of ca. 7.5 L/min to a maximum of 90 to 120 L/min (\rightarrow C3). Both the respiratory rate (40–60 min⁻¹ max; \rightarrow C2) and the tidal volume (ca. 2 L max.) contribute to this increase. Because of the high \dot{V}_E and increased CO, oxygen consumption (\dot{V}_{O_2}) can increase from ca. 0.3 L/min at rest to a maximum (\dot{V}_{O_2} max) of ca. 3 L/min in UT (\rightarrow C4 and p. 76). Around 25 L of air has to be ventilated to take up 1 L of O₂ at rest, corresponding to a respiratory equivalent (\dot{V}_E/\dot{V}_{O_2}) of 25. During physical exercise, \dot{V}_E/\dot{V}_{O_2} rises beyond the endurance limit to a value of 40–50.

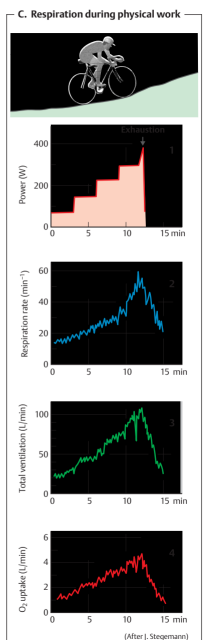
Increased O_2 extraction in the tissues also contributes to the large increase in \dot{V}_{O_2} during exercise. The decreasing pH and increasing temperature shift the O_2 binding curve towards the right (\rightarrow p. 129 B). O_2 extraction is calculated as the arteriovenous difference in O_2 concentration (avDo₂ in L/L blood) times the blood flow in L/min. The maximum O_2 consumption (\dot{V}_{O_2} max) is therefore defined as:

 \dot{V}_{O_2} max = HRmax · SVmax · avD_{O2}max

where HR is the heart rate and SV is the stroke volume. \dot{V}_{O_2} max per body weight is an ideal measure of *physical exercise capacity* (\rightarrow p. 76).







Physical Fitness and Training

The **physical exercise capacity** can be measured using simple yet standardized techniques of **ergometry**. This may be desirable in athletes, for example, to assess the results of training, or in patients undergoing rehabilitation therapy. Ergometry assesses the effects of exercise on physiological parameters such as O_2 consumption (\dot{V}_{O_2}), respiration rate, heart rate (\rightarrow p. 74), and the plasma lactate concentration (\rightarrow A). The measured physical power (performance) is expressed in watts (W) or W/kg body weight (BW).

In bicycle ergometry, a brake is used to adjust the watt level. In "uphill" ergometry on a treadmill set at an angle α , exercise performance in watts is calculated as a factor of body mass (kg) × gravitational acceleration g ($m \cdot s^{-2}$) × distance traveled (m) × sin $\alpha \times 1$ / time required (s^{-1}). In the Margaria step test, the test subject is required to run up a staircase as fast as possible after a certain starting distance. Performance is then measured as body mass (kg) × g ($m \cdot s^{-2}$) × height/time ($m \cdot s^{-1}$).

Short-term performance tests $(10-30 \, s)$ measure performance achieved through the rapidly available energy reserves (creatine phosphate, glycogen). Medium-term performance tests measure performance fueled by anaerobic glycolysis (\rightarrow p. 72). The **maximum O2 consumption** ($\dot{\mathbf{V}}_{02}$ **max**) is used to measure longer term aerobic exercise performance achieved through oxidation of glucose and free fatty acids (\rightarrow p. 74).

In strenuous exercise (roughly 2/3 the maximum physical capacity or more), the aerobic mechanisms do not produce enough energy, so anaerobic metabolism must continue as a parallel energy source. This results in lactacidosis and a sharp increase in the plasma lactate concentration (\rightarrow **A**). **Lactate concentrations** of up to $2 \, \text{mmol/L}$ (aerobic threshold) can be tolerated for prolonged periods of exercise. Lactate concentrations above $4 \, \text{mmol/L}$ (anaerobic threshold) indicate that the performance limit will soon be reached. Exercise must eventually be interrupted, not because of the increasing lactate concentration, but because of the increasing level of acidosis (\rightarrow p. 74).

Physical training raises and maintains the physical exercise capacity. There are three

types of physical training strategies, and most training programs use a combination of them.

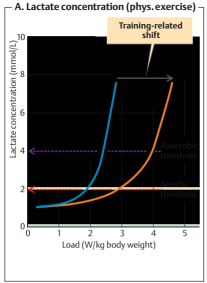
Motor learning, which increases the rate and accuracy of motor skills (e.g., typewriting). These activities primarily involve the CNS.

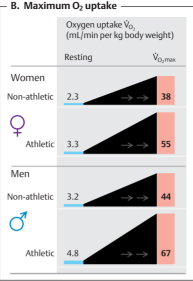
Endurance training, which improves submaximal long-term performance (e.g., running a marathon). The main objectives of endurance training are to increase the oxidative capacity of slow-twitch motor units (\rightarrow p. 58). e.g., by increasing the mitochondrial density, increase the cardiac output and, consequently, to increase \dot{V}_{0} , max (\rightarrow **B**, **C**). The resulting increase in heart weight allows higher stroke volumes $(\rightarrow \mathbf{C})$ as well as higher tidal volumes. resulting in very low resting heart rates and respiratory rates. Trained athletes can therefore achieve larger increases in cardiac output and ventilation than untrained subjects (\rightarrow C). The \dot{V}_{0_2} max of a healthy individual is limited by the cardiovascular capacity, not the respiratory capacity. In individuals who practice endurance training, the exercise-related rise in the lactate concentraton is also lower and occurs later than in untrained subjects $(\rightarrow A)$.

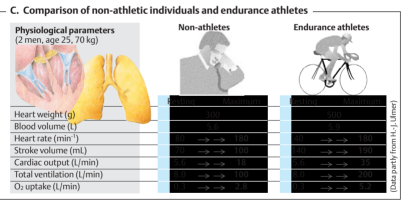
Strength training improves the maximum short-term performance level (e.g., in weight lifting). The main objectives are to increase the muscle mass by increasing the size of the muscle fibers (hypertrophy) and to increase the glycolytic capacity of type motor units (\rightarrow p.58).

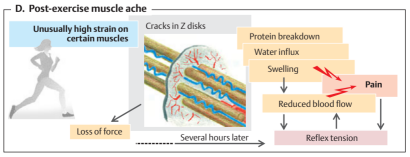
Excessive physical exercise causes **muscle soreness and stiffness.** The underlying cause is not lactic acid accumulation, but sarcomere microtrauma, which leads to muscle swelling and pain. The muscle ache, is a sign of microinflammation (\rightarrow **D**).

Muscle fatigue may be peripheral or central. *Peripheral fatigue* ist caused by the exhaustion of energy reserves and the accumulation of metabolic products in the active muscle. This is particularly quick to occur during postural work (→ p. 66). *Central fatigue* is characterized by work-related pain in the involved muscles and joints that prevents the continuation of physical exercise or decreased the individual's motivation to continue the exercise.









78

3

Autonomic Nervous System (ANS)

Organization of the Autonomic Nervous System

In the somatic nervous system, nerve fibers extend to and from the skeletal muscles, skin and sense organs. They usually emit impulses in response to stimuli from the outside environment, as in the withdrawal reflex (\rightarrow p. 320). Much somatic nervous activity occurs consciously and under voluntary control. In contrast, the autonomic nervous system (ANS) is mainly concerned with regulation of circulation and internal organs. It responds to changing outside conditions by triggering orthostatic responses, work start reactions, etc. to regulate the body's internal environment (\rightarrow p. 2). As the name implies, most activities of the ANS are not subject to voluntary control.

For the most part, the autonomic and somatic nervous systems are anatomically and functionally separate in the periphery $(\rightarrow A)$, but closely connected in the central nervous system, CNS $(\rightarrow p. 266)$. The peripheral ANS is efferent, but most of the nerves containing ANS fibers hold also afferent neurons. These are called visceral afferents because their signals originate from visceral organs, such as the esophagus, gastrointestinal (GI) tract, liver, lungs, heart, arteries, and urinary bladder. Some are also named after the nerve they accompany (e.g., vagal afferents).

Autonomic nervous activity is usually regulated by the reflex arc, which has an afferent limb (visceral and/or somatic afferents) and an efferent limb (autonomic and/or somatic efferents). The afferent fibers convey stimuli from the skin (e.g. nociceptive stimuli; \rightarrow p. 316) and nocisensors, mechanosensors and chemosensors in organs such as the lungs, gastrointestinal tract, bladder, vascular system and genitals. The ANS provides the autonomic efferent fibers that convey the reflex response to such afferent information, thereby inducing smooth muscle contraction (\rightarrow p. 70) in organs such as the eye, lung, digestive tract and bladder, and influencing the function of the heart $(\rightarrow p. 194)$ and glands. Examples of somatic nervous system involvement are afferent stimuli from the skin and sense organs (e.g., light stimuli) and efferent impulses to the skeletal muscles (e.g., coughing and vomiting).

Simple reflexes can take place within an organ (e.g., in the gut, \rightarrow p. 244), but complex reflexes are controlled by **superordinate autonomic centers** in the CNS, primarily in the *spinal cord* (\rightarrow A). These centers are controlled by the *hypothalamus*, which incorporates the ANS in the execution of its programs (\rightarrow p. 330). The *cerebral cortex* is an even higher-ranking center that integrates the ANS with other systems.

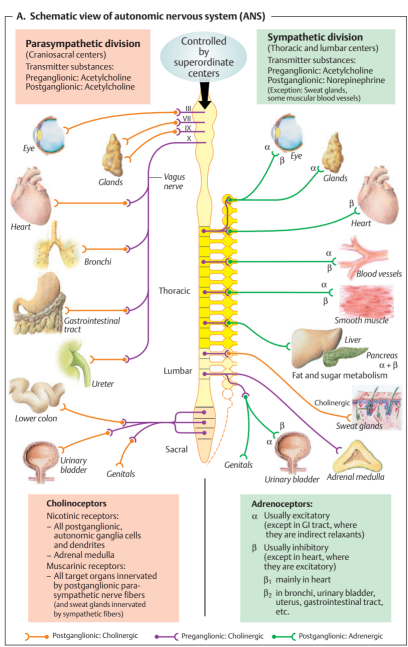
The **peripheral ANS** consists of a sympathetic division and a parasympathetic division (\rightarrow **A**) which, for the most part, are separate entities (\rightarrow also p. 80ff.). The autonomic centers of the sympathetic division lie in the thoracic and lumbar levels of the spinal cord, and those of the parasympathetic division lie in the brain stem (eyes, glands, and organs innervated by the vagus nerve) and sacral part of the spinal cord (bladder, lower parts of the large intestine, and genital organs). (\rightarrow **A**). Preganglionic fibers of both divisions of the ANS extend from their centers to the **ganglia**, where they terminate at the postganglionic neurons.

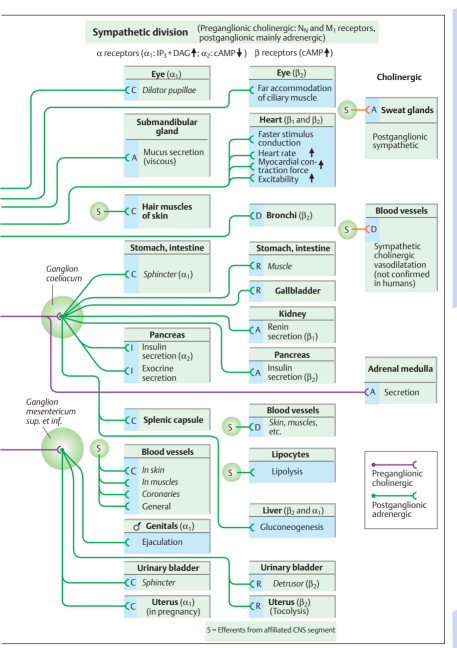
Preganglionic **sympathetic neurons** arising from the spinal cord terminate either in the *paravertebral ganglionic chain*, in the *cervical or abdominal ganglia* or in so-called terminal ganglia. Transmission of stimuli from preganglionic to postganglionic neurons is *cholinergic*, that is, mediated by release of the neurotransmitter *acetylcholine* (\rightarrow p. 82). Stimulation of all effector organs except sweat glands by the postganglionic sympathetic fibers is *adrenergic*, i.e., mediated by the release of *norepinephrine* (\rightarrow A and p. 84ff.).

Parasympathetic ganglia are situated near or within the effector organ. Synaptic transmissions in the parasympathetic ganglia and at the effector organ are *cholinergic* (\rightarrow **A**).

Most organs are innervated by sympathetic and parasympathetic nerve fibers. Nonetheless, the organ's response to the two systems can be either antagonistic (e.g., in the heart) or complementary (e.g., in the sex organs).

The **adrenal medulla** is a ganglion and hormone gland combined. Preganglionic sympathetic fibers in the adrenal medulla release acetylcholine, leading to the secretion of *epinephrine* (and some norepinephrine) *into the bloodstream* (\rightarrow p.86).





Acetylcholine (ACh) serves as a neurotransmitter not only at motor end plates (\rightarrow p. 56) and in the central nervous system, but also in the **autonomic nervous system**, ANS (\rightarrow p. 78ff.), where it is active

- in all preganglionic fibers of the ANS;
- in all parasympathetic postganglionic nerve endings;
- and in some sympathetic postganglionic nerve endings (sweat glands).

Acetylcholine synthesis. ACh is synthesized in the cytoplasm of nerve terminals, and acetyl coenzyme A (acetyl-CoA) is synthesized in mitochondria. The reaction acetyl-CoA + choline is catalyzed by *choline acetyltransferase*, which is synthesized in the soma and reaches the nerve terminals by axoplasmic transport (\rightarrow p. 42). Since choline must be taken up from extracellular fluid by way of a carrier, this is the ratelimiting step of ACh synthesis.

Acetylcholine release. Vesicles on presynaptic nerve terminals empty their contents into the synaptic cleft when the cytosolic Ca^{2+} concentration rises in response to incoming action potentials (AP) (\rightarrow A, p. 50ff.). Epinephrine and norepinephrine can *inhibit ACh release* by stimulating presynaptic α_2 -adrenoceptors (\rightarrow p. 84). In postganglionic parasympathetic fibers, ACh blocks its own release by binding to presynaptic autoreceptors (M-receptors; see below), as shown in B.

ACh binds to postsynaptic **cholinergic receptors** or **cholinoceptors** in autonomic ganglia and organs innervated by parasympathetic fibers, as in the heart, smooth muscles (e.g., of the eye, bronchi, ureter, bladder, genitals, blood vessels, esophagus, and gastrointestinal tract), salivary glands, lacrimal glands, and (sympathetically innervated) sweat glands (\rightarrow p. 80ff.). Cholinoceptors are nicotinic (N) or muscarinic (M). **N-cholinoceptors** (*nicotinic*) can be stimulated by the alkaloid nicotine, whereas *M-cholinoceptors* (*muscarinic*) can be stimulated by the alkaloid mushroom poison muscarine.

Nerve-specific N_N -cholinoceptors on autonomic ganglia (\rightarrow **A**) differ from musclespecific N_M -cholinoceptors on motor end plates (\rightarrow p.56) in that they are formed by

different subunits. They are similar in that they are both *ionotropic receptors*, i.e., they act as cholinoceptors and cation channels at the same time. ACh binding leads to rapid Na^+ and Ca^{2+} influx and in early (rapid) excitatory post-synaptic potentials (EPSP; \rightarrow p. 50ff.), which trigger postsynaptic action potentials (AP) once they rise above threshold (\rightarrow A, left panel).

M-cholinoceptors (M_1-M_5) indirectly affect synaptic transmission through G-proteins (metabotropic receptors).

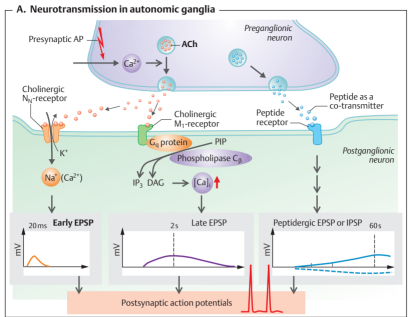
M₁-**cholinoceptors** occur mainly on *autonomic ganglia* (\rightarrow **A**), *CNS*, and *exocrine gland cells*. They activate phospholipase Cβ (PLCβ) via G_q protein in the postganglionic neuron. and inositol *tris*-phosphate (IP₃) and diacylglycerol (DAG) are released as second messengers (\rightarrow p. 276) that stimulate Ca²⁺ influx and a *late EPSP* (\rightarrow **A**, **middle panel**). Synaptic signal transmission is modulated by the late EPSP as well as by co-transmitting peptides that trigger *peptidergic EPSP or IPSP* (\rightarrow **A**, **right panel**).

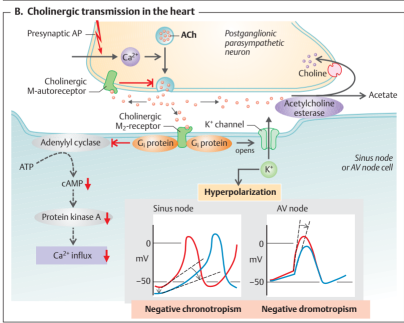
M₂-cholinoceptors occur in the *heart* and function mainly via a G_i protein $(\rightarrow p.274 \text{ ff})$. The G_i protein *opens specific* K^+ *channels* located mainly in the sinoatrial node, atrioventricular (AV) node, and atrial cells, thereby exerting negative chronotropic and dromotropic effects on the heart $(\rightarrow B)$. The G_i protein also *inhibits adenylate cyclase*, thereby reducing Ca^{2+} influx $(\rightarrow B)$.

 M_3 -cholinoceptors occur mainly in smooth muscles. Similar to M_1 -cholinoceptors (\rightarrow A, middle panel), M_3 -cholinoceptors trigger contractions by stimulating Ca^{2+} influx (\rightarrow p. 70). However, they can also induce relaxation by activating Ca^{2+} -dependent NO synthase, e.g., in endothelial cells (\rightarrow p. 278).

Termination of ACh action is achieved by *acetylcholinesterase*-mediated cleavage of ACh molecules in the synaptic cleft (\rightarrow p.56). Approximately 50% of the liberated choline is reabsorbed by presynaptic nerve endings (\rightarrow **B**).

Antagonists. Atropine blocks all M-cholinoceptors, whereas pirenzepine selectively blocks M_1 -cholinoceptors, tubocurarine blocks N_M -cholinoceptors (\rightarrow p. 56), and trimetaphan blocks N_N -cholinoceptors.





Certain neurons can enzymatically produce *L-dopa* (L-dihydroxyphenylalanine) from the amino acid L-tyrosine. L-dopa is the parent substance of dopamine, norepinephrine, and epinephrine—the three natural **cate-cholamines**, which are enzymatically synthesized in this order. Dopamine (DA) is the final step of synthesis in neurons containing only the enzyme required for the first step (the *aromatic L-amino acid decarboxylase*). Dopamine is used as a transmitter by the dopaminergic neurons in the *CNS* and by autonomic neurons that innervate the *kidney*.

Norepinephrine (NE) is produced when a second enzyme (dopamine-β-hydroxylase) is also present. In most sympathetic postganglionic nerve endings and noradrenergic central neurons, NE serves as the neurotransmitter along with the co-transmitters adenosine triphosphate (ATP), somatostatin (SIH), or neuropeptide Y (NPY).

Within the adrenal medulla (see below) and adrenergic neurons of the medulla oblongata, *phenylethanolamine N-methyltransferase* transforms norepinephrine (NE) into **epinephrine (E)**.

The endings of unmyelinated sympathetic postganglionic neurons are knobby or varicose $(\rightarrow A)$. These knobs establish synaptic contact, albeit not always very close, with the effector organ. They also serve as sites of NE synthesis and **storage**. L-tyrosine $(\rightarrow A1)$ is actively taken up by the nerve endings and transformed into dopamine. In adrenergic stimulation, this step is accelerated by protein kinase A-mediated (PKA; \rightarrow A2) phosphorylation of the responsible enzyme. This yields a larger dopamine supply. Dopamine is transferred to chromaffin vesicles, where it is transformed into NE (\rightarrow A3). Norepinephrine, the end product, inhibits further dopamine synthesis (negative feedback).

NE release. NE is exocytosed into the synaptic cleft after the arrival of action potentials at the nerve terminal and the initiation of Ca^{2+} influx (\rightarrow **A4** and p. 50).

Adrenergic receptors or **adrenoceptors** $(\rightarrow \mathbf{B})$. Four main types of adrenoceptors $(\alpha_1, \alpha_2, \alpha_3, \alpha_4, \alpha_5)$

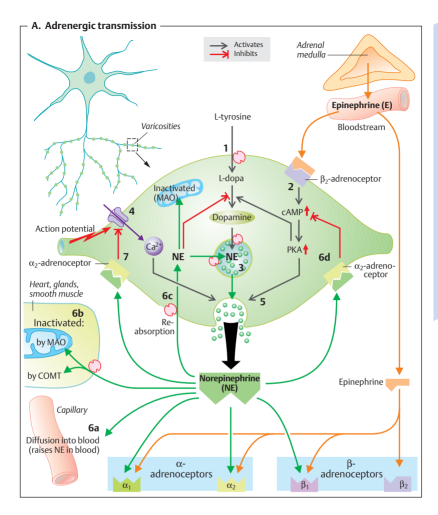
 α_2 , β_1 and β_2) can be distinguished according to their affinity to E and NE and to numerous agonists and antagonists. All adrenoceptors respond to E, but NE has little effect on β_2 -adrenoceptors. Isoproterenol (isoprenaline) activates only β -adrenoceptors, and phentolamine only blocks α -adrenoceptors. The activities of all adrenoceptors are mediated by G proteins (\rightarrow p. 55).

Different subtypes $(\alpha_{1A}, \alpha_{1B}, \alpha_{1D})$ of α_{1} -adrenoceptors can be distinguished $(\rightarrow B1)$. Their location and function are as follows: CNS (sympathetic activity \uparrow), salivary glands, liver (glycogenolysis \uparrow), kidneys (alters threshold for renin release; \rightarrow p. 184), and smooth muscles (trigger contractions in the arterioles, uterus, deferent duct, bronchioles, urinary bladder, gastrointestinal sphincters, and dilator pupillae).

Activation of α_1 -adrenoceptors (\rightarrow **B1**), mediated by G_q proteins and phospholipase $C\beta$ (PLC β), leads to formation of the second messengers inositol tris-phosphate (IP₃), which increases the cytosolic Ca²⁺ concentration, and diacylglycerol (DAG), which activates protein kinase C (PKC; see also p. 276). G_q protein-mediated α_1 -adrenoceptor activity also activates Ca²⁺-dependent K^+ channels. The resulting K^+ outflow hyperpolarizes and relaxes target smooth muscles, e.g., in the gastrointestinal tract.

Three subtypes (α_{2A} , α_{2B} , α_{2C}) of α_{2} -adreno**ceptors** (\rightarrow **B2**) can be distinguished. Their location and action are as follows: CNS (sympathetic activity \downarrow , e.g., use of the α_2 agonist clonidine to lower blood pressure), salivary glands (salivation ↓), pancreatic islets (insulin secretion ↓), lipocytes (lipolysis ↓), platelets (aggregation ↑), and neurons (presynaptic autoreceptors, see below). Activated α_2 -adrenoceptors (\rightarrow **B2**) link with G_i protein and inhibit (via α_i subunit of G_i) adenylate cyclase (cAMP synthesis \downarrow , \rightarrow p. 274) and, at the same time, increase (via the $\beta\gamma$ subunit of G_i) the openprobability of voltage-gated K+ channels (hyperpolarization). When coupled with Go proteins, activated α2-adrenoceptors also inhibit voltage-gated Ca^{2+} channels ($[Ca^{2+}]_i \downarrow$).

All β -adrenoceptors are coupled with a G_S protein, and its α_S subunit releases cAMP as a second messenger. cAMP then activates pro-



tein kinase A (PKA), which phosphorylates different proteins, depending on the target cell type (\rightarrow p. 274).

NE and E work via β_1 -adrenoceptors (\rightarrow B3) to open L-type Ca²⁺ channels in *cardiac cell* membranes. This increases the $[Ca^{2+}]_i$ and therefore produces *positive chronotropic, dromotropic, and inotropic effects*. Activated G_s protein can also directly increase the open-

probability of voltage-gated Ca^{2+} channels in the heart. In the *kidney*, the basal renin secretion is increased via β_1 -adrenoceptors.

Activation of β_2 -adrenoceptors by epinephrine $(\rightarrow B4)$ increases cAMP levels, thereby lowering the $[Ca^{2+}]_i$ (by a still unclear mechanism). This *dilates* the bronchioles and blood vessels of skeletal muscles and *relaxes* the muscles of the uterus, deferent duct, and

Heat production is increased via β_3 -adrenoceptors on brown lipocytes (\rightarrow p. 222).

NE in the synaptic cleft **is deactivated by** $(\rightarrow A6 \ a - d)$:

- diffusion of NE from the synaptic cleft into the blood:
- extraneuronal NE uptake (in the heart, glands, smooth muscles, glia, and liver), and subsequent intracellular degradation of NE by catecholamine-O-methyltransferase (COMT) and monoamine oxidase (MAO);
- active re-uptake of NE (70%) by the presynaptic nerve terminal. Some of the absorbed NE enters intracellular vesicles (→A3) and is reused, and some is inactivated by MAO;
- ♦ stimulation of presynaptic α_2 -adrenoceptors (*autoreceptors*; → **A 6d, 7**) by NE in the synaptic cleft, which inhibits the further release of NE.

Presynaptic α_2 -adrenoceptors can also be found on cholinergic nerve endings, e.g., in the gastrointestinal tract (motility \downarrow) and cardiac atrium (negative dromotropic effect), whereas presynaptic M-cholinoceptors are present on noradrenergic nerve terminals. Their mutual interaction permits a certain degree of peripheral ANS regulation.

Adrenal Medulla

After stimulation of preganglionic sympathetic nerve fibers (cholinergic transmission; → p. 81), 95% of all cells in the adrenal medulla secrete the endocrine hormone **epinephrine** (E) into the blood by exocytosis, and another 5% release norepinephrine (NE). Compared to noradrenergic neurons (see above), *NE synthesis* in the adrenal medulla is similar, but most of the NE leaves the vesicle and is enzymatically metabolized into E in the cytoplasm. Special vesicles called *chromaffin bodies* then actively store E and get ready to release it and co-transmitters (enkephalin, neuropeptide Y) by exocytosis.

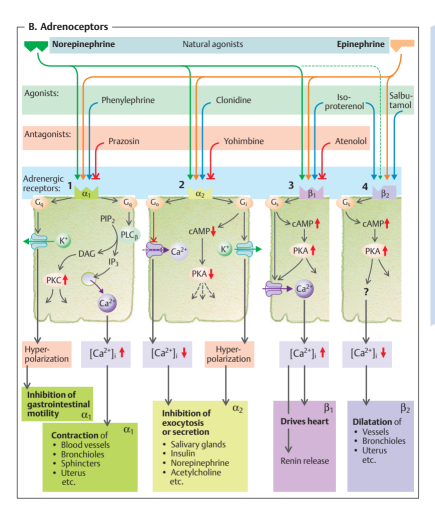
In alarm reactions, secretion of E (and some NE) from the adrenal medulla increases substantially in response to physical and mental or emotional stress. Therefore, cells not sympathetically innervated are also activated in such stress reactions. E also increases neuronal NE release via presynaptic β2-adrenoceptors $(\rightarrow A2)$. Epinephrine secretion from the adrenal medulla (mediated by increased sympathetic activity) is stimulated by certain triggers, e.g., physical work, cold, heat, anxiety, anger (stress), pain, oxygen deficiency, or a drop in blood pressure. In severe hypoglycemia (<30 mg/dL), for example, the plasma epinephrine concentration can increase by as much as 20-fold, while the norepinephrine concentration increases by a factor of only 2.5, resulting in a corresponding rise in the E/NE ratio.

The main task of epinephrine is to mobilize stored chemical energy, e.g., through *lipolysis* and *glycogenolysis*. Epinephrine enhances the uptake of glucose into skeletal muscle (\rightarrow p. 282) and activates enzymes that accelerate glycolysis and lactate formation (\rightarrow p. 72ff.). To enhance the blood flow in the muscles involved, the body increases the cardiac output while curbing gastrointestinal blood flow and activity (\rightarrow p. 75 A). Adrenal epinephrine and neuronal NE begin to stimulate the secretion of hormones responsible for replenishing the depleted energy reserves (e.g., ACTH; \rightarrow p. 297 A) while the alarm reaction is still in process.

Non-cholinergic, Non-adrenergic Transmitters

In humans, gastrin-releasing peptide (GRP) and vasoactive intestinal peptide (VIP) serve as co-transmitters in *preganglionic* sympathetic fibers; neuropeptide Y (NPY) and somatostatin (SIH) are the ones involved in *postganglionic* fibers. Postganglionic parasympathetic fibers utilize the peptides enkephalin, substance P (SP) and/or NPY as co-transmitters.

Modulation of postsynaptic neurons seems to be the primary goal of preganglionic peptide secretion. There is substantial evidence demonstrating that ATP (adenosine triphosphate), NPY and VIP also function as independent neu-



rotransmitters in the autonomic nervous system. VIP and acetylcholine often occur jointly (but in separate vesicles) in the parasympathetic fibers of blood vessels, exocrine glands,

thetic fibers of blood vessels, exocrine glands, and sweat glands. Within the gastrointestinal tract, VIP (along with nitric oxide) induces the slackening of the circular muscle layer and sphincter muscles and (with the co-transmitters dynorphin and galanin) enhances intesti-

nal secretion. **Nitric oxide (NO)** is liberated from nitrergic neurons (\rightarrow p. 278)

Composition and Function of Blood

The blood volume of an adult correlates with his or her (fat-free) body mass and amounts to ca. 4-4.5 L in women (\mathfrak{P}) and 4.5-5 L in men of 70 kg BW (δ : \rightarrow table). The functions of blood include the transport of various molecules (O2. CO2, nutrients, metabolites, vitamins, electrolytes, etc.), heat (regulation of body temperature) and transmission of signals (hormones) as well as buffering and immune defense. The blood consists of a fluid (plasma) formed elements: Red blood cells (RBCs) transport O2 and play an important role in pH regulation. White blood cells (WBCs) can be divided into eosinophilic and neutrophilic. granulocytes, monocytes, and lymphocytes. Neutrophils play a role in nonspecific immune defense, whereas monocytes and lymphocytes participate in specific immune responses. Platelets (thrombocvtes) are needed for hemostasis. Hematocrit (Hct) is the volume ratio of red cells to whole blood ($\rightarrow \mathbf{C}$ and Table). Plasma is the fluid portion of the blood in which electrolytes, nutrients, metabolites, vitamins, hormones, gases, and proteins are dissolved.

Plasma proteins (→ Table) are involved in humoral immune defense and maintain oncotic pressure, which helps to keep the blood volume constant. By binding to plasma proteins, compounds insoluble in water can be transported in blood, and many substances

Blood volume in liters relative to body weight (BW) \circlearrowleft 0.041 \times BW (kg) + 1.53, \circlearrowleft 0.047 \times BW (kg) + 0.86

Hematocrit (cell volume/ blood volume): ♂ 0.40-0.54 Females: 0.37-0.47

Erythrocytes ($10^{12}/L$ of blood = $10^{6}/\mu L$ of blood): $3 \cdot 4.6 - 5.9 \quad 9 \cdot 4.2 - 5.4$

Hemoglobin (g/L of blood): ♂ 140–180 ♀ 120–160

MCH, MCV, MCHC—mean corpuscular (MC), hemoglobin (Hb), MC volume, MC Hb concentration \rightarrow **C**

Leukocytes (10^9 /L of blood = 10^3 / μ L of blood): 3–11 (64% granulocytes, 31% lymphocytes, 6% monocytes)

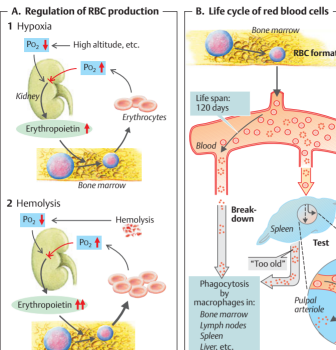
Platelets (10⁹/L of blood = 10³/ μ L of blood): \$\displant 170-360 \quad \text{\$\cong 180-400}\$

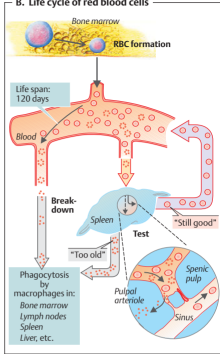
Plasma proteins (g/L of serum): 66–85 (including 55–64% albumin)

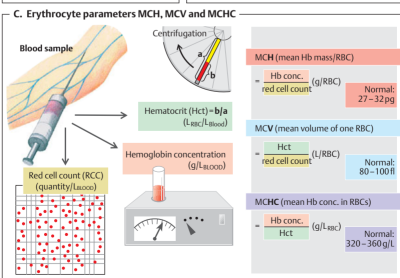
(e.g., heme) can be protected from breakdown and renal excretion. The binding of small molecules to plasma proteins reduces their osmotic efficacy. Many plasma proteins are involved in blood clotting and fibrinolysis. Serum forms when fibrinogen separates from plasma in the process of blood clotting.

The formation of blood cells occurs in the red bone marrow of flat bone in adults and in the spleen and liver of the fetus. Hematopoietic tissues contain pluripotent stem cells which, with the aid of hematopoietic growth factors (see below), develop into myeloid, erythroid and lymphoid precursor cells. Since pluripotent stem cells are autoreproductive, their existence is ensured throughout life. In lymphocyte development, lymphocytes arising from lymphoid precursor cells first undergo special differentiation (in the thymus or bone marrow) and are later formed in the spleen and lymph nodes as well as in the bone marrow. All other precursor cells are produced by myelocytopoiesis, that is, the entire process of proliferation, maturation, and release into the bloodstream occurs in the bone marrow. Two hormones, erythropoietin and thrombopoietin, are involved in myelopoiesis. Thrombopoietin mainly in the liver) promotes the maturation and development of megakaryocytes from which the platelets are split off. A number of other growth factors affect blood cell formation in bone marrow via paracrine mechanisms.

Erythropoietin promotes the maturation and **proliferation of red blood cells.** It is secreted by the liver in the fetus, and chiefly by the kidney (ca. 90%) in postnatal life. In response to an oxygen deficiency (due to high altitudes, hemolysis, etc.; \rightarrow A), erythropoietin secretion increases, larger numbers of red blood cells are produced, and the fraction of reticulocytes (voung erythrocytes) in the blood rises. The life span of a red blood cell is around 120 days. Red blood cells regularly exit from arterioles in the splenic pulp and travel through small pores to enter the splenic sinus $(\rightarrow B)$, where old red blood cells are sorted out and destroyed (hemolysis). Macrophages in the spleen, liver, bone marrow, etc. engulf and break down the cell fragments. Heme, the iron-containing group of hemoglobin (Hb) released during hemolysis, is broken down into bilirubin







Iron Metabolism and Erythropoiesis

Roughly $\frac{2}{3}$ of the body's iron pool (ca. 2 g in women and 5 g in men) is bound to hemoglobin (Hb). About 1/4 exists as stored iron (ferritin, hemosiderin), the rest as functional iron (myoglobin, iron-containing enzymes). Iron losses from the body amount to about 1 mg/day in men and up to 2 mg/day in women due to menstruation, birth, and pregnancy, Iron absorption occurs mainly in the duodenum and varies according to need. The absorption of iron supplied by the diet usually amounts to about 3 to 15% in healthy individuals, but can increase to over 25% in individuals with iron deficiency (→A1). A minimum daily iron intake of at least 10-20 mg/day is therefore recommended (women > children > men).

Iron absorption (\rightarrow **A2**). Fe(II) supplied by the diet (hemoglobin, myoglobin found chiefly in meat and fish) is absorbed relatively efficiently as a heme-Fe(II) upon protein cleavage. With the aid of *heme oxygenase*, Fe in mucosal cells cleaves from heme and oxidizes to Fe(III). The triferric form either remains in the mucosa as a ferritin-Fe(III) complex and returns to the lumen during cell turnover or enters the bloodstream. Non-heme-Fe can only be absorbed as Fe2+. Therefore, non-heme Fe(III) must first be reduced to Fe2+ by ferrireductase (FR: \rightarrow A2) and ascorbate on the surface of the luminal mucosa (\rightarrow A2). Fe²⁺ is probably absorbed through secondary active transport via an Fe²⁺-H⁺ symport carrier (DCT1) (competition with Mn²⁺, Co²⁺, Cd²⁺, etc.). A low chymous pH is important since it (a) increases the H+ gradient that drives Fe2+ via DCT1 into the cell and (b) frees dietary iron from complexes. The absorption of iron into the bloodstream is regulated by the intestinal mucosa. When an iron deficiency exists, aconitase (an iron-regulating protein) in the cytosol binds with ferritin-mRNA, thereby inhibiting mucosal ferritin translation. As a result, larger quantities of absorbed Fe(II) can enter the bloodstream. Fe(II) in the blood is oxidized to Fe(III) by ceruloplasmin (and copper). It then binds to apotransferrin, a protein responsible for iron transport in plasma ($\rightarrow A2$, 3). Transferrin (=apotransferrin loaded with 2 Fe(III)), is taken up by endocytosis into erythroblasts and cells of the liver, placenta, etc. with the aid of

transferrin receptors. Once iron has been released to the target cells, apotransferrin again becomes available for uptake of iron from the intestine and macrophages (see below).

Iron storage and recycling $(\rightarrow A3)$. Ferritin, one of the chief forms in which iron is stored in the body, occurs mainly in the intestinal mucosa, liver, bone marrow, red blood cells, and plasma. It contains binding pockets for up to 4500 Fe³⁺ ions and provides rapidly available stores of iron (ca. 600 mg), whereas iron mobilization from hemosiderin is much slower (250 mg Fe in macrophages of the liver and bone marrow). Hb-Fe and heme-Fe released from malformed erythroblasts (so-called inefficient erythropoiesis) and hemolyzed red blood cells bind to haptoglobin hemopexin, respectively. They are then engulfed by macrophages in the bone marrow or in the liver and spleen, respectively, resulting in 97% iron recycling (\rightarrow A3).

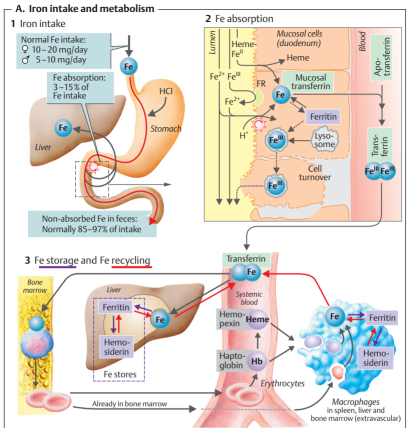
An **iron deficiency** inhibits Hb synthesis, leading to hypochromic microcytic anemia: MCH < 26 pg, MCV < 70 fL, Hb < 110 g/L. The primary causes are:

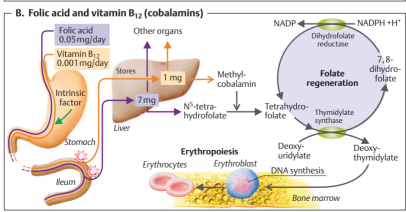
- blood loss (most common cause); 0.5 mg Fe are lost with each mL of blood;
- insufficient iron intake or absorption;
- increased iron requirement due to growth, pregnancy, breast-feeding, etc.;
- decreased iron recycling (due to chronic infection);
- apotransferrin defect (rare cause).

Iron overload most commonly damages the liver, pancreas and myocardium (hemochromatosis). If the iron supply bypasses the intestinal tract (iron injection), the transferrin capacity can be exceeded and the resulting quantities of free iron can induce iron poisoning.

 B_{12} vitamin (cobalamins) and folic acid are also required for erythropoiesis (\rightarrow B). Deficiencies lead to hyperchromic anemia (decreased RCC, increased MCH). The main causes are lack of intrinsic factor (required for cobalamin resorption) and decreased folic acid absorption due to malabsorption (see also p.260) or an extremely unbalanced diet. Because of the large stores available, decreased cobalamin absorption does not lead to symptoms of deficiency until many years later, whereas folic acid deficiency leads to symptoms within a few months.

91





Flow Properties of Blood

The **viscosity** (η) of blood is higher than that of plasma due to its erythrocyte (RBC) content. Viscosity $(\eta) = 1/\text{fluidity} = \text{shearing force}$ (τ) /shearing action (γ) [Pa·s]. The viscosity of blood rises with increasing hematocrit and decreasing flow velocity. Erythrocytes lack the major organelles and, therefore, are highly deformable. Because of the low viscosity of their contents, the liquid film-like characteristics of their membrane, and their high surface/ volume ratio, the blood behaves more like an emulsion than a cell suspension, especially when it flows rapidly. The viscosity of flowing blood (η_{blood}) passing through small arteries (\varnothing 20 µm) is about 4 relative units (RU). This is twice as high as the viscosity of plasma (nplasma = 2 RU: water: 1 RU = $0.7 \text{ mPa} \cdot \text{s}$ at $37 \,^{\circ}\text{C}$).

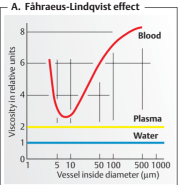
Because they are highly deformable, normal RBCs normally have no problem passing through capillaries or pores in the splenic vessels (see p. 89 B), although their diameter (Ø $< 5 \,\mu\text{m}$) is smaller than that of freely mobile RBCs (7 µm). Although the slowness of flow in small vessels causes the blood viscosity to increase, this is partially compensated for $(\eta_{blood}\downarrow)$ by the passage of red cells in single file through the center of small vessels (diame $ter < 300 \mu m$) due to the Fåhraeus-Lindqvist **effect** $(\rightarrow A)$. Blood viscosity is only slightly higher than plasma viscosity in arterioles $(\emptyset \approx 7 \,\mu\text{m})$, but rises again in capillaries $(\emptyset \approx 4 \,\mu\text{m})$. A critical increase in blood viscosity can occur a) if blood flow becomes too sluggish and/or b) if the fluidity of red cells decreases due to hyperosmolality (resulting in crenation), cell inclusion, hemoglobin malformation (e.g., sickle-cell anemia), changes in the cell membrane (e.g., in old red cells), and so forth. Under such circumstances, the RBCs undergo aggregation (rouleaux formation), increasing the blood viscosity tremendously (up to 1000 RU). This can quickly lead to the cessation of blood flow in small vessels (\rightarrow p. 218).

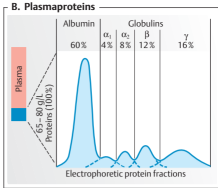
Plasma, Ion Distribution

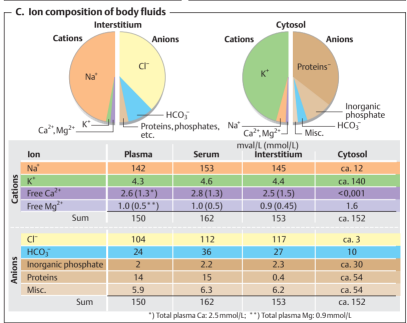
Plasma is obtained by preventing the blood from clotting and extracting the formed elements by centrifugation (\rightarrow p. 89 C). High molecular weight proteins $(\rightarrow B)$ as well as ions and non-charged substances with low molecular weights are dissolved in plasma. The sum of the concentrations of these particles yields a *plasma osmolality* of 290 mOsm/kgH₂O $(\rightarrow$ pp. 164, 377). The most abundant cation in plasma is Na⁺, and the most abundant anions are Cl⁻ and HCO₃⁻. Although plasma proteins carry a number of anionic net charges $(\rightarrow C)$, their osmotic efficacy is smaller because the number of particles, not the ionic valency, is the determining factor.

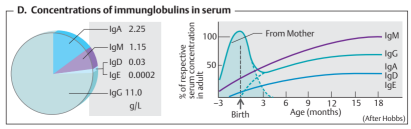
The fraction of proteins able to leave the blood vessels is small and varies from one organ to another. Capillaries in the liver, for example, are much more permeable to proteins than those in the brain. The composition of *interstitial fluid* therefore differs significantly from that of plasma, especially with respect to protein content $(\rightarrow \mathbf{C})$. A completely different composition is found in the *cytosol*, where K⁺ is the prevailing cation, and where phosphates, proteins and other organic anions comprise the major fraction of anions $(\rightarrow \mathbf{C})$. These fractions vary depending on cell type.

Sixty percent of all plasma protein $(\rightarrow B)$ is albumin (35-46 g/L). Albumin serves as a vehicle for a number of substances in the blood. They are the main cause of colloidal osmotic pressure or, rather, oncotic pressure (\rightarrow pp. 208. 378), and they provide a protein reserve in times of protein deficiency. The α_1 , α_2 and β globulins mainly serve to transport lipids (apolipoproteins), hemoglobin (haptoglobin), iron (apotransferrin), cortisol (transcortin), cobalamins (transcobalamin). plasma factors for coagulation and fibrinolysis are also proteins. Most plasma immunoglobulins (Ig, \rightarrow **D**) belong to the group of γ globulins and serve as defense proteins (antibodies). IgG, the most abundant immunoglobulin (7-15 g/L), can cross the placental barrier (maternofetal transmission; \rightarrow **D**). Each Ig consists of two group-specific, heavy protein chains (IgG: γ chain, IgA: α chain, IgM: μ chain, IgD: δ chain, IgE: ϵ chain) and two light protein chains (λ or κ chain) linked by disulfide bonds to yield a characteristic Y-shaped configuration (see p. 95 A).









Immune System

Fundamental Principles

The body has nonspecific (innate) immune defenses linked with specific (acquired) immune defenses that counteract bacteria. viruses, fungi, parasites and foreign (non-self) macromolecules. They all function as antigens. i.e., substances that stimulate the specific immune system, resulting in the activation of antigen-specific T lymphocytes (T cells) and B lymphocytes (B cells). In the process, Blymphocytes differentiate into plasma cells that secrete antigen-specific antibodies (immuno**globulins, Iq**) (\rightarrow **C**). Ig function to neutralize and opsonize antigens and to activate the complement system (\rightarrow p. 96). These mechanisms ensure that the respective antigen is specifically recognized, then eliminated by relatively nonspecific means. Some of the T and B cells have an immunologic memory.

Precursor lymphocytes without an antigenbinding receptor are preprocessed within the thymus (T) or bone marrow (B). These organs produce up to 108 monospecific T or B cells, each of which is directed against a specific antigen. Naive T and B cells which have not previously encountered antigen circulate through the body (blood → peripheral lymphatic tissue \rightarrow lymph \rightarrow blood) and undergo clonal expansion and selection after contact with its specific antigen (usually in lymphatic tissue). The lymphocyte then begins to divide rapidly, producing numerous monospecific daughter cells. The progeny differentiates into plasma cells or "armed" T cells that initiate the elimination of the antigen.

Clonal deletion is a mechanism for eliminating lymphocytes with receptors directed against autologous (self) tissue. After first contact with their specific self-antigen, these lymphocytes are eliminated during the early stages of development in the thymus or bone marrow. Clonal deletion results in central immunologic tolerance. The ability of the immune system to distinguish between endogenous and foreign antigens is called self/nonself recognition. This occurs around the time of birth. All substances encountered by that time are recognized as endogenous (self); others are identified as foreign (nonself). The inability to distinguish self from nonself results in autoimmune disease.

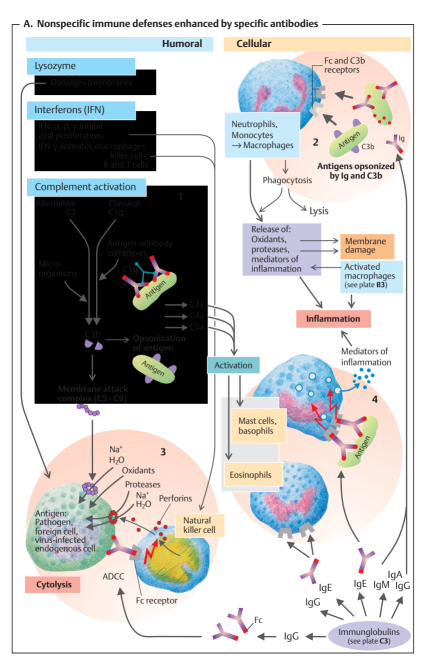
At first contact with a virus (e.g., measles virus), the nonspecific immune system usually cannot prevent the viral proliferation and the development of the measles. The specific immune system, with its T-killer cells (\rightarrow **B2**) and Ig (first IgM, then IgG: \rightarrow C3), responds slowly: primary immune response or sensitization. Once activated, it can eliminate the pathogen, i.e., the individual recovers from the measles. Secondary immune response: When infected a second time, specific IgG is produced much more rapidly. The virus is quickly eliminated, and the disease does not develop a second time. This type of protection against infectious disease is called immunity. It can be achieved by vaccinating the individual with a specific antigen (active immunization). Passive immunization can be achieved by administering ready-made Ig (immune serum).

Nonspecific Immunity

Lysozyme and complement factors dissolved in plasma (\rightarrow A1) as well as natural killer cells (NK cells) and phagocytes, especially neutrophils and macrophages that arise from monocytes that migrate into the tissues $(\rightarrow A2)$ play an important role in nonspecific immunity. Neutrophils, monocytes, and eosinophils circulate throughout the body. They have chemokine receptors (e.g., CXCR1 and 2 for IL-8) and are attracted by various chemokines (e.g., IL-8) to the sites where microorganisms have invaded (chemotaxis). These cells are able to migrate. With the aid of selectins, they dock onto the endothelium (margination), penetrate the endothelium (diapedesis), and engulf and damage the microorganism with the aid of lysozyme, oxidants such as H2O2, oxygen radicals $(O_2^-, OH_1, {}^1O_2)$, and *nitric oxide* (NO). This is followed by digestion (lysis) of the microorganism with the aid of lysosomal enzymes. If the antigen (parasitic worm, etc.) is too large for digestion, other substances involved in nonspecific immunity (e.g., proteases and cvtotoxic proteins) are also exocytosed by these cells.

Reducing enzymes such as catalase and superoxide dismutase usually keep the oxidant concentration low. This is often discontinued, especially when macrophages are activated (—below and B3), to fully exploit the bactericidal effect of the oxidants. However,

▶



the resulting inflammation (\rightarrow A2, 4) also damages cells involved in nonspecific defense and, in some cases, even other endogenous cells.

Opsonization (\rightarrow A1, 2) involves the binding of opsonins, e.g., IgG or complement factor C3b, to specific domains of an antigen, thereby enhancing phagocytosis. It is the only way to make bacteria with a polysaccharide capsule phagocytable. The phagocytes have receptors on their surface for the (antigen-independent) Fc segment of IgG as well as for C3b. Thus, the antigen-bound IgG and C3b bind to their respective receptors, thereby linking the rather unspecific process of phagocytosis with the specific immune defense system, Carbohydrate-binding proteins (lectins) of plasma. called collectins (e.g. mannose-binding protein), which dock onto microbial cell walls, also acts as unspecific opsonins.

The **complement cascade** is activated by antigens opsonized by Ig (classical pathway) as well as by non-opsonophilic antigens (alternative pathway) (\rightarrow A1). Complement components C3a, C4a and C5a activate basophils and eosinophils (\rightarrow A4). Complement components C5 – C9 generate the *membrane-attack complex* (MAC), which perforates and kills (Gram-negative) bacteria by **cytolysis** (\rightarrow A3). This form of defense is assisted by **lysozyme** (= muramidase), an enzyme that breaks down murein-containing bacterial cell walls. It occurs in granulocytes, plasma, lymph, and secretions.

Natural killer (NK) cells are large, granular lymphocytes specialized in nonspecific defense against viruses, mycobacteria, tumor cells etc. They recognize infected cells and tumor cells on "foreign surfaces" and dock via their Fc receptors on IgG-opsonized surface antigens (antibody-dependent cell-mediated cytotoxicity, ADCC; \rightarrow A3). Perforins exocytosed by NK cells form pores in target cell walls, thereby allowing their subsequent lysis (cytolysis). This not only makes the virus unable to proliferate (enzyme apparatus of the cell), but also makes it (and other intracellular pathogens) subject to attack from other defense mechanisms.

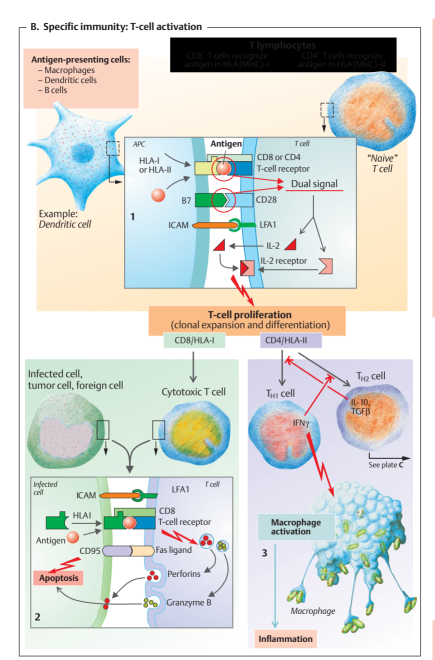
Various **interferons (IFNs)** stimulate NK cell activity: IFN- α , IFN- β and, to a lesser degree, IFN- γ . IFN- α and IFN- β are released mainly from leukocytes and fibro-

blasts, while IFN- γ is liberated from activated T cells and NK cells. Virus-infected cells release large quantities of IFNs, resulting in heightened viral resistance in non-virus-infected cells. **Defensins** are cytotoxic peptides released by phagocytes. They can exert unspecific cytotoxic effects on pathogens resistant to NK cells (e.g., by forming ion channels in the target cell membrane).

Macrophages arise from monocytes that migrate into the tissues. Some macrophages are freely mobile (free macrophages), whereas others (fixed macrophages) remain restricted to a certain area, such as the hepatic sinus (Kupffer cells), the pulmonary alveoli, the intestinal serosa, the splenic sinus, the lymph nodes, the skin (Langerhans cells), the synovia (synovial A cells), the brain (microglia), or the endothelium (e.g., in the renal glomeruli). The mononuclear phagocytic system (MPS) is the collective term for the circulating monocytes in the blood and macrophages in the tissues. Macrophages recognize relatively unspecific carbohydrate components on the surface of bacteria and ingest them by phagocytosis. The macrophages have to be activated if the pathogens survive within the phagosomes (→ below and **B3**).

Specific Immunity: Cell-Mediated Immune Responses

Since specific cell-mediated immune responses through "armed" T effector cells need a few days to become effective, this is called delayed-type immune response. It requires the participation of professional antigen-presenting cells (APCs): dendritic cells, macrophages and B cells. APCs process and present antigenic peptides to the T cells in association with MHC-I or MHC-II proteins, thereby delivering the co-stimulatory signal required for activation of naive T cells. (The gene loci for these proteins are the class I (MHC-I) and class II (MHC-II) major histocompatibility complexes (MHC)), **HLA** (human leukocyte antigen) is the term for MHC proteins in humans. Virusinfected dendritic cells, which are mainly located in lymphatic tissue, most commonly serve as APCs. Such HLA-restricted antigen presentation $(\rightarrow B1)$ involves the insertion of an antigen in the binding pocket of an HLA protein. An ICAM (intercellular adhesion molecule) on the surface of the APC then binds to



98

LFA1 (lymphocyte function-associated antigen 1) on the T cell membrane. When a T cell specific for the antigen in question docks onto the complex, the bond is strengthened and the *APC dual signal* stimulates the activation and clonal selection of the T cell (\rightarrow **B1**).

The APC dual signal consists of 1) recognition of the antigen (class I or class II HLA-restricted antigen) by the *T cell receptor* and its *co-receptor* and 2) a costimulatory signal, that is, the binding of the B7 protein (on the APC) with the CD28 protein on the T cell (\rightarrow B1). CD8 molecules on T cytotoxic cells (Tc cells = T-killer cells) and CD4 molecules on T helper cells (T_H cells) function as the co-receptors. When antigen binding occurs without co-stimulation (e.g., in the liver, where there are no APCs), the lymphocyte is inactivated, i.e., it becomes anergic, and peripheral immunologic tolerance develops.

The T cell can receive APC dual signals from infected macrophages or B cells, provided their receptors have bound the antigen in question (e.g., insect or snake venom or allergens). The APC dual signal induces the T cell to express *interleukin-2* (IL-2) and to bind the respective *IL-2 receptor* (\rightarrow B1). IL-2 is the actual *signal for clonal expansion* of these monospecific T cells. It functions through autocrine and paracrine mechanisms. Potent *immunosuppression*, e.g., for organ transplantation, can be achieved with IL-2 inhibitors like *cyclosporin A*.

During clonal expansion, the T cells differentiate into three "armed" subtypes, i.e., T cytotoxic cells (Tc cells or T killer cells) and T helper cells type 1 (TH1 cells) and type 2 (TH2 cells). These cells no longer require costimulation and express a different type of adhesion molecule (VLA-4 instead of L-selectins) by which they now dock onto the endothelium of inflamed tissues (rather than to lymphatic tissue like their naïve precursors).

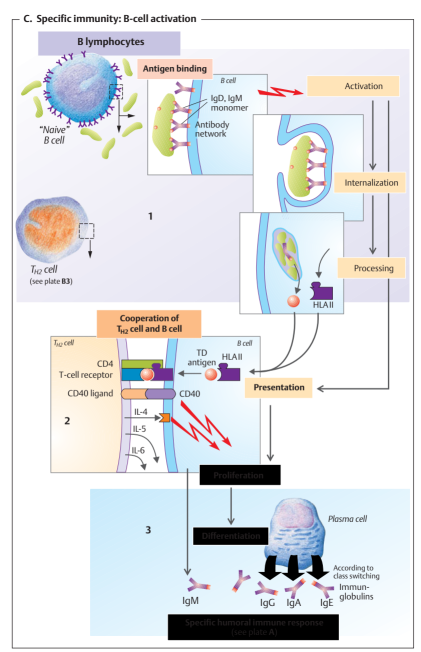
T killer cells develop from naive CD8-containing (CD8⁺) T cells after HLA-I-restricted antigen presentation (\rightarrow **B2**). **Endogenous antigen presentation** occurs when the HLA-I protein takes up the antigen (virus, cytosolic protein) from the cytosol of the APC, which is usually the case. With its CD8-associated T-cell receptor, the T killer cell is able to recognize HLA-I-restricted antigens on (virus) infected endogenous cells and tumor cells as well as on cells of transplanted organs. It subsequently drives the cells into **apoptosis** (programmed cell death) or **necrosis**. Binding of the *Fas lig-*

and to CD95 (=Fas) plays a role, as does granzyme B (protease), which enters the cell through pores created by exocytosed *perforins* (\rightarrow B2).

Once HLA-II-restricted presentation (\rightarrow B1) of antigens from intracellular vesicles (e.g., phagocytosed bacteria or viral envelope proteins = exogenous antigen presentation) has occurred, naive CD4⁺ T cells transform into immature T helper cells (T_{H0}), which differentiate into T_{H1} or T_{H2} cells. T_{H1} cells induce inflammatory responses and promote the activation of macrophages with the aid of IFN- γ (\rightarrow B3), while T_{H2} cells are required for B-cell activation (\rightarrow C2). T_{H1} and T_{H2} cells mutually suppress each other, so only one of the two types will predominate in any given cell-mediated immune response (\rightarrow B3).

Specific Immunity: Humoral Immune Responses

Humoral immunity arise from B cells $(\rightarrow C1)$. Numerous IgD and IgM monomers anchored onto the B-cell surface bind with the respective antigen (dissolved IgM occurs in pentameric form). A resulting network of antigenbound Ig leads to internalization and processing of the antigen-antibody complex in B cells. However, B-cell activation requires a second signal, which can come directly from a thymus-independent (TI) antigen (e.g., bacterial polysaccharide) or indirectly from a T_{H2} cell in the case of a thymus-dependent (TD) antigen. In the latter case, the B cell presents the HLA-II-restricted TD antigen to the T_{H2} cell $(\rightarrow$ C2). If the CD4-associated T-cell receptor (TCR) of the T_{H2} cell recognizes the antigen, CD40 ligands are expressed on the T_{H2} surface (CD40 ligands bind with CD40 proteins on B cells) and IL-4 is secreted. The CD40 ligand and IL-4 (later also IL-5 and IL-6) stimulate the B cell to undergo clonal selection, IgM secretion, and differentiation into plasma cells (\rightarrow C3). Before differentiation, class switching can occur, i.e., a different type of Ig heavy chain $(\rightarrow p.92)$ can be expressed by altered DNA splicing (\rightarrow p. 8f.). In this manner, IgM is converted into IgA, IgG or IgE (\rightarrow p. 92). All Ig types arising from a given B-cell clone remain monospecific for the same antigen. The plasma cells formed after class switching produce only a single type of Ig.



Hypersensitivity Reactions (Allergies)

Allergy is a specific, exaggerated immune response to a (usually harmless) foreign substance or antigen (\rightarrow p. 94ff.). Allergens are antigens that induce allergies. Small molecules conjugated to endogenous proteins can also have antigenic effects. In this case, they are referred to as incomplete antigens or haptens. The heightened immune response to secondary antigen contact ($\rightarrow p.94ff$.) normally has a protective effect. In allergies, however, the first contact with an antigen induces sensitization (allergization), and subsequent exposure leads to the destruction of healthy cells and intact tissue. This can also result in damage to endogenous proteins and autoantibody production. *Inflammatory reactions* are the main causes of damage.

Types of hypersensitivity reactions: Type I reactions are common. On first contact, the allergen internalized by B cells is presented to T_{H2} cells. The B cell then proliferates and differentiates into plasma cells (see p. 98), which release immunoglobulin E (IqE). The Fc fragment of IgE binds to mast cells and basophils. On subsequent contact, the antigens bind to the already available IgE-linked mast cells $(\rightarrow A)$. Due to the rapid release of mostly vasoactive mediators of inflammation such as histamine. leukotrienes and platelet-activating factor (PAF), an immediate reaction (anaphylaxis) occurs within seconds or minutes: immediate type hypersensitivity. This is the mechanism by which allergens breathed into the lungs trigger hay fever and asthma attacks. The vasodilatory effect of a generalized type I reaction can lead to anaphylactic shock (see p. 218).

In **type II** reactions, the immune system mainly attacks *cells* with antigenic properties. This can be attributable to the transfusion of the erythrocytes of the wrong blood group or the binding of *haptens* (e.g., medications) to endogenous cells. The binding of haptens to platelets can, for example, result in throm-bocytopenia.

Type III reactions are caused by antigen-antibody complexes. If more antigen than antibody is available, *soluble antigen-antibody complexes* circulate in blood for a long time (\rightarrow B) and settle mainly in the capillaries, making the capillary wall subject to attack

by the complement system. This leads to the development of **serum sickness** $(\rightarrow \mathbf{B})$, the main symptoms of which are joint pain and fever.

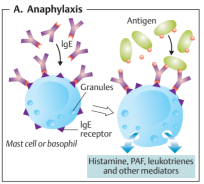
Type IV reactions are mainly mediated by T_{H1} cells, T_C cells, and macrophages. Since symptoms peak 2 to 4 days after antigen contact, this is called **delayed type hypersensitivity**. The main triggers are mycobacteria (e.g. Tbc), other foreign proteins, and haptens, such as medications and plant substances, such as poison ivy. Primary *transplant rejection* is also a type IV hypersensitivity reaction. *Contact dermatitis* is also a type IV reaction caused by various haptens (e.g., nickel in jewelry).

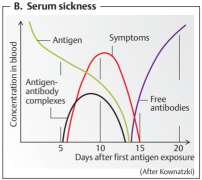
Blood Groups

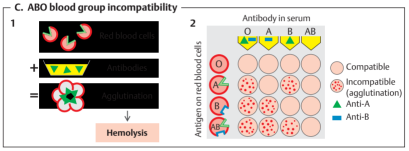
A person's blood group is determined by the type of antigen (certain glycolipids) present on the red blood cells (RBCs). In the **ABO system**, the antigens are A and B (\rightarrow **C**). In blood type A, antigen A (on RBC) and anti-B antibody (in serum) are present; in type B, B and anti-A are present; in type AB, A and B are present, no antibody; in type 0 (zero), no antigen but anti-A and anti-B are present.

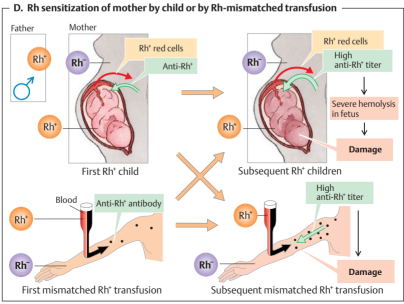
When giving a **blood transfusion**, it is important that the blood groups of donor and recipient match, i.e. that the RBCs of the donor (e.g. A) do not come in contact with the respective antibodies (e.g. anti-A) in the recipient. If the donor's blood is the wrong type, agglutination (cross-linking by IgM) and hemolysis (bursting) of the donor's RBCs will occur (\rightarrow C1). Donor and recipient blood types must therefore be determined and cross-matched (\rightarrow C2) prior to a blood transfusion. Since ABO antibodies belong to the IgM class, they usually do not cross the placenta.

In the **Rh system**, antibodies against rhesus antigens (C, D, E) on RBCs do not develop unless *prior sensitization* has occurred. D is by far the most antigenic. A person is Rh-positive (Rh+) when D is present on their RBCs (most people), and Rh-negative (Rh-) when D is absent. Anti-D antibodies belong to the IgG class of immunoglobulins, which are capable of crossing the placenta (\rightarrow p. 93 D). Rh- individuals can form anti-Rh+ (= anti-D) antibodies, e.g., after sensitization by a mismatched blood transfusion or of an Rh- mother by an Rh+ fetus. Subsequent exposure to the mismatched blood leads to a severe antigen-antibody reaction characterized by intravascular agqlutination and hemolysis (\rightarrow **D**).









Hemostasis

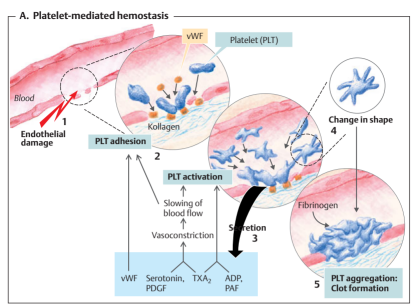
The hemostatic system stops bleeding. Thrombocytes (platelets), coagulation (or clotting) factors in plasma and vessel walls interact to seal leaks in blood vessels. The damaged vessel constricts (release of endothelin), and platelets aggregate at the site of puncture (and attract more platelets) to seal the leak by a platelet plug. The time required for sealing (ca. 2 to 4 min) is called the bleeding time. Subsequently, the coagulation system produces a fibrin meshwork. Due to covalent cross-linking of fibrin, it turns to a fibrin clot or thrombus that retracts afterwards, thus reinforcing the seal. Later recanalization of the vessel can be achieved by fibrinolysis.

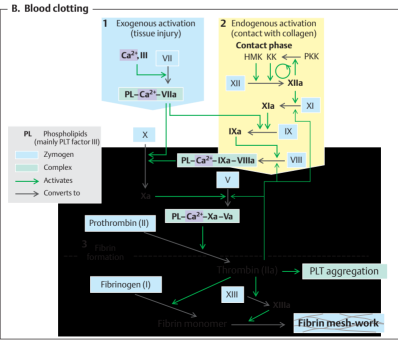
Platelets $(170-400 \cdot 10^3 \text{ per } \mu\text{L of blood};$ half-life ≈ 10 days) are small non-nucleated bodies that are pinched off from megakaryocytes in the bone marrow. When an endothelial injury occurs, platelets adhere to subendothelial collagen fibers ($\rightarrow A1$) bridged by von Willebrand's factor (vWF), which is formed by endothelial cells and circulates in the plasma complexed with factor VIII. Glycoprotein complex GP Ib/IX on the platelets are vWF receptors. This adhesion activates plate**lets** $(\rightarrow A2)$. They begin to release substances $(\rightarrow A3)$, some of which promote platelet adhesiveness (vWF). Others like serotonin, platelet-derived growth factor (PDGF) and thromboxane A2 (TXA2) promote vasoconstriction. Vasoconstriction and platelet contraction slow the blood flow. Mediators released by platelets enhance platelet activation and attract and activate more platelets: ADP, TXA2, platelet-activating factor (PAF). The shape of activated platelets change drastically ($\rightarrow A4$). Discoid platelets become spherical and exhibit pseudopodia that intertwine with those of other platelets. This platelet aggregation $(\rightarrow A5)$ is further enhanced by thrombin and stabilized by GP IIb/IIIa. Once a platelet changes its shape, GP IIb/IIIa is expressed on the platelet surface, leading to fibrinogen binding and platelet aggregation. GPIIb/IIIa also increases the adhesiveness of platelets, which makes it easier for them to stick to subendothelial fibronectin.

I	Fibrinogen I	Half-life (h):	96
II ^K	Prothrombin		72
Ш	Tissue thromboplastin		
IV	Ionized calcium (Ca ²⁺)		
V	Proaccelerin		20
VII^{K}	Proconvertin		5
VIII	Antihemophilic factor A		12
IX K	Antihemophilic factor B; planthromboplastin component Christmas factor		24
χк	Stuart–Prower factor		30
XI	Plasma thromboplastin ante	cedent	50
,	(PTA)	ccaciic	48
XII	Hageman factor		50
XIII	Fibrin-stabilizing factor (FSF)		250
-	Prekallikrein (PKK); Fletcher	factor	
-	High-molecular-weight kinin (HMK); Fitzgerald factor	ogen	

Several coagulation factors are involved in the clotting process. Except for Ca²⁺, they are proteins formed in the *liver* (\rightarrow **B** and Table). Factors labeled with a "K" in the table (as well as protein C and protein S, see below) are produced with vitamin K, an essential cofactor in posttranslational y-carboxylation of glutamate residues of the factors. These γ-carboxyglutamyl groups are chelators of Ca2+. They are required for Ca2+-mediated complex formation of factors on the surface of phospholipid layers (PL), particularly on the platelet membrane (platelet factor 3). Vitamin K is oxidized in the reaction and has to be re-reduced by liver epoxide reductase (vitamin K recycling). Ca²⁺ ions are required for several steps in the clotting process $(\rightarrow B)$. When added to blood samples in vitro, citrate, oxalate, and EDTA bind with Ca2+ ions, thereby preventing the blood from clotting. This effect is desirable when performing various blood tests.

Activation of blood clotting $(\rightarrow B)$. Most coagulation factors normally are not active, or *zymogenic*. Their activation requires a cascade of events (An "a" added to the factor number means "activated"). Thus, even small amounts of a trigger factor lead to rapid blood clotting. The trigger can be endogenous (within a vessel) or exogenous (external). **Endogenous activation** $(\rightarrow B2)$ occurs at an *endothelial defect*. XII is activated to XIIa by the contact with





negative charges of subendothelial collagen and sulfatide groups. This stimulates the conversion of prekallikrein (PKK) to kallikrein (KK), which enhances XII activation (positive feedback). Next, XIIa activates XI to XIa, which converts IX to IXa and, subsequently, VIII to VIIIa. Complexes formed by conjugation of IXa and VIIIa with Ca^{2+} on phospholipid (PL) layers activate X. **Exogenous activation** now merges with endogenous activation (\rightarrow **B1**). In relatively large injuries, *tissue thrombokinase* (factor III), present on of nonvascular cells, is exposed to the blood, resulting in activation of VII. VII forms complexes with Ca^{2+} and phospholipids. thereby activating X (and IX).

Fibrin formation (\rightarrow **B3**). After activation of X to Xa by endogenous and/or exogenous activation (the latter is faster). Xa activates V and conjugates with Va and Ca2+ on the surface of membranes. This complex, called prothrombinase, activates prothrombin (II) to thrombin (IIa). In the process, Ca2+ binds with phospholipids, and the N-terminal end of prothrombin splits off. The thrombin liberated in the process now activates (a) fibrinogen (I) to fibrin (Ia), (b) fibrin-stabilizing factor (XIII), and (c) V, VIII and XI (positive feedback). The single (monomeric) fibrin threads form a soluble meshwork (fibrins: "s" for soluble) which XIIIa ultimately stabilizes to insoluble fibrin (fibrini). XIIIa is a transamidase that links the side chains of the fibrin threads via covalent bonds.

Fibrinolysis and Thromboprotection

To prevent excessive clotting and occlusion of major blood vessels (thrombosis) and embolisms due to clot migration, fibrins is re-dissolved (fibrinolysis) and inhibitory factors are activated as soon as vessel repair is initiated.

Fibrinolysis is mediated by **plasmin** (→ C). Various factors in blood (plasma kallikrein, factor XIIa), tissues (tissue plasminogen activator, tPA, endothelial etc.), and urine (urokinase) activate *plasminogen* to plasmin. Streptokinase, staphylokinase and tPA are used therapeutically to activate plasminogen. This is useful for dissolving a fresh thrombus located, e.g., in a coronary artery. Fibrin is split into fibrinopeptides which inhibit thrombin

formation and polymerization of fibrin to prevent further clot formation. *Alpha*₂-antiplasmin is an endogenous inhibitor of fibrinolysis. *Tranexamic acid* is administered therapeutically for the same purpose.

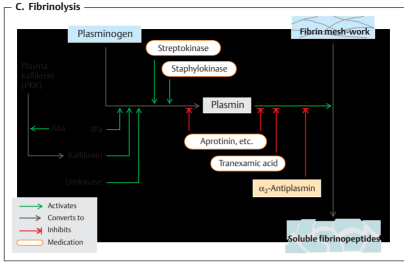
Thromboprotection. Antithrombin III, a serpin, is the most important thromboprotective plasma protein $(\rightarrow D)$.

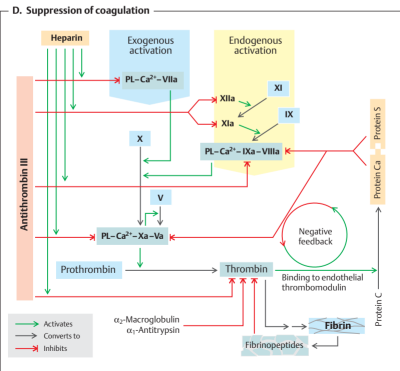
It inactivates the protease activity of thrombin and factors IXa, Xa, Xla and Xlla by forming complexes with them. This is enhanced by **heparin** and heparinlike endothelial glucosaminoglycans. Heparin is produced naturally by mast cells and granulocytes, and synthetic heparin is injected for therapeutic purposes.

The binding of thrombin with endothelial **thrombomodulin** provides further thromboprotection. Only in this form does thrombin have anticoagulant effects (\rightarrow **D**, negative feedback). Thrombomodulin activates protein C to Ca which, after binding to protein S, deactivates coagulation factors Va and VIIIa. The synthesis of proteins C and S is vitamin K-dependent. Other plasma proteins that inhibit thrombin are α_2 -macroglobulin and α_1 -antitrypsin (\rightarrow **D**). Endothelial cells secrete **tissue thromboplastin inhibitor**, a substance that inhibits exogenous activation of coagulation, and **prostacy-clin** (= prostaglandin 1_2), which inhibits platelet adhesion to the normal endothelium.

Anticoagulants are administered for thromboprotection in patients at risk of blood clotting. Injected heparin has immediate action. Oral coumarin derivatives (phenprocoumon, warfarin, acenocoumarol) are vitamin K antagonists that work by inhibiting liver epoxide reductase, which is necessary for vitamin D recycling. Therefore, these drugs do not take effect until the serum concentration of vitamin K-dependent coagulation factors has decreased. Cyclooxygenase inhibitors, such as aspirin (acetylsalicylic acid), inhibit platelet aggregation by blocking thromboxane A₂ (TXA2) synthesis (→ p. 269).

Hemorrhagic diatheses can have the following causes: a) Congenital deficiency of certain coagulation factors. Lack of VIII or IX, for example, leads to hemophilia A or B, respectively. b) Acquired deficiency of coagulation factors. The main causes are liver damage as well as vitamin K deficiency due to the destruction of vitamin K-producing intestinal flora or intestinal malabsorption. c) Increased consumption of coagulation factors, by disseminated intravascular coagulation. d) Platelet deficiency (thrombocytopenia) or platelet defect (thrombocytopathy). e) Certain vascular diseases, and f) excessive fibrinolysis.





Lung Function, Respiration

The lung is mainly responsible for respiration, but it also has *metabolic functions*, e.g. conversion of angiotensin I to angiotensin II (\rightarrow p. 184). In addition, the pulmonary circulation *buffers the blood volume* (\rightarrow p. 204) and *filters* out small blood clots from the venous circulation before they obstruct the arterial circulation (heart, brain!).

External respiration is the exchange of gases between the body and the environment. (Internal or tissue respiration involves nutrient oxidation, \rightarrow p. 228). Convection (bulk flow) is the means by which the body transports gases over long distances (\rightarrow p. 24) along with the flow of air and blood. Both flows are driven by a pressure difference. Diffusion is used to transport gases over short distances of a few µme.g., through cell membranes and other physiological barriers ($\rightarrow p$, 20ff.). The exchange of gas between the atmosphere and alveoli is called ventilation. Oxygen (O2) in the inspired air is convected to the alveoli before diffusing across the alveolar membrane into the bloodstream. It is then transported via the bloodstream to the tissues, where it diffuses from the blood into the cells and, ultimately, to the intracellular mitochondria. Carbon dioxide (CO₂) produced in the mitochondria returns to the lung by the same route.

The **total ventilation per unit time.** \dot{V}_T (also called minute volume) is the volume (V) of air inspired or expired per time. As the expiratory volume is usually measured, it is also abbreviated **V**E. (The dot means "per unit time"). At rest, the body maintains a $\dot{\mathbf{V}}_{E}$ of about 8 L/min, with a corresponding oxygen consumption rate (Vo₂) of about 0.3 L/min and a CO₂ elimination rate (Vco₂) of about 0.25 L/min. Thus, about 26 L of air have to be inspired and expired to supply 1 L of O₂ (respiratory equivalent = 26). The tidal volume (VT) is the volume of air moved in and out during one respiratory cycle. VE is the product of VT (ca. 0.5 L at rest) and respiration rate f (about 16/min at rest) (see p. 74 for values during physical work). Only around 5.6 L/min (at $f = 16 \text{ min}^{-1}$) of the \dot{V}_E of 8 L/min reaches the alveoli; this is known as alveolar ventilation (VA). The rest fills airways space not contributing to gas exchange (dead space venThe human body contains around 300 million **alveoli**, or thin-walled air sacs (ca. 0.3 mm in diameter) located on the terminal branches of the bronchial tree. They are surrounded by a dense network of **pulmonary capillaries** and have a total surface area of about 100 m². Because of this and the small air/blood diffusion distances of only a few μ m (\rightarrow p. 22, Eq. 1.7), sufficient quantities of O₂ can **diffuse** across the alveolar wall into the blood and CO₂ towards the alveolar space (\rightarrow p. 120ff.), even at a tenfold increased O₂ demand (\rightarrow p. 74). The oxygen-deficient "venous" blood of the pulmonary artery is thus oxygenated ("arterialized") and pumped back by the left heart to the periphery.

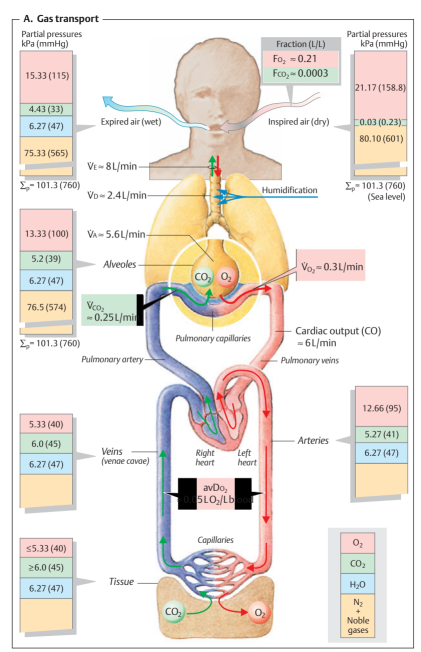
The **cardiac output (CO)** is the volume of blood pumped through the pulmonary and systemic circulation per unit time (5–6 L/min at rest). CO times the arterial-venous O_2 difference ($avDO_2$)—i.e., the difference between the arterial O_2 fraction in the aorta and in mixed venous blood of the right atrium (ca. $0.05 \, \text{L}$ of O_2 per L of blood)—gives the O_2 volume transported per unit time from the lungs to the periphery. At rest, it amounts to $(6 \times 0.05 =) 0.3 \, \text{L/min}$, a value matching that of \dot{V}_{O_2} (see above). Inversely, if \dot{V}_{O_2} and $avDO_2$ have been measured, CO can be calculated (**Fick's principle**):

 $CO = \dot{V}_{O_2} / avD_{O_2}$ [5.1] The *stroke volume* (**SV**) is obtained by dividing CO by the heart rate (pulse rate).

According to *Dalton's law*, the total pressure (P_{total}) of a **mixture of gases** is the sum of the **partial pressures (P)** of the individual gases. The volume **fraction** (F, in L/L; \rightarrow p. 376), of the individual gas relative to the total volume times P_{total} gives the partial pressure—in the case of O_2 , for example, $P_{O_2} = F_{O_2} \times P_{total}$. The atmospheric partial pressures in dry ambient air at sea level ($P_{total} = 101.3 \text{ kPa} = 760 \text{ mmHg}$) are: $F_{O_2} = 0.209$, $F_{CO_2} = 0.0004$, and $F_{N_2} + \text{noble}$ gases = 0.79 (\rightarrow **A**, top right).

If the mixture of gases is "wet", the **partial pressure** of water, PH₂O has to be subtracted from P_{total} (usually = atmospheric pressure). The other partial pressures will then be lower, since $P_X = F_X$ ($P_{total} - P_{H2}O$). When passing through the respiratory tract (37 °C), inspired air is fully saturated with water. As a result, $P_{H2}O$ rises to 6.27 kPa (47 mmHg), and P_{O2} drops 1.32 kPa lower than the dry atmospheric air (\rightarrow p. 112). The partial pressures in the alveoli, arteries, veins (mixed venous blood), tissues, and expiratory air (all "wet") are listed in **A**.

106



Mechanics of Breathing

Pressure differences between the alveoli and the environment are the *driving "forces"* for the exchange of gases that occurs during ventilation. **Alveolar pressure** ($PA = intrapulmonary pressure; <math>\rightarrow B$) must be lower than the barometric pressure (PB) during **inspiration** (breathing in), and higher during **expiration** (breathing out). If PB is defined as zero, the alveolar pressure is negative during inspiration and positive during expiration ($\rightarrow B$). These pressure differences are created through coordinated movement of the diaphragm and chest (thorax), resulting in an increase in lung volume (V_{pulm}) during inspiration and a decrease during expiration ($\rightarrow A1,2$).

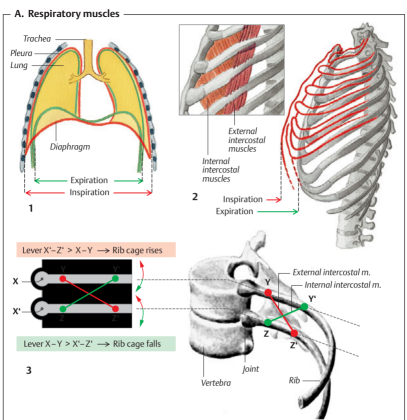
The inspiratory muscles consist of the diaphragm, scalene muscles, and external intercostal muscles. Their contraction lowers (flattens) the diaphragm and raises and expands the chest, thus expanding the lungs. Inspiration is therefore active. The external intercostal muscles and accessory respiratory muscles are activated for deep breathing. During expiration, the diaphragm and other inspiratory muscles relax, thereby raising the diaphragm and lowering and reducing the volume of the chest and lungs. Since this action occurs primarily due to the intrinsic elastic recoil of the lungs (\rightarrow p. 116), expiration is passive at rest. In deeper breathing, active mechanisms can also play a role in expiration: the internal intercostal muscles contract, and the diaphragm is pushed upward by abdominal pressure created by the muscles of the abdominal wall.

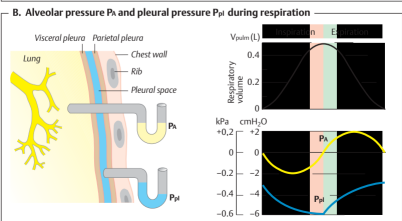
Two adjacent ribs are bound by internal and external intercostal muscle. Counteractivity of the muscles is due to variable leverage of the upper and lower rib (\rightarrow A3). The distance separating the insertion of the external intercostal muscle on the upper rib (Y) from the axis of rotation of the upper rib (X) is smaller than the distance separating the insertion of the muscles on the lower rib (Z') and the axis of rotation of the lower rib (X'). Therefore, X'–Z' is longer and a more powerful lever than X–Y. The chest generally rises when the external intercostal muscles contract, and lowers when the opposing internal intercostal muscles contract.

To exploit the motion of the diaphragm and chest for ventilation, the lungs must be able to follow this motion without being completely attached to the diaphragm and chest. This is achieved with the aid of the **pleura**, a thin fluid-covered sheet of cells that invests each lung (visceral pleura), thereby separating it from the adjacent organs, which are covered by the pleura as well (parietal pleura).

In its natural state, the lung tends to shrink due to its intrinsic elasticity and alveolar surface tension (\rightarrow p. 118). Since the fluid in the pleural space cannot expand, the lung sticks to the inner surface of the chest, resulting in suction (which still allows tangential movement of the two pleural sheets). Pleural pressure (Ppl) is then negative with respect to atmospheric pressure. Ppl. also called intrapleural (P_{in}) or intrathoracic pressure, can be measured during breathing (dynamically) using an esophageal probe ($\approx P_{pl}$). The intensity of suction (negative pressure) increases when the chest expands during inspiration, and decreases during expiration (\rightarrow **B**). P_{pl} usually does not become positive unless there is very forceful expiration requiring the use of expiratory muscles. The difference between the alveolar and the pleural pressure (PA - Ppl) is called **transpulmonary pressure** (\rightarrow p. 114).

Characterization of breathing activity. The terms hyperpnea and hypopnea are used to describe abnormal increases or decreases in the depth and rate of respiratory movements. Tachypnea (too fast), bradypnea (too slow), and apnea (cessation of breathing) describe abnormal changes in the respiratory rate. The terms hyperventilation and hypoventilation imply that the volume of exhaled CO2 is larger or smaller, respectively, than the rate of CO2 production, and the arterial partial pressure of CO₂ (Paco₂) decreases or rises accordingly $(\rightarrow p. 142)$. Dyspnea is used to describe difficult or labored breathing, and orthopnea occurs when breathing is difficult except in an upright position.





Purification of Respiratory Air

Inhaled foreign particles are trapped by **mucus** in the nose, throat, trachea, and bronchial tree. The entrapped particles are engulfed by macrophages and are driven back to the trachea by the cilia lining the bronchial epithelium. Cilial escalator: The cilia move at a rate of 5-10 s⁻¹ and propel the mucus towards the mouth at a rate of 1 cm/min on a film of fluid secreted by the epithelium. Heavy smoking, mucoviscidosis genetic defects can impair cilial transport. A volume of 10-100 mL of mucus is produced each day, depending on the type and frequency of local irritation (e.g., smoke inhalation) and vagal stimulation. Mucus is usually swallowed, and the fluid fraction is absorbed in the gastrointestinal tract.

Artificial Respiration

Mouth-to-mouth resuscitation is an emergency measure performed when someone stops breathing. The patient is placed flat on the back. While pinching the patient's nostrils shut, the aid-giver places his or her mouth on the patient's mouth and blows forcefully into the patient's lungs (\rightarrow A3). This raises the alveolar pressure (\rightarrow p. 108) in the patient's lungs relative to the atmospheric pressure outside the chest and causes the lungs and chest to expand (inspiration). The rescuer then removes his or her mouth to allow the patient to exhale. Expulsion of the air blown into the lungs (expiration) occurs due to the intrinsic elastic recoil of the lungs and chest (\rightarrow p. 109 A2). This process can be accelerated by pressing down on the chest. The rescuer should ventilate the patient at a rate of about 16 min⁻¹. The exspiratory O_2 fraction (\rightarrow p. 107 A) of the rescuer is high enough to adequately oxygenate the patient's blood. The color change in the patient's skin from blue (cyanosis; \rightarrow p. 130) to pink indicates that a resuscitation attempt was successful

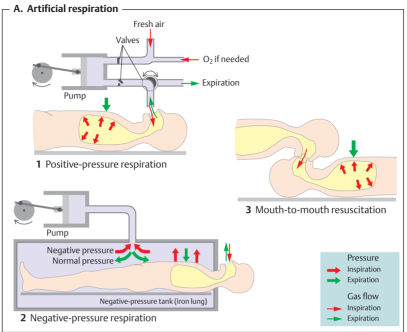
Mechanical ventilation. Mechanical *intermittent positive pressure ventilation* (**IPPV**) works on the same principle. This technique is used when the respiratory muscles are paralyzed due to disease, anesthesia, etc. The pump of the respirator drives air into the patient's lung during inspiration (\rightarrow **A1**). The external inspiratory and expiratory pathways are separatory than the patient of the pati

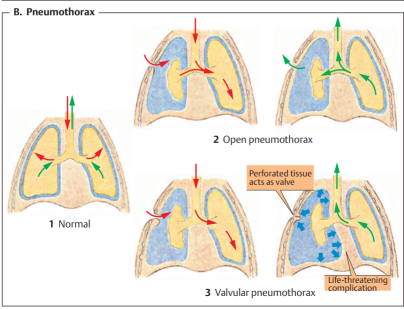
rated by a valve (close to the patient's mouth as possible) to prevent enlargement of dead space (\rightarrow p. 114). Ventilation frequency, tidal volume, inspiratory flow, as well as duration of inspiration and expiration can be preselected at the respirator. The drawback of this type of ventilation is that venous return to the heart is impaired to some extent (\rightarrow p. 204). Today, the standard technique of mechanical respiration is continuous positive pressure ventilation (CPPV). In contrast to IPPV, the endexpiratory pressure is kept positive (PEEP) in CPPV. In any case, all ventilated patients should be continuously monitored (expiratory gas fraction; blood gas composition, etc.).

The **iron lung** (Drinker respirator) makes use of negative-pressure respiration (\rightarrow A2). The patient's body is enclosed from the neck down in a metal tank. To achieve inhalation, pressure in the tank is decreased to a level below ambient pressure and, thus, below alveolar pressure. This pressure difference causes the chest to expand (inspiratory phase), and the cessation of negative pressure in the tank allows the patient to breathe out (expiratory phase). This type of respirator is used to ventilate patients who require long-term mechanical ventilation due to paralytic diseases, such as polio.

Pneumothorax

Pneumothorax occurs when air enters the pleural space and P_{pl} falls to zero ($\rightarrow p$. 108), which can lead to collapse of the affected lung due to elastic recoil and respiratory failure (\rightarrow **B**). The contralateral lung is also impaired because a portion of the inspired air travels back and forth between the healthy and collapsed lung and is not available for gas exchange. Closed pneumothorax, i.e., the leakage of air from the alveolar space into the pleural space, can occur spontaneously (e.g., lung rupture due to bullous emphysema) or due to lung injury (e.g., during mechanical ventilation = barotrauma; → p. 134). Open pneumo**thorax** $(\rightarrow B2)$ can be caused by an open chest wound or blunt chest trauma (e.g., penetration of the pleura by a broken rib). **Valvular pneumothorax** $(\rightarrow B3)$ is a life-threatening form of pneumothorax that occurs when air enters the pleural space with every breath and can no longer be expelled. A flap of acts like a valve. Positive pressure develops in the pleural space on the affected side, as well as in the rest of the thoracic cavity. Since the tidal volume increases due to hypoxia, high pressure levels (4 kPa = 30 mmHg) quickly develop. This leads to increasing impairment of cardiac filling and compression of the healthy contralateral lung. Treatment of valvular pneumothorax consists of slow drainage of excess pressure and measures to prevent further valvular action.





Lung Volumes and their Measurement

At the end of normal quiet expiration, the lung-chest system returns to its intrinsic resting position. About 0.5 L of air is taken in with each breath during normal quiet respiration; this is called the resting tidal volume (VT). Inspiration can be increased by another 3 L or so on forced (maximum) inspiration; this is called the inspiratory reserve volume (IRV). Likewise, expiration can be increased by about 1.7 L more on forced (maximum) expiration. This is called the expiratory reserve volume (ERV). These reserve volumes are used during strenuous physical exercise (\rightarrow p. 74) and in other situations where normal tidal volumes are insufficient. Even after forced expiration. about 1.3 L of air remains in the lungs; this is called the residual volume (RV). Lung capacities are sums of the individual lung volumes. The vital capacity (VC) is the maximum volume of air that can be moved in and out in a single breath. Therefore, VC = VT + IRV + ERV. The average 20-year-old male with a height of 1.80 m has a VC of about 5.3 L. Vital capacity decreases and residual volume increases with age (1.5 \Rightarrow 3 L). The **total lung capacity** is the sum of VC and RV-normally 6 to 7L. The functional residual capacity is the sum of ERV and RV (\rightarrow A and p. 114). The inspiratory capacity is the sum of VT and IRV. All numerical values of these volumes apply under body temperature-pressure saturation (BTPS) conditions (see below).

Spirometry. These lung volumes and capacities (except FRC, RV) can be measured by routine spirometry. The *spirometer* (\rightarrow **A**) consists usually of a water-filled tank with a bell-shaped floating device. A tube connects the air space within the spirometer (\rightarrow **A**) with the airways of the test subject. A counterweight is placed on the bell. The position of the bell indicates how much air is in the spirometer and is calibrated in volume units (L_{ATPS} ; see below). The bell on the spirometer rises when the test subject blows into the device (expiration), and falls during inspiration (\rightarrow **A**).

If the spirometer is equipped with a recording device (*spirograph*), it can be also used for graphic measurement of the total ventilation per unit time ($\dot{V}E$; \rightarrow pp. 106 and 118),

compliance (\rightarrow p. 116), O₂ consumption (\dot{V} O₂), and in dynamic lung function tests (\rightarrow p. 118).

Range of normal variation. Lung volumes and capacities vary greatly according to age, height, physical constitution, sex, and degree of physical fitness. The range of normal variation of VC, for example, is 2.5 to 7 L. Empirical formulas were therefore developed to create normative values for better interpretation of lung function tests. For instance, the following formulas are used to calculate the range of normal values for VC in Caucasians:

Men:
$$VC = 5.2 h - 0.022 a - 3.6 (\pm 0.58)$$

Women: $VC = 5.2 h - 0.018 a - 4.36 (\pm 0.42)$,

where h = height (in meters) and a = age (in years); the standard deviation is given in parentheses. Because of the broad range of normal variation, patients with mild pulmonary disease may go undetected. Patients with lung disease should ideally be monitored by recording baseline values and observing *changes* over the course of time.

Conversion of respiratory volumes. The volume, V, of a gas (in L or m^3 ; $1 m^3 = 1000 L$) can be obtained from the amount, M, of the gas (in mol), absolute temperature, T (in K), and total pressure, P (in Pa), using the *ideal gas equation*:

$$V = M \cdot R \cdot T/P, \qquad [5.2]$$

where P is barometric pressure (P_B) minus water partial pressure (PH₂o; \rightarrow p. 106) and R is the universal gas constant = 8.31 J K⁻¹ mol⁻¹.

Volume conditions

STPD: Standard temperature pressure dry (273 K, 101 kPa, PH₂O = 0)

ATPS: Ambient temperature pressure H_2O -saturated $(T_{amb}, P_B, P_{H_2O} \text{ at } T_{Amb})$

BTPS: Body temperature pressure-saturated (310 K, P_B , $P_{H>0}$ = 6.25 kPa)

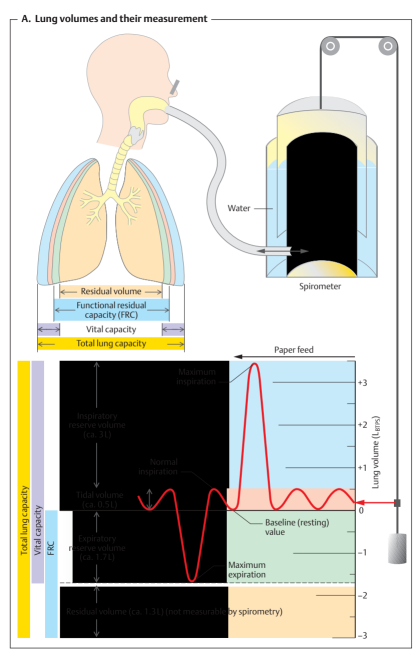
It follows that:

 $V_{STPD} = M \cdot R \cdot 273/101000 [m^3]$

 $V_{ATPS} = M \cdot R \cdot T_{Amb}/(P_B - P_{H_2O}) [m^3]$

 $V_{BTPS} = M \cdot R \cdot 310/(P_B - 6250) [m^3].$

Conversion factors are derived from the respective quotients (M · R is a reducing factor). Example: $V_{BTPS}/V_{STPD} = 1.17$. If V_{ATPS} is measured by spirometry at room temperature ($T_{Amb} = 20\,^{\circ}\text{C}$; $P_{H2}O_{sat} = 2.3\,\text{kPa}$) and $P_{B} = 101\,\text{kPa}$, $V_{BTPS} \approx 1.1\,V_{ATPS}$ and $V_{STPD} \approx 0.9\,V_{ATPS}$.



114

Dead Space, Residual Volume, Airway Resistance

The exchange of gases in the respiratory tract occurs in the alveoli. Only a portion of the tidal volume (VT) reaches the alveoli: this is known as the alveolar part (VA). The rest goes to dead space (not involved in gas exchange) and is therefore called **dead space** volume (Vp). The oral, nasal, and pharyngeal cavities plus the trachea and bronchi are jointly known as physiological dead space or conducting zone of the airways. The physiological dead space (ca. 0.15 L) is approximately equal to the functional dead space, which becomes larger than physiological dead space when the exchange of gases fails to take place in a portion of the alveoli (\rightarrow p. 120). The functions of dead space are to conduct incoming air to the alveoli and to purify (\rightarrow p. 110), humidify, and warm inspired ambient air. Dead space is also an element of the vocal organ (\rightarrow p. 370).

The **Bohr equation** $(\rightarrow A)$ can be used to **esti**mate the dead space.

Derivation: The expired tidal volume V_T is equal to the sum of its alveolar part V_A plus dead space $V_D (\rightarrow A$, top). Each of these three variables has a characteristic CO₂ fraction (\rightarrow p. 376): FE_{CO₂} in V_T, FA_{CO₂} in V_A, and FICO, in VD. FICO, is extremely small and therefore negligible. The product of each of the three volumes and its corresponding CO₂ fraction gives the volume of CO2 for each. The CO2 volume in the expired air (VT · FECO₂) equals the sum of the CO₂ volumes in its two components, i.e. in Va and VD $(\rightarrow A)$.

Thus, three values must be known to **determine** the dead space: VT, FECO, and FACO. VT can be measured using a spirometer, and FECO, and FACO, can be measured using a Bunte glass burette or an infrared absorption spectrometer. FACO, is present in the last expired portion of V_T—i.e., in alveolar gas. This value can be measured using a Rahn valve or similar device.

The functional residual capacity (FRC) is the amount of air remaining in the lungs at the end of normal quiet expiration, and the residual volume (RV) is the amount present after forced maximum expiration (\rightarrow p. 112). About 0.35 L of air (V_A) reaches the alveolar space with each breath during normal quiet respiration. Therefore, only about 12% of the 3 L total FRC is renewed at rest. The composition of gases in the alveolar space therefore remains relatively

constant.

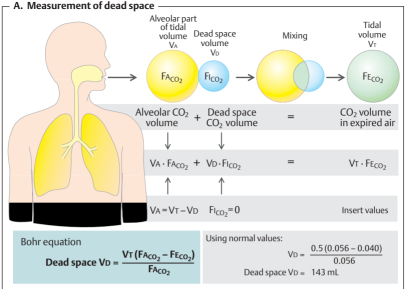
indirect techniques such as helium dilution $(\rightarrow B)$. Helium (He) is a poorly soluble inert gas. The test subject is instructed to repeatedly inhale and exhale a known volume (VSp) of a helium-containing gas mixture (e.g., $F_{He_0} = 0.1$) out of and into a spirometer. The helium distributes evenly in the lungs (VL) and spirometer $(\rightarrow B)$ and is thereby diluted $(F_{He_x} < F_{He_0})$. Since the total helium volume does not change, the known initial helium volume (V_{Sp} · F_{Hen}) is equal to the final helium volume $(V_{Sp} + V_L) \cdot F_{He_x}$. V_L can be determined once F_{He}, in the spirometer has been measured at the end of the test $(\rightarrow B)$. VL will be equivalent to RV if the test was started after a forced expiration, and will be equivalent to FRC if the test was started after normal expiration, i.e. from the resting position of lung and chest. The helium dilution method measures gases in ventilated airways only.

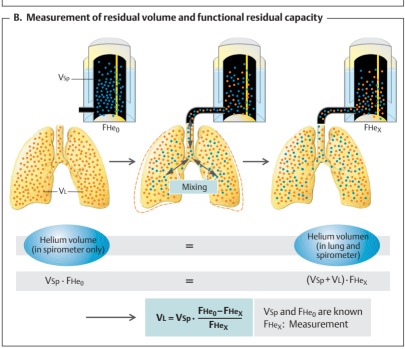
Body plethysmography can also detect gases in encapsulated spaces (e.g., cysts) in the lung. The test subject is placed in an airtight chamber and instructed to breathe through a pneumotachygraph (instrument for recording the flow rate of respired air). At the same time, respiration-dependent changes in air pressure in the subject's mouth and in the chamber are continuously recorded. FRC and RV can be derived from these measurements.

Such measurements can also be used to determine airway resistance. Rt. which is defined as the driving pressure gradient between the alveoli and the atmosphere divided by the air flow per unit time. Airway resistance is very low under normal conditions, especially during inspiration when (a) the lungs become more expanded (lateral traction of the airways), and (b) the transpulmonary pressure (PA-Ppl) rises $(\rightarrow p. 108)$. PA-P_{pl} represents the transmural pressure of the airways and widens them more and more as it increases. Airway resistance may become too high when the airway is narrowed by mucus-e.g., in chronic obstructive pulmonary disease, or when its smooth muscle contracts, e.g. in asthma (\rightarrow p.118).

The residual volume (RV) fraction of the total luna capacity (TLC) is clinically significant (\rightarrow p. 112). This fraction normally is no more than 0.25 in healthy subjects and somewhat higher in old age. It can rise to 0.55 and higher when pathological enlargement of the alveoli has occurred due, for example, to emphysema. The RV/TLC fraction is therefore a rough measure of the severity of such diseases.

Measurement of FRC and RV cannot be per-





Pressure-Volume Curve, Respiratory Work

Resting position (RP) is the position to which the lung-chest system returns at the end of normal quiet expiration; this lung volume equals the functional residual capacity (FRC, \rightarrow p. 114). Its value is set at zero ($V_{pulm} = 0$) in **A-C.** RP $(\rightarrow A1)$ is a stable central position characterized by the reciprocal cancellation of two passive forces: chest expansion force (CEF) and lung contraction force (LCF). As we breathe in and out, the lung - chest system makes excursions from resting position; thus, LCF > CEF during inspiration, and CEF > LCF during expiration. The difference between LCF and CEF, i.e. the net force (\rightarrow blue arrows in A2. **3, 5, 6**), is equal to the alveolar pressure (PA → p. 108) if the airway is closed (e.g., by turning a stopcock, as in A1-3, 5, 6) after a known air volume has been inhaled $(V_{pulm} > 0, \rightarrow A2)$ from a spirometer or expelled into it ($V_{pulm} < 0$, → A3). (In the resting position, CEF = LCF, and PA = 0). Therefore, the relationship between V_{pulm} and PA in the lung-chest system can be determined as illustrated by the static resting pressure—volume (PV) curve (→ blue curve in A-C) ("static" = measured while holding the breath; "resting" = with the respiratory muscles relaxed).

(Compression and expansion of V_{pulm} by a positive or negative PA during measurement has to be taken into account; \rightarrow **A**, dark-gray areas).

The **slope** of the static resting PV curve, $\Delta V_{pulm}/\Delta P_A$, represents the (static) **compliance** of the lung-chest system (\rightarrow B). The steepest part of the curve (range of greatest compliance; ca. 1 L/kPa in an adult) lies between RP and $V_{pulm}=1$ L. This is in the normal respiratory range. The curve loses its steepness, i.e. compliance decreases, in old age or in the presence of lung disease. As a result, greater effort is needed to breathe the same tidal volume.

The above statements apply to lung-and-chest compliance. It is also possible to calculate compliance for the chest wall alone $(\Delta V_A/\Delta P_{pl} = 2 L/kPa)$ or for the lung alone $(\Delta V/\Delta [PA - P_{pl}] = 2 L/kPa)$ when the pleural pressure (P_{pl}) is known $(\rightarrow p. 108)$.

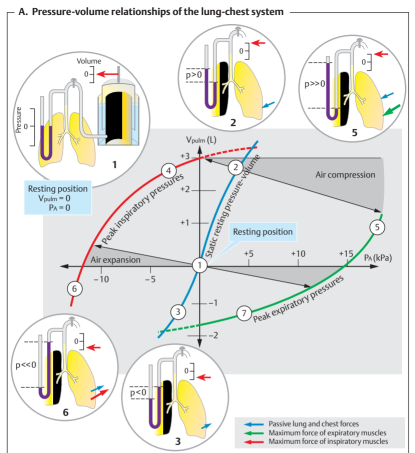
PV relationships can also be plotted during maximum expiratory and inspiratory effort to

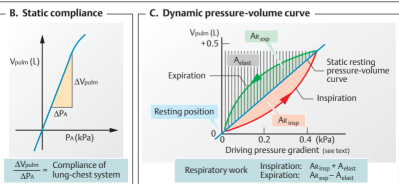
determine the **peak expiratory** and **inspiratory pressures** (\rightarrow **A**, red and green curves). Only a very small pressure can be generated from a position of near-maximum expiration ($V_{pulm} \ll 0$; \rightarrow **A7**) compared to a peak pressure of about 15 kPa (\approx 110 mmHg) at $V_{pulm} \gg 0$ (Valsalva's maneuver; \rightarrow **A5**). Likewise, the greetest negative pressure (suction) (ca. - 10 kPa = 75 mmHg) can be generated from a position of maximum expiration (Müller's maneuver; \rightarrow **A6**), but not from an inspiratory position (\rightarrow **A4**).

A **dynamic PV curve** is obtained *during* respiration (\rightarrow **C**). The result is a loop consisting of the opposing inspiratory (red) and expiratory (green) curves transected by the resting curve (blue) because airway flow resistance (RL) must be overcome (mainly in the upper and middle airways) while inhaling in the one direction and exhaling in the other. The *driving* pressure gradients (ΔP) also oppose each other (inspiratory PA < 0; expiratory PA > 0; \rightarrow p. 109 B). As in Ohm's law, $\Delta P = RL \cdot \text{respiratory}$ flow rate (\hat{V}). Therefore, ΔP must increase if the bronchial tubes narrow and/or if the respiratory flow rate increases (\rightarrow **C**).

In **asthma**, the airway radius (r) decreases and a very high ΔP is needed for normal ventilation (RL $\approx 1/r^4$ l). During exspiration, a high ΔP decreases the *transulmonary pressure* (= PA - P_{pl}) and thereby squeezes the airways (R_L ↑). The high RL results in a pressure decrease along the expiratory airway (P_{airway} \downarrow) until P_{airway} - P_{pl} < 0. At this point, the airway will collapse. This is called **dynamic airway compression**, which often results in a life-threatening vicious cycle: $r \downarrow \Rightarrow \Delta P \uparrow \Rightarrow r \downarrow \downarrow \Rightarrow \Delta P \uparrow \uparrow \uparrow \dots$

Respiratory work. The colored areas within the loop (A_{Rinsp} and A_{Rexp} ; \rightarrow C) represent the inspiratory and expiratory PV work (\rightarrow p. 374) exerted to overcome flow resistance. The cross-hatched area (\rightarrow C) is the work required to overcome the intrinsic elastic force of the lungs and chest (A_{elast}). *Inspiratory work* is defined as A_{Rinsp} + A_{elast} . The inspiratory muscles (\rightarrow p. 108) must overcome the elastic force, whereas the same elastic force provides the (passive) driving force for expiration at rest (sign reverses for A_{elast}). Thus, *expiratory work* is A_{Rexp} - A_{elast} . Expiration can also require muscle energy if A_{Rexp} becomes larger than A_{elast} —e.g., during forced respiration or if R_L is elevated.





Surface Tension, Surfactant

Surface tension is the main factor that determines the *compliance* of the lung-chest system (\rightarrow p. 116) and develops at gas-liquid interfaces or, in the case of the lungs, on the *gas exchange surface of the alveoli* (ca. 100 m²).

The effectiveness of these forces can be demonstrated by filling an isolated and completely collapsed lung with (a) air or (b) liquid. In example (a), the lung exerts a much higher resistance, especially at the beginning of the filling phase. This represents the **opening pressure**, which raises the alveolar pressure (PA) to about 2 kPa or 15 mmHg when the total lung capacity is reached (\rightarrow p. 113 A). In example (b), the resistance and therefore PA is only one-fourth as large. Accordingly, the larger pressure requirement in example (a) is required to overcome surface tension.

If a gas bubble with radius r is surrounded by liquid, the surface tension γ (N·m⁻¹) of the liquid raises the pressure inside the bubble relative to the outside pressure (transmural pressure $\Delta P > 0$). According to Laplace's law (cf. p. 188):

$$\Delta P = 2\gamma/r$$
 (Pa). [5.3] Since γ normally remains constant for the respective liquid (e.g., plasma: $10^{-3} \text{ N} \cdot \text{m}^{-1}$), ΔP becomes larger and larger as r decreases.

Soap bubble model. If a flat soap bubble is positioned on the opening of a cylinder, r will be relatively large (\rightarrow A1) and Δ P small. (Since two air-liquid interfaces have to be considered in this case, Eq. 5.3 yields $\Delta P = 4\gamma/r$). For the bubble volume to expand, r must initially decrease and ΔP must increase ($\rightarrow A2$). Hence, a relatively high "opening pressure" is required. As the bubble further expands, r increases again (\rightarrow A3) and the pressure requirement/volume expansion ratio decreases. The alveoli work in a similar fashion. This model demonstrates that, in the case of two alveoli connected with each other $(\rightarrow A4)$, the smaller one (ΔP_2 high) would normally become even smaller while the larger one (ΔP_1 low) becomes larger due to pressure equalization.

Surfactant (surface-active agent) lining the inner alveolar surface prevents this problem by lowering γ in smaller alveoli more potently than in larger alveoli. Surfactant is a mixture of

proteins and phospholipids (chiefly dipalmitoyl lecithin) secreted by alveolar type II cells. Respiratory distress syndrome of the newborn, a serious pulmonary gas exchange disorder, is caused by failure of the immature lung to produce sufficient quantities of surfactant. Lung damage related to O_2 toxicity (\rightarrow p. 136) is also partly due to oxidative destruction of surfactant, leading to reduced compliance. This can ultimately result in alveolar collapse (atelectasis) and pulmonary edema.

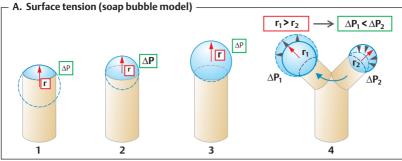
Dynamic Lung Function Tests

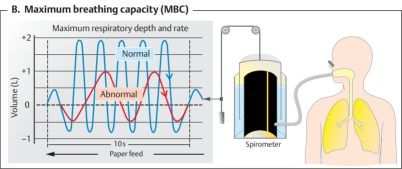
The **maximum breathing capacity** (MBC) is the greatest volume of gas that can be breathed (for $10\,\mathrm{s}$) by voluntarily increasing the tidal volume and respiratory rate (\rightarrow B). The MBC normally ranges from 120 to 170 L/min. This capacity can be useful for monitoring diseases affecting the respiratory muscles, e.g., myasthenia gravis.

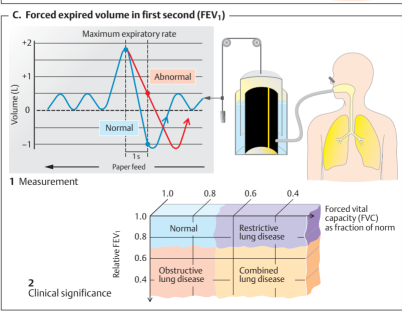
The **forced expiratory volume** (FEV or Tiffeneau test) is the maximum volume of gas that can be expelled from the lungs. In clinical medicine, FEV in the first second (FEV₁) is routinely measured. When its absolute value is related to the *forced vital capacity* (FVC), the *relative FEV*₁ (normally > 0.7) is obtained. (FVC is the maximum volume of gas that can be expelled from the lungs as quickly and as forcefully as possible from a position of full inspiration; \rightarrow C). It is often slightly lower than the vital capacity VC (\rightarrow p. 112). **Maximum expiratory flow**, which is measured using a pneumotachygraph during FVC measurement, is around 10 L/s.

Dynamic lung function tests are useful for distinguishing restrictive lung disease (RLD) from obstructive lung disease (OLD). RLD is characterized by a functional reduction of lung volume, as in pulmonary edema, pneumonia and impaired lung inflation due to spinal curvature, whereas OLD is characterized by physical narrowing of the airways, as in asthma, bronchitis, emphysema, and vocal cord paralysis (\rightarrow C2).

As with VC (\rightarrow p. 112), empirical formulas are also used to standardize FVC for age, height and sex.







120

Pulmonary Gas Exchange

Alveolar ventilation. Only the alveolar part (VA) of the tidal volume (VT) reaches the alveoli. The rest goes to dead space (VD). It follows that VA = VT - VD (L) (\rightarrow p. 114). Multiplying these volumes by the *respiratory rate* (**f** in min⁻¹) results in the respective ventilation, i.e., \dot{V}_A , \dot{V}_E (or \dot{V}_T), and \dot{V}_D . Thus, $\dot{V}_A = \dot{V}_E - \dot{V}_D$ (L·min⁻¹). Since V_D is anatomically determined, \dot{V}_D (= $V_D \cdot$ f) rises with f. If, at a given total ventilation ($\dot{V}_E = V_T \cdot f$), the breathing becomes more frequent (f †) yet more shallow ($V_T \downarrow$), \dot{V}_A will decrease because \dot{V}_D increases.

Example: At a \dot{V}_E of 8 L·min⁻¹, a VD of 0.15 L and a normal respiratory rate f of 16 min⁻¹ \dot{V}_A = 5.6 L·min⁻¹ or 70% of \dot{V}_E . When f is doubled and V \dot{V} drops to onehalf, \dot{V}_A drops to 3.2 L·min⁻¹ or 40% of V \dot{V}_A , although \dot{V}_E (8 L·min⁻¹) remains unchanged.

Alveolar gas exchange can therefore decrease due to flat breathing and panting (e.g., due to a painful rib fracture) or artificial enlargement of $V_D (\rightarrow p. 134)$.

 O_2 **consumption** (\dot{V}_{O_2}) is calculated as the *difference between* the *inspired* O_2 volume/time $(=\dot{V}_E\cdot F_{IO_2}, \text{ and the } expired \ O_2 \text{ volume/time}$ $(=\dot{V}_E\cdot F_{EO_2}. \text{ Therefore, } \dot{V}_{O_2}=\dot{V}_E \text{ (F}_{IO_2}-F_{EO_2}). \text{ At } \text{rest, } \dot{V}_{O_3}\approx 8 \text{ (0.21-0.17)}=0.32 \text{ L} \cdot \text{min}^{-1}.$

The eliminated CO₂ volume (\dot{V} co₂) is calculated as \dot{V} T·Fe_{CO₂} (\approx 0.26 L·min⁻¹ at rest; Fi_{CO₂} \approx 0). \dot{V} o₂ and \dot{V} co₂ increase about tenfold during strenuous physical work (\rightarrow p. 74). The \dot{V} co₂ to \dot{V} o₂ ratio is called the **respiratory quotient** (**RQ**), which depends on a person's nutritional state. RQ ranges from 0.7 to 1.0 (\rightarrow p. 228).

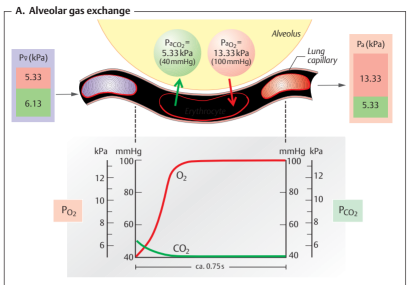
The **exchange of gases** between the alveoli and the blood occurs by *diffusion*, as described by Fick's law of diffusion (\rightarrow Eq. 1.7, p. 22,). The driving "force" for this diffusion is provided by the *partial pressure differences* between alveolar space and erythrocytes in pulmonary capillary blood (\rightarrow A). The mean alveolar partial pressure of O₂ (PA_{O2}) is about 13.3 kPa (100 mmHg) and that of CO₂ (PA_{CO2}) is about 5.3 kPa (40 mmHg). The mean partial pressures in the "venous" blood of the pulmonary artery are approx. 5.3 kPa (40 mmHg) for O₂ (PV_{O2}) and approx. 6.1 kPa (46 mmHg) for CO₂ (PV_{CO2}). Hence, the mean **partial pressure difference** between alveolus and capillary is

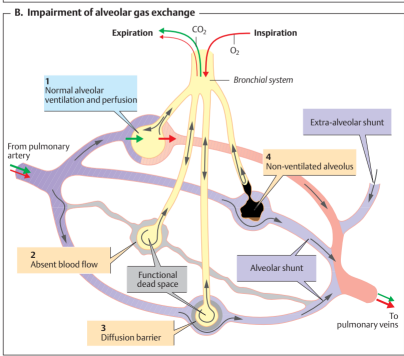
about 8 kPa (60 mmHg) for O_2 and about 0.8 kPa (6 mmHg) for CO_2 , although regional variation occurs (\rightarrow p. 122). PAO_2 , will rise when $PACO_2$, falls (e.g., due to hyperventilation) and vice versa (\rightarrow alveolar gas equation, p. 136).

O2 diffuses about 1-2 um from alveolus to bloodstream (diffusion distance). Under normal resting conditions, the blood in the pulmonary capillary is in contact with the alveolus for about 0.75 s. This **contact time** $(\rightarrow A)$ is long enough for the blood to equilibrate with the partial pressure of alveolar gases. The capillary blood is then arterialized. Po₂ and Pco₂ in arterialized blood (Pao, and Paco,) are about the same as the corresponding mean alveolar pressures (PAO2 and PACO2). However, venous blood enters the arterialized blood through arteriovenous shunts in the lung and from bronchial and thebesian veins $(\rightarrow B)$. This extra-alveolar shunt as well as ventilation-perfusion inequality ($\rightarrow p.122$) make the Pa₀ decrease from 13.3 kPa (after alveolar passage) to about 12.0 kPa (90 mmHg) in the aorta (Pa_{CO_2} increases slightly; $\rightarrow A$ and p. 107).

The small pressure difference of about 0.8 kPa is large enough for alveolar CO_2 exchange, since Krogh's diffusion coefficient K for CO_2 ($Kco_2 \approx 2.5 \cdot 10^{-16} \, \text{m}^2 \cdot \text{s}^{-1} \cdot \text{Pa}^{-1}$ in tissue) is 23 times larger than that for O_2 (\rightarrow p.22). Thus, CO_2 diffuses much more rapidly than O_2 . During physical work (high cardiac output), the contact time falls to a third of the resting value. If diffusion is impaired (see below), alveolar equilibration of O_2 partial pressure is less likely to occur during physical exercise than at rest.

Impairment of alveolar gas exchange can occur for several reasons: (a) when the blood flow rate along the alveolar capillaries decreases (e.g., due to pulmonary infarction; \rightarrow B2), (b) if a diffusion barrier exists (e.g., due to a thickened alveolar wall, as in pulmonary edema; \rightarrow B3), and (c) if alveolar ventilation is reduced (e.g., due to bronchial obstruction; \rightarrow B4). Cases B2 and B3 lead to an increase in functional dead space (\rightarrow p. 114); cases B3 and B4 lead to inadequate arterialization of the blood (alveolar shunt, i.e. non-arterialized blood mixing towards arterial blood). Gradual impairments of type B2 and B4 can occur even in healthy individuals (\rightarrow p. 122).





Pulmonary Blood Flow, Ventilation–Perfusion Ratio

Neglecting the slight amount of blood that reaches the lungs via the bronchial arteries, the mean **pulmonary perfusion** ($\dot{\mathbf{Q}}$), or blood flow to the lungs, is equal to the cardiac output (CO = 5–6 L/min). The **pulmonary arterial pressure** is about 25 mmHg in systole and 8 mmHg in diastole, with a mean ($\dot{\mathbf{P}}$) of about 15 mmHg. $\dot{\mathbf{P}}$ decreases to about 12 mmHg ($\dot{\mathbf{P}}$) in the precapillary region (up to the origin of the pulmonary capillaries) and about 8 mmHg in the postcapillary region ($\dot{\mathbf{P}}$). These values apply to the areas of the lung located at the level of the pulmonary valve.

Uneven distribution of blood flow within the lung $(\rightarrow A)$. Due to the additive effect of hydrostatic pressure (up to 12 mmHg), Pprecap increases in blood vessels below the pulmonary valves (near the base of the lung) when the chest is positioned upright. Near the apex of the lung, P_{precap} decreases in vessels above the pulmonary valve (\rightarrow A, zone 1). Under these conditions, P_{precap} can even drop to subatmospheric levels, and the mean alveolar pressure (PA) is atmospheric and can therefore cause extensive capillary compression (PA > P_{precap} > P_{postcap}; \rightarrow **A**). \dot{Q} per unit of lung volume is therefore very small. In the central parts of the lung (\rightarrow A, zone 2), luminal narrowing of capillaries can occur at their venous end, at least temporarily ($P_{precap} > P_A > P_{postcap}$), while the area near the base of the lung ($\rightarrow A$, zone 3) is continuously supplied with blood ($P_{precap} > P_{postcap} > P_A$). O per unit of lung volume therefore decreases from the apex of the lung to the base ($\rightarrow A$, B, red line).

Uneven distribution of alveolar ventilation. Alveolar ventilation (\mathring{V}_A) per unit of lung volume also increases from the apex to the base of the lungs due to the effects of gravity $(\rightarrow \mathbf{B}$, orange line), although not as much as \mathring{Q} . Therefore, the $\mathring{\mathbf{V}}_A/\mathring{\mathbf{Q}}$ ratio decreases from the apex to the base of the lung $(\rightarrow \mathbf{B})$, green curve and top scale).

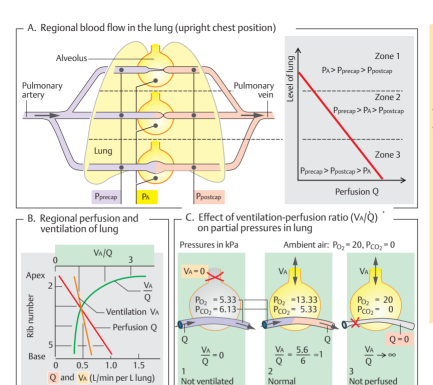
 \dot{V}_A/\dot{Q} **imbalance.** The mean \dot{V}_A/\dot{Q} for the entire lung is 0.93 (\rightarrow C2). This value is calculated from the mean alveolar ventilation \dot{V}_A (ca. 5.6 L/min) and total perfusion \dot{Q} (ca. 6 L/min), which is equal to the cardiac output (CO). Under extreme conditions in which one part of the lung is not ventilated at all, $\dot{V}_A/\dot{Q} = 0$ (\rightarrow C1). In the other extreme in which blood flow is absent (\dot{V}_A/\dot{Q} approaches infinity; \rightarrow C3), fresh air conditions will prevail in the alveoli (functional dead space; \rightarrow p. 120). \dot{V}_A/\dot{Q} can

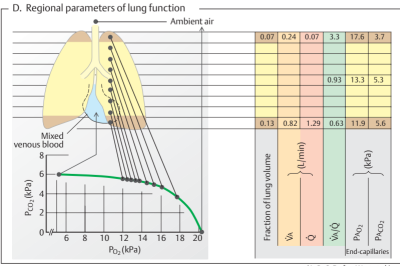
vary tremendously—theoretically, from 0 to ∞ . In this case, the PA_{O_2} will fluctuate between mixed venous PV_{O_2} and PI_{O_2} of (humidified) fresh air (\rightarrow **D**). In a healthy upright lung, \dot{V}_A/\dot{Q} decreases greatly (from 3.3 to 0.63) from apex to base at rest (\rightarrow **B**, green line); PA_{O_2} (PA_{CO_2}) is therefore 17.6 (3.7) kPa in the "hyperventilated" lung apex, 13.3 (5.3) kPa in the normally ventilated central zone, and 11.9 (5.6) kPa in the hypoventilated lung base. These changes are less pronounced during physical exercise because \dot{Q} also increases in zone 1 due to the corresponding increase in P_{DDCGD} .

 \dot{V}_A/\dot{Q} imbalance decreases the efficiency of the lungs for gas exchange. In spite of the high $P_{A_{0_2}}$ at the apex of the lung (ca. 17.6 kPa; \rightarrow **D**, right panel) and the fairly normal mean $P_{A_{0_2}}$ value, the relatively small \dot{Q} fraction of zone 1 contributes little to the total \dot{Q} of the pulmonary veins. In this case, $P_{A_{0_2}} < P_{A_{0_2}}$ and an alveolar–arterial O_2 difference ($A_aD_{0_2}$) exists (normally about 1.3 kPa). When a total arteriovenous shunt is present ($\dot{V}_A/\dot{Q} = 0$), even oxygen treatment will not help the patient, because it would not reach the pulmonary capillary bed (\rightarrow **C1**).

Hypoxic vasoconstriction regulates alveolar perfusion and prevents the development of extreme \dot{V}_A/\dot{Q} ratios. When the P_{AO_2} decreases sharply, receptors in the alveoli emit local signals that trigger constriction of the supplying blood vessels. This throttles shunts in poorly ventilated or non-ventilated regions of the lung, thereby routing larger quantities of blood for gas exchange to more productive regions.

 \dot{V}_A/\dot{Q} imbalance can cause severe complications in many lung diseases. In *shock lung*, for example, shunts can comprise 50% of \dot{Q} . Lifethreatening lung failure can quickly develop if a concomitant pulmonary edema, alveolar diffusion barrier, or surfactant disorder exists (\rightarrow p. 118).





(A, B, C, D after West et al.)

123

CO₂ Transport in Blood

Carbon dioxide (CO₂) is the end-product of energy metabolism (\rightarrow p. 228). CO₂ produced by cells of the body undergoes **physical dissolution** and diffuses into adjacent blood capillaries. A small portion of CO₂ in the blood remains dissolved, while the rest is **chemically bound** in form of HCO₃⁻ and carbamate residues of hemoglobin (\rightarrow A, lower panel, blue arrows; \rightarrow arteriovenous CO₂ difference given in the table). Circulating CO₂-loaded blood reaches the pulmonary capillaries via the right heart. CO₂ entering the pulmonary capillaries is released from the compounds (\rightarrow A, red arrows), diffuses into the alveoli, and is expired into the atmosphere (\rightarrow A and p. 106).

The enzyme **carbonic anhydrase** (carbonate dehydratase) catalyzes the reaction

 $HCO_3^- + H^+$ $CO_2 + H_2O$ in erythrocytes (\rightarrow **A5**, **7**). Because it accelerates the establishment of equilibrium, the short contact time (<1s) between red blood cells and alveolus or peripheral tissue is sufficient for the transformation CO_2 HCO_3^- .

 CO_2 diffusing from the **peripheral cells** (\rightarrow **A**, bottom panel: "Tissue") increases PCO2 (approx. 5.3 kPa = 40 mmHg in arterial blood) to a mean venous P_{CO_2} of about 6.3 kPa = 47 mmHg. It also increases the concentration of CO2 dissolved in plasma. However, the major portion of the CO₂ diffuses into red blood cells, thereby increasing their content of dissolved CO2. CO2 (+ H2O) within the cells is converted to $HCO_3^-(\rightarrow A5, 2)$ and hemoglobin carbamate (\rightarrow A3). The HCO₃⁻ concentration in erythrocytes therefore becomes higher than in plasma. As a result, about three-quarters of the HCO3- ions exit the erythrocytes by way of an HCO3-/Cl- antiporter. This anion exchange is also called Hamburger shift $(\rightarrow A4)$.

 H^+ ions are liberated when CO_2 in red cells circulating in the periphery is converted to HCO_3^- and hemoglobin (Hb) carbamate.

Bicarbonate formation:

$$CO_2 + H_2O HCO_3^- + H^+,$$
 [5.4]

Hemoglobin carbamate formation:

 $Hb-NH_2+CO_2$ $Hb-NH-COO^-+H^+$. [5.5] **Hemoglobin (Hb)** is a key **buffer for H**⁺ ions in the red cells (\rightarrow **A6**; see also p. 140, "Non-bicarbonate buffers"). Since the removal of H⁺ ions

in reactions 5.4 and 5.5 prevents the rapid establishment of equilibrium, large quantities of CO_2 can be incorporated in HCO_3^- and Hb carbamate. Deoxygenated hemoglobin (Hb) can take up more H^+ ions than oxygenated hemoglobin (Oxy-Hb) because Hb is a weaker acid (\rightarrow A). This promotes CO_2 uptake in the peripheral circulation ($Haldane\ effect$) because of the simultaneous liberation of O_2 from erythrocytes, i.e. deoxygenation of Oxy-Hb to Hb.

In the **pulmonary capillaries**, these reactions proceed in the opposite direction (\rightarrow **A**, top panel, red and black arrows). Since the P_{CO_2} in alveoli is lower than in venous blood, CO_2 diffuses into the alveoli, and reactions 5.4 and 5.5 proceed to the left. CO_2 is released from HCO_3^- and Hb carbamate whereby H^* ions (released from Hb) are bound in both reactions (\rightarrow **A7**, **A8**), and the direction of HCO_3^-/Cl^- exchange reverses (\rightarrow **A9**). Reoxygenation of Hb to Oxy-Hb in the lung promotes this process by increasing the supply of H^* ions ($Haldane\ effect$).

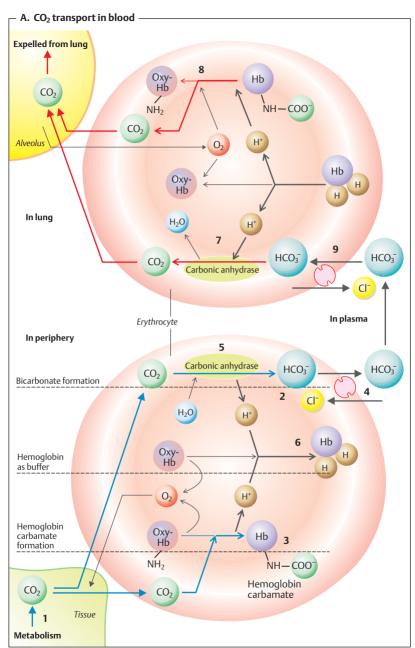
 CO_2 distribution in blood (mmol/L blood, 1 mmol = 22.26 mL CO_2)

	Dis- solved CO ₂	HCO ₃ -	Carba- mate	Total		
Arterial blood:						
Plasma*	0.7	13.2	0.1	14.0		
Erythrocytes**	0.5	6.5	1.1	8.1		
Blood	1.2	19.7	1.2	22.1		
Mixed venous blood:						
Plasma*	0.8	14.3	ca. 0.1	15.2		
Erythrocytes **	0.6	7.2	1.4	9.2		
Blood	1.4	21.5	1.5	24.4		
Arteriovenous CO ₂ difference in blood						

Arteriovenous CO₂ difference in blood
0.2 1.8 0.3 2.3

Percentage of total arteriovenous difference
9% 78% 13% 100%

^{*} Approx 0.55 L plasma/L blood; ** ca. 0.45 L erythrocytes/L blood



CO₂ Binding in Blood, CO₂ in CSF

The **total carbon dioxide** concentration (= chemically bound "CO₂" + dissolved CO₂) of mixed venous blood is about 24–25 mmol/L; that of arterial blood is roughly 22–23 mmol/L. Nearly 90% of this is present as HCO_3^- (\rightarrow **A**, right panel, and table on p. 124). The partial pressure of CO_2 (P_{CO_2}) is the chief factor that determines the CO_2 content of blood. The **CO₂ dissociation curve** illustrates how the total CO_2 concentration depends on P_{CO_2} (\rightarrow **A**).

The concentration of **dissolved CO₂**, [CO₂], in plasma is directly proportional to the Pco₂ in plasma and can be calculated as follows:

 $[CO_2] = \alpha_{CO_2} \cdot P_{CO_2} \text{ (mmol/L plasma}$ or mL/L plasma), [5.6]

where α_{CO_2} is the (Bunsen) solubility coefficient for CO_2 . At 37 °C,

 $\alpha_{CO_2} = 0.225 \text{ mmol} \cdot L^{-1} \cdot \text{kPa}^{-1}$,

After converting the amount of CO_2 into volume CO_2 (mL = mmol \cdot 22.26), this yields $\alpha_{CO_2} = 5 \text{ mL} \cdot \text{L}^{-1} \cdot \text{kPa}^{-1}$.

The *curve for dissolved* CO_2 is therefore linear $(\rightarrow A$, green line).

Since the buffering and carbamate formation capacities of hemoglobin are limited, the relation between **bound "CO₂"** and P_{CO_2} is curvilinear. The *dissociation curve for total* CO₂ is calculated from the sum of dissolved and bound CO₂ (\rightarrow **A**, red and violet lines).

CO2 binding with hemoglobin depends on the degree of oxygen saturation (S_{0_2}) of hemoglobin. Blood completely saturated with O2 is not able to bind as much CO2 as O2-free blood at equal P_{CO_2} levels ($\rightarrow A$, red and violet lines). When venous blood in the lungs is loaded with O2, the buffer capacity of hemoglobin and, consequently, the levels of chemical CO₂ binding decrease due to the Haldane effect (\rightarrow p. 124). Venous blood is never completely void of O2, but is always O2-saturated to a certain degree, depending on the degree of O_2 extraction (\rightarrow p. 130) of the organ in question. The S₀, of mixed venous blood is about 0.75. The CO_2 dissociation curve for S_{O_2} = 0.75 therefore lies between those for $S_{0_2} = 0.00$ and 1.00 (\rightarrow A, dotted line). In arterial blood, $P_{CO_2} \approx 5.33$ kPa and $S_{O_2} \approx 0.97$ (\rightarrow **A**, point a). In mixed venous blood, $P_{CO_2} \approx 6.27 \, \text{kPa}$ and S_{O_2} \approx 0.75 (\rightarrow **A**, point \triangledown). The normal range of CO₂

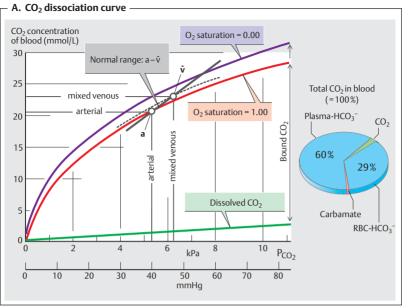
dissociation is determined by connecting these two points by a line called "physiologic CO₂ dissociation curve."

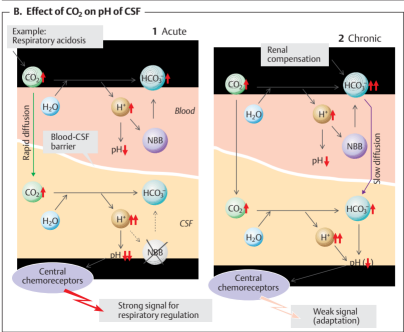
The concentration ratio of HCO_3^- to dissolved CO_2 in plasma and red blood cells differs (about 20:1 and 12:1, respectively). This reflects the difference in the pH of plasma (7.4) and erythrocytes (ca. 7.2) (\rightarrow p. 138ff.).

CO₂ in Cerebrospinal Fluid

Unlike HCO₃⁻ and H⁺, CO₂ can cross the bloodcerebrospinal fluid (CSF) barrier with relative ease (\rightarrow **B1** and p. 310). The P_{CO2} in CSF therefore adapts quickly to acute changes in the Pco. in blood. CO2-related (respiratory) pH changes in the body can be buffered by non-bicarbonate buffers (NBBs) only (\rightarrow p. 144). Since the concentration of non-bicarbonate buffers in CSF is very low, an acute rise in Pco2 (respiratory acidosis: $\rightarrow p. 144$) leads to a relatively sharp decrease in the pH of CSF (\rightarrow **B1**, pH $\downarrow\downarrow$). This decrease is registered by central chemosensors (or chemoreceptors) that adjust respiratory activity accordingly ($\rightarrow p$, 132). (In this book, sensory receptors are called sensors in order to distinguish them from hormone and transmitter receptors.)

The concentration of non-bicarbonate buffers in blood (hemoglobin, plasma proteins) is high. When the CO2 concentration increases, the liberated H+ ions are therefore effectively buffered in the blood. The actual HCO₃⁻ concentration in blood then rises relatively slowly, to ultimately become higher than in the CSF. As a result, HCO3- diffuses (relatively slowly) into the CSF (\rightarrow **B2**), resulting in a renewed increase in the pH of the CSF because the HCO3-/CO2 ratio increases $(\rightarrow p. 140)$. This, in turn, leads to a reduction in respiratory activity (via central chemosensors), a process enhanced by renal compensation, i.e., a pH increase through HCO3- retention (\rightarrow p. 144). By this mechanism, the body ultimately adapts to chronic elevation in Pco.i.e., a chronically elevated PCO2 will no longer represent a respiratory drive (cf. p. 132).





Binding and Transport of O2 in Blood

Hemoglobin (Hb) is the O₂-carrying protein of red blood cells (RBCs) (mol. mass: 64500 Da). Hb is also involved in CO₂ transport and is an important blood pH buffer (\rightarrow pp. 124 and 138ff.). Hb is a tetramer with 4 subunits (adults: 98%: $2\alpha + 2\beta = HbA$; $2\% 2\alpha + 2\delta =$ HbA2), each with its own heme group, Heme consists of porphyrin and Fe(II). Each of the four Fe(II) atoms (each linked with one histidine residue of Hb) binds reversibly with an O₂ molecule. This is referred to as **oxygenation** (not oxidation) of Hb to oxyhemoglobin (Oxy-Hb). The amount of O2 which combines with Hb depends on the partial pressure of $O_2(P_{O_2})$: **oxygen dissociation curve** (\rightarrow **A**, red line). The curve has a sigmoid shape, because initially bound O2 molecules change the conformation of the Hb tetramer (positive cooperativity) and thereby increase hemoglobin-O2 affinity.

When fully saturated with O_2 , 1 mol of tetrameric Hb combines with 4 mol O_2 , i.e., 64 500 g of Hb combine with $4 \times 22.4 \, \text{L}$ of O_2 . Thus, 1g Hb can theoretically transport 1.39 mL O_2 , or 1.35 mL in vivo (Hüfner number). The total Hb concentration of the blood ([Hb]_{total}) is a mean 150 g/L (\rightarrow p. 88), corresponding to a maximum O_2 concentration of 9.1 mmol/L or an O_2 fraction of 0.203 L O_2 /L blood. This **oxygen-carrying capacity** is a function of [Hb]_{total} (\rightarrow **A**, yellow and purple curves as compared to the red curve).

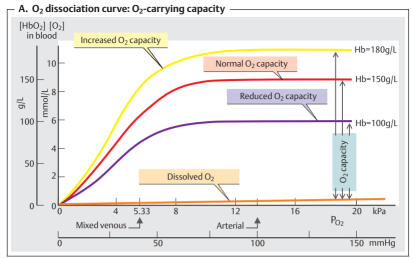
The O_2 content of blood is virtually equivalent to the amount of O_2 bound by Hb since only 1.4% of O_2 in blood is dissolved at a P_{O_2} of 13.3 kPa (\rightarrow A, orange line). The solubility coefficient (α_{O_2}), which is $10~\mu$ mol·[L of plasma]⁻¹·kPa⁻¹, is 22 times smaller than α_{CO_2} (\rightarrow p. 126).

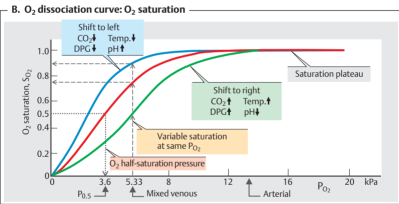
Oxygen saturation (S_{O_2}) is the fraction of Oxy-Hb relative to $[Hb]_{total}$, or the ratio of actual O_2 concentration/ O_2 -carrying capacity. At normal P_{O_2} in arterial blood (e.g., $P_{AO_2} = 12.6$ kPa or 95 mmHg), S_{O_2} will reach a saturation plateau at approx. 0.97, while S_{O_2} will still amount to 0.73 in mixed venous blood $(P\overline{V}_{O_2} = 5.33$ kPa or 40 mmHg). The venous S_{O_2} values in different organs can vary greatly $(\rightarrow p. 130)$.

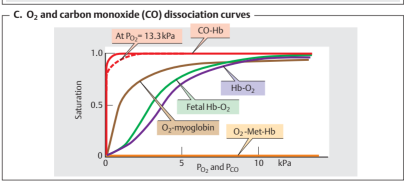
 O_2 dissociation is independent of total Hb if plotted as a function of $S_{O_2}(\rightarrow \mathbf{B})$. Changes in O_2 affinity to Hb can then be easily identified as **shifting of the O_2 dissociation curve**. A shift to

the right signifies an affinity decrease, and a shift to the left signifies an affinity increase, resulting in flattening and steepening, respectively, of the initial part of the curve. Shifts to the left are caused by increases in pH (with or without a Pco, decrease) and/or decreases in P_{CO₂}, temperature and 2,3-bisphosphoglycerate (BPG; normally 1 mol/mol Hb tetramer). Shifts to the right occur due to decreases in pH and/or increases in Pco, temperature and 2,3-BPG (\rightarrow B). The half-saturation pressure (P_{0.5} or P_{50}) of $O_2 (\rightarrow \mathbf{B}$, dotted lines) is the P_{02} at which S_{0} , is 0.5 or 50%. The $P_{0.5}$, which is normally 3.6 kPa or 27 mmHg, is a measure of shifting to the right $(P_{0.5} \uparrow)$ or left $(P_{0.5} \downarrow)$. Displacement of the O2 dissociation curve due to changes in pH and P_{CO_2} is called the **Bohr effect**. A shift to the right means that, in the periphery (pH \downarrow). $P_{CO_2} \uparrow$), larger quantities of O_2 can be absorbed from the blood without decreasing the Po, which is the driving force for O_2 diffusion ($\rightarrow B$. broken lines). A higher affinity for O₂ is then re-established in the pulmonary capillaries $(pH \uparrow, P_{CO_2} \downarrow)$. A shift to the left is useful when the PAO_2 is decreased (e.g., in altitude hypoxia), a situation where arterial So, lies to the left of the S_{0_2} plateau.

Myoglobin is an Fe(II)-containing muscle protein that serves as a short-term storage molecule for $O_2 (\rightarrow p.72)$. As it is monomeric (no positive cooperativity), its O2 dissociation curve at low Po2 is much steeper than that of HbA (\rightarrow **C**). Since the O₂ dissociation curve of **fetal Hb** $(2\alpha + 2\gamma = \text{HbF})$ is also steeper, S_{02} values of 45 to 70% can be reached in the fetal umbilical vein despite the low Po2 (3-4 kPa or 22-30 mmHg) of maternal placental blood. This is sufficient, because the fetal [Hb]total is 180 g/L. The carbon monoxide (CO) dissociation curve is extremely steep. Therefore, even tiny amounts of CO in the respiratory air will dissociate O2 from Hb. This can result in carbon monoxide poisoning $(\rightarrow C)$. Methemoglobin, Met-Hb (normally 1% of Hb), is formed from Hb by oxidation of Fe(II) to Fe(III) either spontaneously or via exogenous oxidants. Met-Hb cannot combine with $O_2 (\rightarrow \mathbb{C})$. Methemoglobin reductase reduces Fe(III) of Met-Hb back to Fe(II); deficiencies of this enzyme can cause methemoglobinemia, resulting in neonatal anoxia.







Internal (Tissue) Respiration, Hypoxia

O₂ diffuses from peripheral blood to adjacent tissues and CO₂ in the opposite direction $(\rightarrow pp. 20ff. and 106)$. Since CO_2 diffuses much faster (\rightarrow p. 120), O₂ diffusion is the limiting factor. Sufficient O2 delivery is ensured by a dense capillary network with a gas exchange area of about 1000 m². The diffusion distance $(\rightarrow R \text{ in } \mathbf{A})$ is only 10–25 µm. The driving force for diffusion is the difference in partial pressures of oxygen (ΔP_{0_2}) in capillary blood and mitochondria, where the Po2 must not fall below 0.1 kPa \approx 1 mmHg. Since P₀, decreases with distance parallel and perpendicular to the course of capillaries, the O2 supply to cells at the venous end far away from the capillaries (large R) is lowest, as shown using Krogh's cylinder model (\rightarrow A1). Since these cells are also the first to be affected by oxygen deficiency (hypoxia), this is sometimes called the "lethal corner" (\rightarrow A2).

Using Fick's principle (\rightarrow p. 106), **oxygen consumption** of a given organ, $\dot{V}o_2$ (in L/min), is calculated as the difference between the *a*rterial supply ($\dot{Q} \cdot [O_2]a$) and non-utilized *v*enous O_2 volume/time ($\dot{Q} \cdot [O_2]v$), where \dot{Q} is rate of blood flow in the organ (L/min) and $[O_2]$ is the oxygen fraction (LO₂/L blood):

 $\hat{\mathbf{V}}_{\mathbf{O_2}} = \hat{\mathbf{Q}} \left([\mathbf{O_2}]_a - [\mathbf{O_2}]_v \right)$ [5.7] To meet **increased** $\mathbf{O_2}$ **demands**, $\hat{\mathbf{Q}}$ *can therefore* be increased by **vasodilatation** in the organ in question and/or by **raising the oxygen extraction** ($\mathbf{E_{O_2}}$). \mathbf{E}_{O_2} describes the $\mathbf{O_2}$ consumption in the organ (= $\hat{\mathbf{Q}} \left([\mathbf{O_2}]_a - [\mathbf{O_2}]_v \right)$; see Eq. 5.7) relative to the arterial $\mathbf{O_2}$ supply ($\hat{\mathbf{Q}} \cdot [\mathbf{O_2}]_a$). Since $\hat{\mathbf{Q}}$ can be simplified.

 $E_{O_2} = ([O_2]_a - [O_2]_v)/[O_2]_a$ [5.8] E_{O_2} varies according to the type and function of the organ under resting conditions: skin 0.04 (4%), kidney 0.07; brain, liver and resting skeletal muscle ca. 0.3, myocardium 0.6. The E_{O_2} of muscle during strenuous exercise can rise to 0.9. Skeletal muscle can therefore meet increased O_2 demands by raising the $E_{O_2}(0.3 \Rightarrow 0.9)$, as can myocardial tissue to a much smaller extent (\rightarrow p. 210).

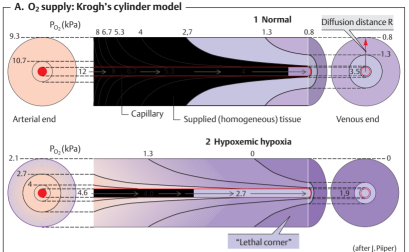
Hypoxia. An abnormally reduced O₂ supply to tissue is classified as follows:

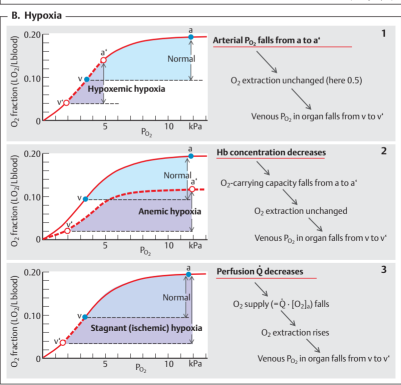
1. Hypoxic hypoxia (\rightarrow **A2, B1**): an insufficient O₂ supply reaches the blood due, for ex-

- ample, to decreased atmospheric P_{0_2} at high altitudes (\rightarrow p. 136), reduced alveolar ventilation, or impaired alveolar gas exchange.
- **2.** Anemic hypoxia (\rightarrow **B2**): reduced O₂-carrying capacity of blood (\rightarrow p. 128), e.g., due to decreased total Hb in iron deficiency anemia (\rightarrow p. 90).
- 3. Stagnant or ischemic hypoxia (\rightarrow B3): insufficient O_2 reaches the tissue due to reduced blood flow ($\dot{Q}\downarrow$). The cause can be systemic (e.g., heart failure) or local (e.g., obstructed artery). The reduction of blood flow must be compensated for by a rise in E_{O_2} to maintain an adequate O_2 delivery (see Eq. 5.7). This is not the case in hypoxic and anemic hypoxia. The influx and efflux of substrates and metabolites is also impaired in stagnant hypoxia. Anaerobic glycolysis (\rightarrow p. 72) is therefore of little help because neither the uptake of glucose nor the discharge of H⁺ ions dissociated from lactic acid is possible.
- **4.** Hypoxia can also occur when the diffusion distance is increased due to tissue thickening without a corresponding increase in the number of blood capilaries. This results in an insufficient blood supply to cells lying outside the O_2 supply radius (R) of the Krogh cylinder (\rightarrow A).
- **5.** Histotoxic or cytotoxic hypoxia occurs due to impaired utilization of O_2 by the tissues despite a sufficient supply of O_2 in the mitochondria, as observed in cyanide poisoning. Cyanide (HCN) blocks oxidative cellular metabolism by inhibiting cytochromoxidase.

Brain tissue is extremely susceptible to hypoxia, which can cause critical damage since dead nerve cells generally cannot be replaced. Anoxia, or a total lack of oxygen, can occur due to heart or respiratory failure. The cerebral survival time is thus the limiting factor for overall survival. Unconsciousness occurs after only 15 s of anoxia, and irreparable brain damage occurs if anoxia lasts for more than 3 min or so.

Cyanosis is a bluish discoloration of the skin, lips, nails, etc. due to excessive arterial deoxyhemoglobin ($>50\,\mathrm{g/L}$). Cyanosis is a sign of hypoxia in individuals with normal or only moderately reduced total Hb levels. When total Hb is extremely low, O_2 deficiencies (anemic hypoxia) can be life-threatening, even in the absence of cyanosis. Cyanosis can occur in absence of significant hypoxia when the Hb level is elevated.





Respiratory Control and Stimulation

The respiratory muscles (\rightarrow p. 108) are innervated by nerve fibers extending from the cervical and thoracic medulla (C4-C8 and T1-T7). The most important control centers are located in the medulla oblongata and cervical medulla (C1-C2), where interactive inspiratory and expiratory neurons on different levels $(\rightarrow A1$, red and green areas). The network of these spatially separate neuron groups form a rhythm generator (respiratory "center") where respiratory rhythm originates $(\rightarrow A1)$. The neuron groups are triggered alternately. resulting in rhythmic inspiration and expiration. They are activated in a tonic (nonrhythm-dependent) manner by the formatio reticularis, which receives signals from respiratory stimulants in the periphery and higher centers of the brain.

Respiratory sensors or receptors are involved in respiratory control circuits ($\rightarrow p.4$). Central and peripheral chemosensors on the medulla oblongata and in the arterial circulation continuously register gas partial pressures in cerebrospinal fluid (CSF) and blood, respectively, and mechanosensors in the chest wall respond to stretch of intercostal muscles to modulate the depth of breathing ($\rightarrow A2$). Pulmonary stretch sensors in the tracheal and bronchial walls respond to marked increases in lung volume, thereby limiting the depth of respiration in humans (Hering-Breuer reflex). Muscle spindles (\rightarrow p. 318) in the respiratory muscles also respond to changes in airway resistance in the lung and chest wall.

Chemical respiratory stimulants. The extent of involuntary ventilation is mainly determined by the partial pressures of O_2 and CO_2 and the pH of blood and CSF. Chemosensors respond to any changes in these variables. Peripheral chemosensors in the glomera aortica and carotica $(\rightarrow$ A3) register changes in the arterial P_{O_2} . If it falls, they stimulate an increase in ventilation via the vagus (X) and glossopharyngeal nerves (IX) until the arterial P_{O_2} rises again. This occurs, for example, at high altitudes $(\rightarrow$ p. 136). The impulse frequency of the sensors increases sharply when the P_{O_2} drops below 13 kPa or 97 mmHg (peripheral ventilatory drive). These changes are even

stronger when P_{CO2} and/or the H⁺ concentration in blood also increase.

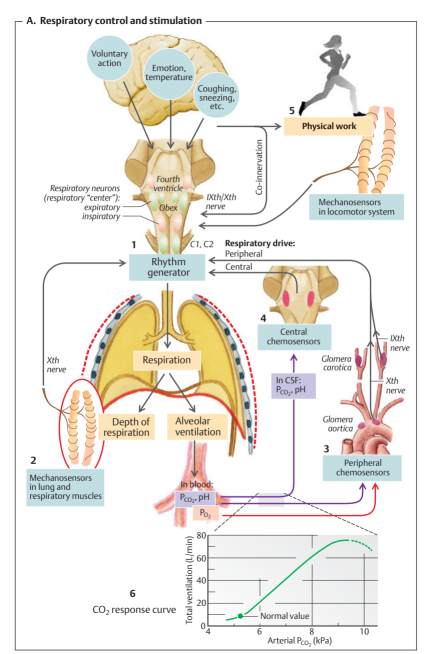
Central chemosensors in the medulla react to CO_2 and H^+ increases (= pH decrease) in the CSF (\rightarrow A4 and p. 126). Ventilation is then increased until P_{CO_2} and the H^+ concentration in blood and CSF decrease to normal values. This mostly **central respiratory drive** is very effective in responding to acute changes. An increase in arterial P_{CO_2} from, say, 5 to 9 kPa increases the total ventilation V_E by a factor of ten, as shown in the CO_2 response curve (\rightarrow A6).

When a **chronic rise in P**_{CO₂} occurs, the previously increased central respiratory drive decreases (\rightarrow p. 126). If O₂ supplied by artificial respiration tricks the peripheral chemosensors into believing that there is adequate ventilation, the residual peripheral respiratory drive will also be in jeopardy.

During **physical work** (\rightarrow **A5**), the total ventilation increases due to (a) co-innervation of the respiratory centers (by collaterals of cortical efferent motor fibers) and (b) through impulses transmitted by proprioceptive fibers from the muscles.

Non-feedback sensors and stimulants also play an important role in modulating the basic rhythm of respiration. They include

- Irritant sensors in the bronchial mucosa, which quickly respond to lung volume decreases by increasing the respiratory rate (deflation reflex or Head's reflex), and to dust particles or irritating gases by triggering the cough reflex.
- ♦ J sensors of free C fiber endings on alveolar and bronchial walls; these are stimulated in pulmonary edema, triggering symptoms such as apnea and lowering the blood pressure.
- Higher central nervous centers such as the cortex, limbic system, hypothalamus or pons. They are involved in the expression of emotions like fear, pain and joy; in reflexes such as sneezing, coughing, yawning and swallowing; and in voluntary control of respiration while speaking, singing, etc.
- ◆ Pressosensors (→ p. 214), which are responsible for increasing respiration when the blood pressure decreases.
- Heat and cold sensors in the skin and thermoregulatory center. Increases (fever) and decreases in body temperature lead to increased respiration.
- Certain hormones also help to regulate respiration.
 Progesterone, for example, increases respiration in the second half of the menstrual cycle and during pregnancy.



Effects of Diving on Respiration

Diving creates a problem for respiration due to the lack of normal ambient air supply and to higher the outside pressures exerted on the body. The total pressure on the body underwater is equal to the water pressure (98 kPa or 735 mmHg for each 10 m of water) plus the atmospheric pressure at the water surface.

A **snorkel** can be used when diving just below the water surface $(\rightarrow A)$, but it increases dead space $(\rightarrow pp. 114$ and 120), making it harder to breathe. The additional pressure load from the water on chest and abdomen must also be overcome with each breath.

The depth at which a snorkel can be used is limited 1) because an intolerable increase in dead space or airway resistance will occur when using an extremely long or narrow snorkel, and 2) because the water pressure at lower depths will prevent inhalation. The maximum suction produced on inspiration is about 11 kPa, equivalent to 112 cm H_2O (peak inspiratory pressure, \rightarrow p. 116). Inspiration therefore is no longer possible at aquatic depths of about 112 cm or more due to the risk of hypoxic anoxia (\rightarrow A).

Scuba diving equipment (scuba = self-contained underwater breathing upparatus) is needed to breathe at lower depths (up to about 70 m). The inspiratory air pressure (from pressurized air cylinders) is automatically adjusted to the water pressure, thereby permitting the diver to breathe with normal effort.

However, the additional water pressure increases the partial pressure of **nitrogen** P_{N_2} (\rightarrow **B**), resulting in higher concentrations of dissolved N2 in the blood. The pressure at a depth of 60 meters is about seven times higher than at the water surface. The pressure decreases as the diver returns to the water surface, but the additional N2 does not remain dissolved. The diver must therefore ascend slowly, in gradual stages so that the excess N2 can return to and be expelled from the lungs. Resurfacing too guickly would lead to the development of N2 bubbles in tissue (pain!) and blood, where they can cause obstruction and embolism of small blood vessels. This is called decompression sickness or caisson disease $(\rightarrow B)$. Euphoria (N2 narcosis?), also called rapture of the deep, can occur when diving at depths of over 40 to 60 meters. Oxygen toxicity can occur at depths of 75 m or more (\rightarrow p. 136).

When **diving unassisted**, i.e., simply by holding one's breath, P_{CO_2} in the blood rises, since the CO_2 produced by the body is not exhaled. Once a certain P_{CO_2} has been reached, chemosensors

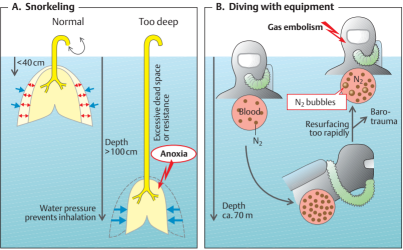
 $(\rightarrow p. 132)$ trigger a sensation of shortness of breath, signaling that it is time to resurface.

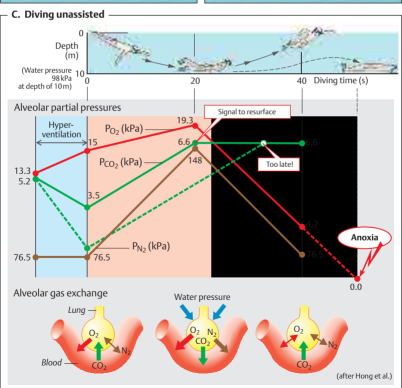
To delay the time to resurface, it is possible to lower the P_{CO₂} in blood by **hyperventilating before diving**. Experienced divers use this trick to stay under water longer. The course of alveolar partial pressures over time and the direction of alveolar gas exchange while diving (depth: 10 m; duration 40 s) is shown in C: Hyperventilating before a dive reduces the Pco, (solid green line) and slightly increases the Po, (red line) in the alveoli (and in blood). Diving at a depth of 10 m doubles the pressure on the chest and abdominal wall. As a result, the partial pressures of gases in the alveoli (PCO2, PO2, PN2) increase sharply. Increased quantities of O2 and CO2 therefore diffuse from the alveoli into the blood, once the P_{CO_2} in blood rises to a certain level, the body signals that it is time to resurface. If the diver resurfaces at this time, the Po, in the alveoli and blood drops rapidly (O2 consumption + pressure decrease) and the alveolar O2 exchange stops. Back at the water surface, the Po, reaches a level that is just tolerable. If the diver excessively hyperventilates before the dive, the signal to resurface will come too late, and the Po, will drop to zero (anoxia) before the person reaches the water surface, which can result in unconsciousness and drowning (\rightarrow **C**, dotted lines).

Barotrauma. The increased pressure associated with diving leads to compression of air-filled organs, such as the lung and middle ear. Their gas volumes are compressed to $^{1}/_{2}$ their normal size at water depths of 10 m, and to $^{1}/_{4}$ at depths of 30 m.

The missing volume of air in the lungs is automatically replaced by the scuba, but not that of the middle ear. The middle ear and throat are connected by the Eustachian tube, which is open only at certain times (e.g., when swallowing) or not at all (e.g., in pharyngitis). If volume loss in the ear is not compensated for during a dive, the increasing water pressure in the outer auditory canal distends the eardrum, causing pain or even eardrum rupture. As a result, cold water can enter the middle ear and impair the organ of equilibrium, leading to nausea, dizziness, and disorientation. This can be prevented by pressing air from the lungs into the middle ear by holding the nose and blowing with the mouth closed.

The air in air-filled organs expand when the diver ascends to the water surface. Resurfacing too quickly, i.e., without expelling air at regular intervals, can lead to complications such as lung laceration and pneumothorax (\rightarrow p. 110) as well as potentially fatal hemorrhage and air embolism.





Effects of High Altitude on Respiration

At sea level, the average barometric pressure $(P_B) \approx 101 \, \text{kPa} \ (760 \, \text{mmHg})$, the O_2 fraction in ambient air (F_{10_2}) is 0.209, and the inspiratory partial pressure of $O_2 \ (P_{10_2}) \approx 21 \, \text{kPa} \ (\rightarrow p. \, 106)$. However, P_B decreases with increasing altitude (h, in km):

 P_B (at h) = P_B (at sea level) $\cdot e^{-0.127 \cdot h}$ [5.9] This results in a drop in $Pl_{O_2} (\rightarrow A, column 1)$, alveolar P_{O_2} (PAO₂) and arterial P_{O_2} (PaO₂). The PA_{O_2} at sea level is about 13 kPa ($\rightarrow A$, column 2). PAO, is an important measure of oxygen supply. If the PAO, falls below a critical level (ca. 4.7 kPa = 35 mmHg), hypoxia ($\rightarrow \text{p.} 130$) and impairment of cerebral function will occur. The critical PAO2 would be reached at heights of about 4000 m above sea level during normal ventilation ($\rightarrow A$, dotted line in column 2). However, the low Pao, triggers chemosensors $(\rightarrow p. 132)$ that stimulate an increase in total ventilation $(\dot{\mathbf{V}}_{E})$; this is called \mathbf{O}_{2} deficiency **ventilation** (\rightarrow **A**, column 4). As a result, larger volumes of CO2 are exhaled, and the PACO2 and Pa_{CO2} decrease (see below). As described by the alveolar gas equation,

$$P_{A_{O_2}} = P_{I_{O_2}} - \frac{P_{A_{CO_2}}}{RQ}$$
 [5.10]

where RQ is the respiratory quotient (\rightarrow pp. 120 and 228), any fall in PA_{CO₂} will lead to a rise in the PA_{O₂}. O₂ deficiency ventilation stops the PA_{O₂} from becoming critical up to altitudes of about 7000 m (*altitude gain*, \rightarrow A).

The maximal increase in ventilation ($\approx 3 \times$ resting rate) during acute O2 deficiency is relatively small compared to the increase (≈ 10 times the resting rate) during strenuous physical exercise at normal altitudes (\rightarrow p. 74, C3) because increased ventilation at high altitudes reduces the Pa_{CO_2} (= hyperventilation, $\rightarrow p$. 108), resulting in the development of respiratory alkalosis (→ p. 144). Central chemosensors (\rightarrow p. 132) then emit signals to lower the respiratory drive, thereby counteracting the signals from O2 chemosensors to increase the respiratory drive. As the mountain climber adapts, respiratory alkalosis is compensated for by increased renal excretion of HCO3- $(\rightarrow p. 144)$. This helps return the pH of the blood toward normal, and the O2 deficiencyrelated increase in respiratory drive can now

prevail. Stimulation of O₂ chemosensors at high altitudes also leads to an increase in the heart rate and a corresponding *increase in cardiac output*, thereby increasing the O₂ supply to the tissues.

High altitude also stimulates **erythropoiesis** (\rightarrow p. 88ff.). Prolonged exposure to high altitudes increases the hematocrit levels, although this is limited by the corresponding rise in blood viscosity (\rightarrow pp. 92, 188).

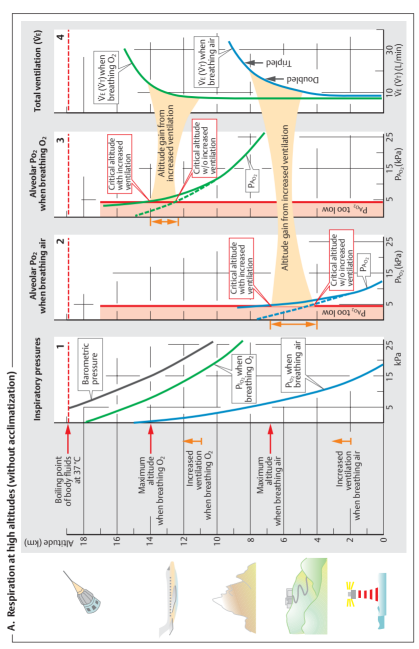
Breathing oxygen from pressurized O_2 cylinders is necessary for survival at altitudes above 7000 m, where P_{IO_2} is almost as high as the barometric pressure $P_B (\rightarrow A, \text{ column } 3)$. The critical P_{AO_2} level now occurs at an altitude of about 12 km with normal ventilation, and about 14 km with increased ventilation. Modern long-distance planes fly slightly below this altitude to ensure that the passengers can survive with an oxygen mask in case the cabin pressure drops unexpectedly.

Survival at altitudes above 14 km is not possible without pressurized chambers or pressurized suits like those used in space travel. Otherwise, the body fluids would begin to boil at altitudes of 20 km or so (\rightarrow **A**), where P_B is lower than water vapor pressure at body temperature (37 °C).

Oxygen Toxicity

Hyperoxia occurs when Pio, is above normal (> 22 kPa or 165 mmHg) due to an increased O₂ fraction (oxygen therapy) or to an overall pressure increase with a normal O2 fraction (e.g. in diving, \rightarrow p. 134). The degree of O₂ toxicity depends on the P102 level (critical: ca. 40 kPa or 300 mmHg) and duration of hyperoxia. Lung dysfunction (→ p. 118, surfactant deficiency) occurs when a PiO2 of about 70 kPa (525 mmHg) persists for several days or 200 kPa (1500 mmHg) for 3-6 hours. Lung dysfunction initially manifests as coughing and painful breathing. Seizures and unconsciousness occur at PiO2 levels above 220 kPa (1650 mmHg), e.g., when diving at a depth of about 100 m using pressurized air.

Newborns will go blind if exposed to P_{10_2} levels much greater than 40 kPa (300 mmHg) for long periods of time (e.g., in an incubator), because the vitreous body then opacifies.



pH, pH Buffers, Acid-Base Balance

The **pH** indicates the hydrogen ion activity or the "effective" H^+ concentration of a solution $(H^+$ activity = $f_H \cdot [H^+]$, where square brackets mean concentration; $\rightarrow p.378$), where

$$pH = -\log(f_H \cdot [\mathbf{H}^+])$$
 [6.1]

In healthy individuals, the **pH of the blood** is usually a mean pH **7.4** (see p. 142 for normal range), corresponding to H⁺ activity of about 40 nmol/L. The maintenance of a constant pH is important for human survival. Large deviations from the norm can have detrimental effects on metabolism, membrane permeability, and electrolyte distribution. Blood pH values below 7.0 and above 7.8 are not compatible with life.

Various **pH buffers** are responsible for maintaining the body at a constant pH (\rightarrow p. 379). One important buffer for blood and other body fluids is the **bicarbonate/carbon dioxide** (HCO₃-/CO₂) buffer system:

$$CO_2 + H_2O HCO_3^- + H^+.$$
 [6.2]

The pK_a value (\rightarrow p. 378f.) determines the prevailing concentration ratio of the buffer base and buffer acid ($[HCO_3^-]$ and $[CO_2]$, respectively in Eq. 6.2) at a given pH (Henderson–Hasselbalch equation; \rightarrow A).

The primary function of the CO_2/HCO_3 -buffer system in blood is to buffer H^+ ions. However, this system is especially important because the concentrations of the two buffer components can be modified largely independent of each other: $[CO_2]$ by respiration and $[HCO_3^-]$ by the liver and kidney (\rightarrow **A**; see also p. 174). It is therefore classified as an **open buffer system** (\rightarrow p. 140).

Hemoglobin in red blood cells (320 g Hb/L erythrocytes! → MCHC, p. 89 C), the second most important buffer in blood, is a **non-bicar-bonate buffer**.

HbH Hb
$$^-$$
 + H $^+$ [6.3]

Oxy-HbH Oxy-Hb
$$^-$$
 + H $^+$ [6.4]

The relatively acidic oxyhemoglobin anion (Oxy-Hb⁻) combines with fewer H⁺ ions than deoxygenated Hb⁻, which is less acidic (see also p. 124). H⁺ ions are therefore liberated upon oxygenation of Hb to Oxy-Hb in the lung. Reaction 6.2 therefore proceeds to the left, thereby promoting the release of CO₂ from its

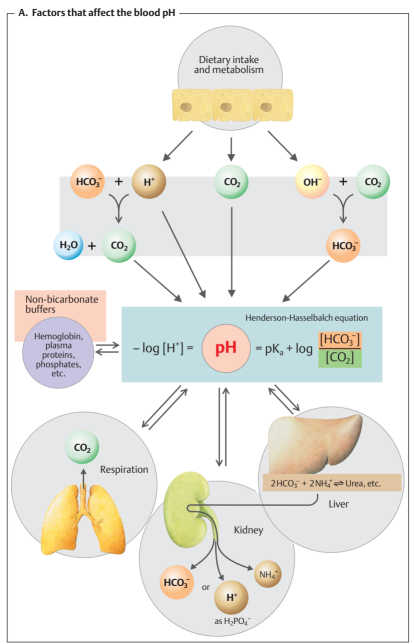
bound forms (\rightarrow p. 124). This, in turn, increases the pulmonary elimination of CO₂.

Other non-bicarbonate buffers of the blood include plasma proteins and inorganic phosphate $(H_2PO_4^- H^+ + HPO_4^{-2})$ as well as organic phosphates (in red blood cells). Intracellular organic and inorganic substances in various tissues also function as buffers.

The **buffer capacity** is a measure of the buffering power of a buffer system (mol·L-1 · $[\Delta pH]^{-1}$). It corresponds to the number of added H⁺ or OH⁻ ions per unit volume that change the pH by one unit. The buffer capacity therefore corresponds to the slope of the titration curve for a given buffer (\rightarrow p. 380, B). The buffer capacity is dependent on (a) the buffer concentration and (b) the pH. The farther the pH is from the pK_a of a buffer system, the smaller the buffer capacity ($\rightarrow p.380$). The buffer capacity of the blood is about 75 mmol·L⁻¹·(Δ pH)⁻¹ at pH 7.4 and constant Pco₂. Since the buffer capacity is dependent on the prevailing Pco, the buffer base concentration of the blood (normally about 48 mEq/L) is normally used as the measure of buffering power of the blood in clinical medicine $(\rightarrow pp. 142 \text{ and } 146)$. The buffer base concentration is the sum of the concentrations of all buffer components that accept hydrogen ions, i.e., HCO₃-, Hb-, Oxy-Hb-, diphosphoglycerate anions, plasma protein anions, HPO₄²⁻, etc.

Changes in the pH of the blood are chiefly due to changes in the following factors (\rightarrow A and p. 142ff.):

- H^+ ions: Direct uptake in foodstuffs (e.g., vinegar) or by metabolism, or removal from the blood (e.g., by the kidney; \rightarrow p. 174 ff.).
- OH⁻ ions: Uptake in foodstuffs containing (basic) salts of weak acids, especially in primarily vegetarian diet.
- ◆ CO₂: Its concentration, [CO₂], can change due to alterations in metabolic production or pulmonary elimination of CO₂. A drop in [CO₂] leads to a rise in pH and vice versa (→ A: [CO₂] is the denominator in the equation).
- ◆ HCO₃⁻: It can be eliminated directly from the blood by the kidney or gut (in diarrhea) (→ pp. 176, 142). A rise or fall in [HCO₃⁻] will lead to a corresponding rise or fall in pH (→ A: [HCO₃⁻] is the numerator in the equation).



Bicarbonate/Carbon Dioxide Buffer

The pH of any buffer system is determined by the concentration ratio of the buffer pairs and the pK_a of the system (\rightarrow p. 378). The pH of a bicarbonate solution is the concentration ratio of bicarbonate and dissolved carbon dioxide ([HCO₃-]/[CO₂]), as defined in the Henderson–Hasselbalch equation (\rightarrow A1). Given [HCO₃-] = 24 mmol/L and [CO₂] = 1.2 mmol/l, [HCO₃-]/[CO₂] = 24/1.2 = 20. Given log20 = 1.3 and pK_a = 6.1, a pH of 7.4 is derived when these values are set into the equation (\rightarrow A2). If [HCO₃-] drops to 10 and [CO₂] decreases to 0.5 mmol/L, the ratio of the two variables will not change, and the pH will remain constant.

When added to a buffered solution, H+ ions combine with the buffer base (HCO₃- in this case), resulting in the formation of buffer acid $(HCO_3^- + H^+ \rightarrow CO_2 + H_2O)$. In a closed system from which CO_2 cannot escape ($\rightarrow A3$), the amount of buffer acid formed (CO2) equals the amount of buffer base consumed (HCO₃-). The inverse holds true for the addition of hydroxide ions (OH⁻ + CO₂ \rightarrow HCO₃⁻). After addition of 2 mmol/L of H+, the aforementioned baseline ratio $[HCO_3^-]/[CO_2]$ of $24/1.2 (\rightarrow A2)$ changes to 22/3.2, making the pH fall to 6.93 $(\rightarrow A3)$. Thus, the buffer capacity of the HCO₃⁻/ CO₂ buffer at pH 7.4 is very low in a closed system, for which the pKa of 6.1 is too far from the target pH of 7.4 (\rightarrow pp. 138, 378 ff).

If, however, the additionally produced CO₂ is eliminated from the system (open system; → A4), only the [HCO₃-] will change when the same amount of H+ is added (2 mmol/L). The corresponding decrease in the [HCO₃-]/[CO₂] ratio (22/1.2) and pH (7.36) is much less than in a closed system. In the body, bicarbonate buffering occurs in an open system in which the partial pressure (PCO2) and hence the concentration of carbon dioxide in plasma ($[CO_2]$ = $\alpha \cdot P_{CO_2}$; \rightarrow p. 126) are regulated by respiration $(\rightarrow B)$. The lungs normally eliminate as much CO₂ as produced by metabolism (15 000-20 000 mmol/day), while the alveolar Pco, remains constant (\rightarrow p. 120ff.). Since the plasma P_{CO2} adapts to the alveolar P_{CO2} during each respiratory cycle, the arterial PCO2 (PaCO2) also remains constant. An increased supply of H+ in the periphery leads to an increase in the P_{CO_2} of

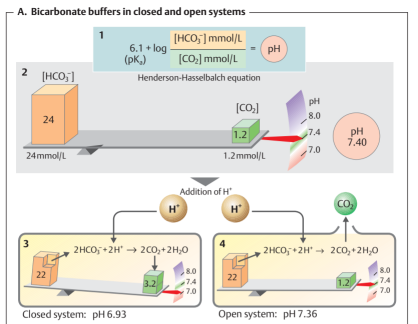
venous blood (H⁺+HCO₃⁻ \rightarrow CO₂+H₂O)(\rightarrow **B1**). The lungs eliminate the additional CO₂ so quickly that the arterial P_{CO_2} remains practically unchanged despite the addition of H⁺ (open system!).

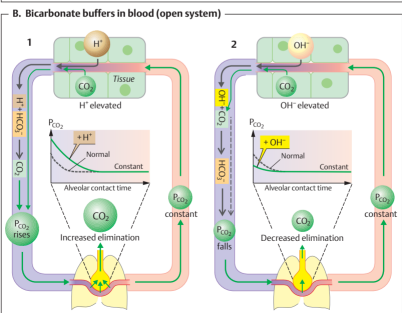
The following example demonstrates the quantitatively small impact of increased **pulmonary CO₂ elimination**. A two-fold increase in the amount of H⁺ ions produced within the body on a given day (normally 60 mmol/day) will result in the added production of 60 mmol more of CO₂ per day (disregarding non-bicarbonate buffers). This corresponds to only about 0.3% of the normal daily CO₂ elimination rate.

An increased supply of OH^- ions in the periphery has basically similar effects. Since $OH^- + CO_2 \rightarrow HCO_3^-$, $[HCO_3^-]$ increases and the venous P_{CO_2} becomes smaller than normal. Because the rate of CO_2 elimination is also reduced, the arterial P_{CO_2} also does not change in the illustrated example $(\rightarrow B2)$.

At a pH of 7.4, the open HCO₃ $^-$ /CO₂ buffer system makes up about two-thirds of the buffer capacity of the blood when the P_{CO_2} remains constant at 5.33 kPa (\rightarrow p. 138). Mainly intracellular non-bicarbonate buffers provide the remaining buffer capacity.

Since **non-bicarbonate buffers** (NBBs) function in *closed systems*, their total concentration ([NBB base] + [NBB acid]) remains constant, even after buffering. The total concentration changes in response to changes in the hemoglobin concentration, however, since hemoglobin is the main constituent of NBBs (→ pp. 138, 146). NBBs supplement the HCO₃-/CO₂ buffer in non-respiratory (metabolic) acid-base disturbances (→ p. 142), but are the *only* effective buffers in respiratory acid-base disturbances (→ p. 144).





Acidosis and Alkalosis

The main objective of acid-base regulation is to keep the pH of blood, and thus of the body, constant. The normal ranges for parameters relevant to acid-base homeostasis, as measured in plasma (arterialized capillary blood) are listed in the table (see table on p. 124 for erythrocyte P_{CO2} and [HCO3⁻] values).

Normal range of acid-base parameters in plasma

	Women	Men
[H ⁺] (nmol/L)	39.8 ± 1.4	40.7 ± 1.4
pН	7.40 ± 0.015	7.39 ± 0.015
P _{CO2} (kPa)	5.07 ± 0.3	5.47 ± 0.3
(mmHg)	38.9 ± 2.3	41.0 ± 2.3
[HCO ₃ -] (mmol/L)	24 ± 2.5	24 ± 2.5

Acid-base homeostasis exists when the following balances are maintained:

- (H⁺ addition or production) (HCO₃⁻ addition or production) = (H⁺ excretion) (HCO₃⁻ excretion) ≈ 60 mmol/day (diet-dependent).
- 2. (CO₂ production) = (CO₂ excretion) $\approx 15\,000-20\,000\,\text{mmol/day}$.

H⁺ production (HCl, H₂SO₄, lactic acid, H₃PO₄, etc.) and adequate renal H⁺ excretion (→ p. 174ff.) are the main factors that influence the first balance. A vegetarian diet can lead to a considerable addition of HCO₃⁻ (metabolism: OH⁻ + CO₂ → HCO₃⁻; → p. 138). HCO₃⁻ is excreted in the urine to compensate for the added supply (the urine of vegetarians therefore tends to be alkaline).

Acid-base disturbances. Alkalosis occurs when the pH of the blood rises above the normal range (see table), and acidosis occurs when it falls below the lower limits of normal. Respiratory acid-base disturbances occur due to primary changes in $P_{CO_2}(\rightarrow p. 144)$, whereas non-respiratory (metabolic) disturbances occur due to a primary change in [HCO₃-]. Acid-base disturbances can be partially or almost completely compensated.

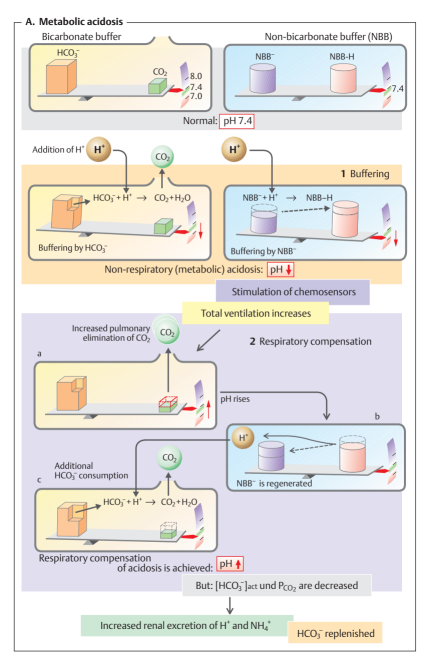
Nonrespiratory (Metabolic) Acid-Base Disturbances

Nonrespiratory acidosis is most commonly caused by (1) renal failure or isolated renal tubular H⁺ secretion defect resulting in inability to eliminate normal quantities of H+ ions (renal acidosis); (2) hyperkalemia (\rightarrow p. 180); (3) increased β-hydroxybutyric acid and acetoacetic acid production (diabetes mellitus. starvation): (4) increased anaerobic conversion of glucose to lactic acid (\rightarrow lactate⁻ + H⁺). e.g., due to strenuous physical work (\rightarrow p. 74) or hypoxia; (5) increased metabolic production of HCl and H2SO4 in individuals with a high intake of dietary proteins; and (6) loss of HCO₃through renal excretion (proximal renal tubular acidosis, use of carbonic anhydrase inhibitors) or diarrhea.

Buffering (\rightarrow A1) of excess hydrogen ions occurs in the first stage of non-respiratory acidosis (every HCO₃⁻ lost results in an H⁺ gained). Two-thirds and one-third of the buffering is achieved by HCO₃⁻ and non-bicarbonate buffer bases (NBB⁻), respectively, and the CO₂ arising from HCO₃⁻ buffering is eliminated from the body by the lungs (open system; \rightarrow p. 140). The standard bicarbonate concentration [HCO₃⁻]_{St}, the actual bicarbonate concentration [HCO₃⁻]_{Act} and the buffer base concentration [BB] *decrease* (negative base excess; \rightarrow p. 146).

Respiratory compensation of non-respiratory acidosis $(\rightarrow A2)$ occurs in the second stage. The total ventilation rises in response to the reduced pH levels (via central chemosensors), leading to a decrease in the alveolar and arterial P_{CO_2} (hyperventilation; $\rightarrow A2a$). This not only helps to return the [HCO3-]/[CO2] ratio towards normal (20:1), but also converts NBB-H back to NBB- (due to the increasing pH) (→ A2b). The latter process also requires HCO₃⁻ and, thus, further compensatory pulmonary elimination of CO_2 (\rightarrow **A2c**). If the cause of acidosis persists, respiratory compensation will eventually become insufficient, and increased renal excretion of H+ ions will occur $(\rightarrow p. 174ff.)$, provided that the acidosis is not of renal origin (see above, cause 1).

Nonrespiratory (metabolic) alkalosis is caused by (1) the administration of bases (e.g., HCO_3 - infusion); (2) increased breakdown of



organic anions (e.g., lactate⁻, α -ketoglutarate²⁻); (3) loss of H⁺ ions due to vomiting (\rightarrow p.238) or hypokalemia; and (4) volume depletion. **Buffering** in metabolic alkalosis is similar to that of non-respiratory acidosis (rise in [HCO₃⁻]_{St}, positive base excess). Nonetheless, the capacity for **respiratory compensation** through hypoventilation is very limited because of the resulting O₂ deficit.

Respiratory Acid-Base Disturbances

Respiratory alkalosis $(\rightarrow B)$ occurs when the lungs eliminate more CO₂ than is produced by metabolism (hyperventilation), resulting in a decrease in plasma P_{CO2} (hypocapnia). Inversely, respiratory acidosis occurs $(\rightarrow B)$ when less CO2 is eliminated than produced (hypoventilation), resulting in an increase in plasma Pco, (hypercapnia). Whereas bicarbonate and non-bicarbonate buffer bases (NBB-) jointly buffer the pH decrease in metabolic acidosis (\rightarrow p. 142), the two buffer systems behave very differently in respiratory alkalosis (\rightarrow **B1**). In the latter case, the HCO₃⁻/ CO₂ system is not effective because the change in P_{CO2} is the primary cause, not the result of respiratory alkalosis.

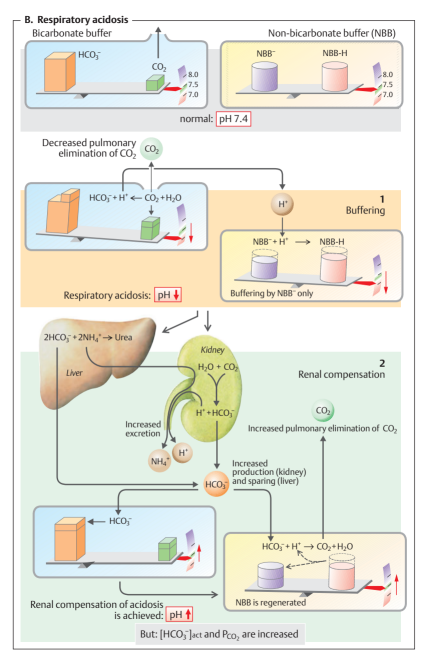
Respiratory acidosis can occur as the result of lung tissue damage (e.g., tuberculosis), impairment of alveolar gas exchange (e.g., pulmonary edema), paralysis of respiratory muscles (e.g., polio), insufficient respiratory drive (e.g., narcotic overdose), reduced chest motility (e.g., extreme spinal curvature), and many other conditions. The resulting increase in plasma CO_2 ($[CO_2] = \alpha \cdot P_{CO_2}$) is followed by increased HCO_3^- and H^+ production (\rightarrow **B1**, left panel). The H⁺ ions are buffered by NBB bases $(NBB^- + H^+ \rightarrow NBB-H; \rightarrow B1, right panel)$ while [HCO₃-]_{Act} increases. Unlike non-respiratory acidosis, [HCO₃-]_{St} remains unchanged (at least initially since it is defined for normal P_{CO2}; → p. 146) and [BB] remains unchanged because the [NBB⁻] decrease equals the [HCO₃⁻]_{Act} increase. Since the percentage increase in [HCO₃-]_{Act} is much lower than the rise in [CO₂], the [HCO₃-]/[CO₂] ratio and pH are lower than normal (acidosis).

If the increased P_{CO_2} persists, **renal compensation** (\rightarrow **B2**) of the respiratory disturbance will occur. The kidneys begin to excrete in-

creased quantities of H+ in form of titratable acidity (\rightarrow p. 174f.) of NH₄⁺ as well and, after a latency period of 1 to 2 days. Each NH₄⁺ ion excreted results in the sparing of one HCO₃- ion in the liver, and each H+ ion excreted results in the tubular cellular release of one HCO3- ion into the blood (\rightarrow p. 174ff.). This process continues until the pH has been reasonably normalized despite the Pcoa increase. A portion of the HCO3- is used to buffer the H+ ions liberated during the reaction NBB-H → NBB- + H+ $(\rightarrow B2, right panel)$. Because of the relatively long latency for renal compensation, the drop in pH is more pronounced in acute respiratory acidosis than in chronic respiratory acidosis. In the chronic form, [HCO₃-]_{Act} can rise by about 1 mmol per 1.34 kPa (10 mmHg) increase in P_{CO_2} .

Respiratory alkalosis is usually caused by hyperventilation due to anxiety or high altitude (oxygen deficit ventilation: $\rightarrow p$, 136), resulting in a fall in plasma P_{CO_2} . This leads to a slight decrease in [HCO3-]Act since a small portion of the HCO3- is converted to CO2 (H++ $HCO_3^- \rightarrow CO_2 + H_2O$); the HCO_3^- required for this reaction is supplied by H+ ions from NBB's (**buffering**: NBB-H \rightarrow NBB⁻ + H⁺). This is also the reason for the additional drop in [HCO₃-]_{Act} when respiratory compensation of non-respiratory acidosis occurs (\rightarrow p. 143 A, bottom panel, and p. 146). Further reduction of [HCO₃-]_{Act} is required for adequate pH normalization (compensation). This is achieved through reduced renal tubular secretion of H+. As a consequence, increased renal excretion of HCO₃⁻ will occur (renal compensation).

In acute respiratory acidosis or alkalosis, CO_2 diffuses more rapidly than HCO_3^- and H^+ from the blood into the cerebrospinal fluid (CSF). The low NBB concentrations there causes relatively strong fluctuations in the pH of the CSF (\rightarrow p. 126), providing an adequate stimulus for central chemosensors (\rightarrow p. 132).



Assessment of Acid-Base Status

The Henderson–Hasselbalch equation for the HCO₃⁻/CO₂ buffer system states:

 $pH = pK_a + log ([HCO_3^-]/[CO_2]). \qquad [6.5] \\ \text{Since } [CO_2] = \alpha \cdot P_{CO_2} \ (\rightarrow p. 126), \ \text{Equation } 6.5 \\ \text{contains two constants } (pK_a \text{ and } \alpha) \ \text{and three} \\ \text{variables } (pH, [HCO_3^-], \text{ and } P_{CO_2}). \ \text{At } 37 \,^{\circ}\text{C in} \\ \text{plasma, } pK_a = 6.1 \ \text{and } \alpha = 0.225 \ \text{mmol} \cdot L^{-1} \cdot kPa^{-1} \ (\text{cf. p. } 126). \ \text{When one of the variables remains constant } (e.g., [HCO_3^-]), \ \text{the other two} \\ \text{(e.g., } P_{CO_2} \ \text{ and } pH) \ \text{ are interdependent. In a graphic representation, this dependency is reflected as a straight line when the logarithm of P_{CO_2} is plotted against the pH <math>(\rightarrow \textbf{A-C})$ and p. 382).

When the P_{CO_2} varies in a **bicarbonate solution** (without other buffers), the pH changes but $[HCO_3^-]$ remains constant (\rightarrow A, solid line). One can also plot the lines for different HCO_3^- concentrations, all of which are parallel (\rightarrow A, B, dotted orange lines). Figures A through C use a scale that ensures that the bicarbonate lines intersect the coordinates at 45° angles. The **Siggaard–Andersen nomogram** (\rightarrow C) does not use the lines, but only the points of intersection of the $[HCO_3^-]$ lines with the normal P_{CO_2} of 5.33 kPa (40 mmHq).

The **blood** contains not only the HCO₃⁻/CO₂ buffer but also non-bicarbonate buffers, NBB $(\rightarrow p. 138)$. Thus, a change in the P_{CO_2} does not alter the pH as much as in a solution containing the HCO_3^-/CO_2 buffer alone (\rightarrow p. 144). In the P_{CO},/pH nomogram, the slope is therefore steeper than 45° (\rightarrow **B**, green and red lines). Hence, the actual bicarbonate concentration, [HCO₃-]_{Act}, in blood changes and shifts in the same direction as the PCO2 if the pH varies $(\rightarrow p. 144)$. Therefore, both the $[HCO_3^-]_{Act}$ and the standard bicarbonate concentration, [HCO₃⁻]_{St}, can be determined in clinical blood tests. By definition, [HCO₃-]_{St} represents the $[HCO_3^-]$ at a normal P_{CO_2} of 5.33 kPa (40 mmHg). [HCO₃-|_{St} therefore permits an assessment of [HCO₃-] independent of P_{CO₂} changes.

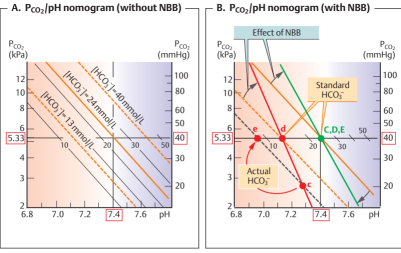
[HCO₃-]_{St} and [HCO₃-]_{Act} are determined using measured P_{CO_2} and pH values obtained with a blood gas analyzer. When plotted on the Siggaard–Andersen nomogram, [HCO₃-]_{St} is read from the line as indicated by the points of intersect of the [HCO₃-] line (\rightarrow B, orange lines) and the P_{CO_3} /pH line (B and C, green and red

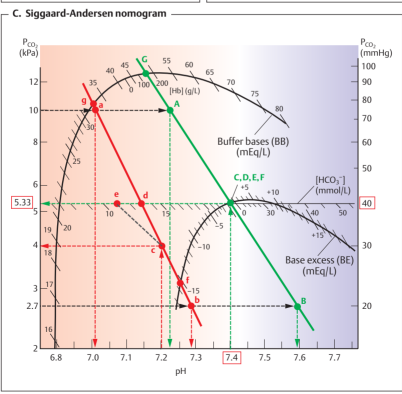
lines) at normal P_{CO_2} = 5.33 (\rightarrow **B** and **C**, points D and d). [HCO₃⁻]_{Act} is read from the [HCO₃⁻] line intersected by the P_{CO_2} /pH line at the level of the actually measured P_{CO_2} . Since the normal and measured P_{CO_2} values agree in normals, their [HCO₃⁻]_{Act} is usually equal to [HCO₃⁻]_{St}. If P_{CO_2} deviates from normal (\rightarrow **B**, **C**, point c), [HCO₃⁻]_{Act} is read at point e on the HCO₃⁻ line (\rightarrow **B**, **C**, interrupted 45° line) on which the actually measured P_{CO_2} lies (\rightarrow **B**, **C**, point c).

Blood Pco, and pH measurement. When using the equilibration method (Astrup method), three pH measurements are taken: (1) in the unchanged blood sample; (2) after equilibration with a high Pco2 (e.g., 10 kPa [75 mmHg]; \rightarrow **C**, points A and a), and (3) after equilibration with a low Pco, (e.g., 2.7 kPa [20 mmHg]; \rightarrow **C**, points *B* and *b*). The P_{CO2} of the original blood sample can then be read from lines A-B and a-b using the pH value obtained in measurement 1. In normals $(\rightarrow C$, upper case letters, green), $[HCO_3^-]_{Act} =$ $[HCO_3^-]_{St} = 24 \text{ mmol/L } (\rightarrow \mathbb{C}, \text{ points } E \text{ and } D).$ Example 2 (\rightarrow **C**, lower case letters, red) shows an acid-base disturbance: The pH is too low (7.2) and $[HCO_3^-]_{St}$ (\rightarrow **C**, point *d*) has dropped to 13 mmol/L (metabolic acidosis). This has been partially compensated (\rightarrow p. 142) by a reduction in Pco2 to 4 kPa, which led to a consequent reduction in [HCO₃-]_{Act} to 11 mmol/L $(\rightarrow \mathbf{C}, point e)$.

Total **buffer bases** (BB) and **base excess** (BE) (\rightarrow p. 142) can also be read from the Siggaard–Andersen nomogram (\rightarrow C). The base excess (points F and f on the curve) is the difference between the measured buffer base value (points *G* or *g*) and the normal buffer base value (point *G*). Point *G* is dependent on the hemoglobin concentration of the blood (\rightarrow C; [Hb]/BB comparison). Like [HCO₃⁻]_{St}, deviation of BB from the norm (0 \pm 2.5 mEq/L) is diagnostic of primary non-respiratory acid–base disturbances.

The P_{CO_2}/pH line of the blood sample in plate ${\bf C}$ can also be determined if (1) the P_{CO_2} (without equilibration), (2) the pH, and (3) the hemoglobin concentration are known. One point (\rightarrow ${\bf C}$, point c) on the unknown line can be drawn using (1) and (2). The line must be drawn through the point in such a way that BB (point g) – BB_{normal} (dependent on Hb value) = BE (point f).





Kidneys, Salt, and Water Balance

Kidney Structure and Function

Three fundamental mechanisms characterize kidney function: (1) large quantities of water and solutes are **filtered** from the blood. (2) This *primary urine* enters the tubule, where most of it is **reabsorbed**, i.e., it exits the tubule and passes back into the blood. (3) Certain substances (e.g., toxins) are not only not reabsorbed but actively **secreted** into the tubule lumen. The non-reabsorbed residual filtrate is **excreted** together with the secreted substances in the *final urine*.

Functions: The kidneys (1) adjust salt and water excretion to maintain a constant *extracellular fluid volume* and *osmolality*; (2) they help to maintain *acid-base homeostasis*; (3) they *eliminate end-products* of metabolism and foreign substances while (4) preserving useful compounds (e.g., glucose) by reabsorption; (5) the produce *hormones* (e.g., erythropoietin) and hormone activators (renin), and (6) have *metabolic functions* (protein and peptide catabolism, gluconeogenesis, etc.).

Nephron Structure

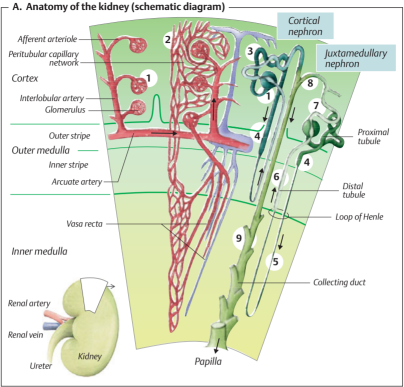
Each kidney contains about 10^6 nephrons, each consisting of the malpighian body and the tubule. The malpighian body is located in the renal cortex $(\rightarrow A)$ and consists of a tuft of capillaries (glomerulus) surrounded by a double-walled capsule (Bowman's capsule). The primary urine accumulates in the capsular space between its two layers $(\rightarrow B)$. Blood enters the glomerulus by an afferent arteriole (vas afferens) and exits via an efferent arteriole (vas efferens) from which the peritubular capillary network arises $(\rightarrow p. 150)$. The glomerular filter $(\rightarrow B)$ separates the blood side from the Bowman's capsular space.

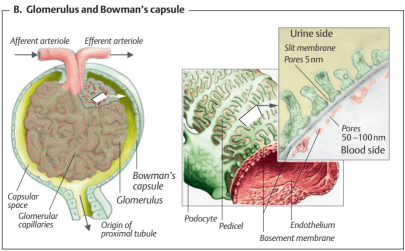
The **glomerular filter** comprises the fenestrated endothelium of the glomerular capillaries (50–100 nm pore size) followed by the basal membrane as the second layer and the visceral membrane of Bowman's capsule on the urine side. The latter is formed by *podocytes* with numerous interdigitating footlike processes (*pedicels*). The slit-like spaces between them are covered by the *slit membrane*, the *pores* of which are about 5 nm in diameter. They are shaped by the protein *nephrine*, which is anchored to the cytoskeleton of the podocytes.

- ♦ The **proximal tubule** (\rightarrow **A**, dark green) is the longest part of a nephron (ca. 10 mm). Its twisted initial segment (*proximal convoluted tubule*, *PCT*; \rightarrow **A3**) merges into a straight part, PST (*pars recta*; \rightarrow **A4**).
- ♦ The **loop of Henle** consists of a *thick descending limb* that extends into the renal medulla (\rightarrow **A4** = PST), a *thin descending limb* (\rightarrow **A5**), a *thin ascending limb* (only in juxtamedullary nephrons which have long loops), and a *thick ascending limb*, TAL (\rightarrow **A6**). It contains the **macula densa** (\rightarrow p. 184), a group of specialized cells that closely communicate with the glomerulus of the respective nephron. Only about 20% of all Henle's loops (those of the deep juxtamedullary nephrons) are long enough to penetrate into the inner medulla. Cortical nephrons have shorter loops (\rightarrow **A** and p. 150).
- The distal tubule (→ A, grayish green) has an initially straight part (=TAL of Henle's loop; → A6) that merges with a convoluted part (distal convoluted tubule, DCT; → A7).

The DCT merges with a connecting tubule $(\rightarrow A8)$. Many of them lead into a **collecting duct**, **CD** $(\rightarrow A9)$ which extends through the renal cortex (cortical CD) and medulla (medullary CD). At the *renal papilla* the collecting ducts opens in the *renal pelvis*. From there, the urine (propelled by peristaltic contractions) passes via the *ureter* into the *urinary bladder* and, finally, into the *urethra*, through which the urine exits the body.

Micturition. Voiding of the bladder is controlled by reflexes. Filling of the bladder activates the smooth detrusor muscle of the bladder wall via stretch sensors and parasympathetic neurons (S_2 – S_4 , \rightarrow p. 78 ff.). At low filling volumes, the wall relaxes via sympathetic neurons (L_1 – L_2) controlled by supraspinal centers (pons). At higher filling volumes (> 0.3 L), the threshold pressure (about 1 kPa) that triggers the *micturition reflex* via a positive feedback loop is reached: The detrusor muscle contracts \rightarrow pressure \uparrow \rightarrow contraction $\uparrow \uparrow$ and so on until the *internal* (smooth m.) and *external sphincter* (striated m.) open so the urine can exit the body.





150

Renal Circulation

The arcuate arteries $(\rightarrow A1)$ pass between the renal cortex and medulla. They branch towards the cortex into the interlobular arteries $(\rightarrow A2)$ from which the afferent arterioles (or vasa afferentia) arise (\rightarrow A3). Unlike other organs, the kidney has two successive capillary networks that are connected with each other by an efferent arteriole (or vas efferens) (\rightarrow **A. B**). Pressure in the first network of glomerular capillaries $(\rightarrow p. 148)$ is a relatively high $(\rightarrow B \text{ and } p. 152)$ and is regulated by adjusting the width of interlobular artery, the afferent and/or efferent arterioles ($\rightarrow A$ 3.4). The second network of **peritubular capillaries** $(\rightarrow A)$ winds around the cortical tubules. It supplies the tubule cells with blood, but it also contributes the to exchange of substances with the tubule lumen (reabsorption, secretion; \rightarrow p. 154 ff.).

The **renal blood flow** (RBF) is relatively high, ca. 1.2 L/min, equivalent to 20-25% of the cardiac output. This is required to maintain a high glomerular filtration rate (GFR; \rightarrow p. 152) and results in a very low arteriovenous O_2 difference (ca. 15 mL/L of blood). In the *renal cortex*, O_2 is **consumed** (ca. 18 mL/min) for oxidative **metabolism** of fatty acids, etc. Most of the ATP produced in the process is used to fuel active transport. In the renal medulla, metabolism is mainly anaerobic (\rightarrow p. 72).

Around 90% of the renal blood supply goes to the cortex. Per gram of tissue, approximately 5, 1.75 and 0.5 mL/min of blood pass through the cortex, external medulla, and internal medulla, respectively. The latter value is still higher than in most organs $(\rightarrow p. 213 \text{ A})$.

The kidney contains **two types of nephrons** that differ with respect to the features of their second network of capillaries $(\rightarrow A)$.

◆ Cortical nephrons are supplied by peritubular capillaries (see above) and have short loops of Henle. ◆ Juxtamedullary nephrons are located near the cortex-medulla junction. Their efferent arterioles give rise to relatively long (≤ 40 mm), straight arterioles (vasa recta) that descend into the renal medulla. The vasa recta supply the renal medulla and can accompany long loops of Henle of juxtamedulary nephrons as far as the tip of the renal papilla (→ p. 148). Their hairpin shape is important for the concentration of urine (→ p. 164ff.).

Any **change in blood distribution** to these two types of nephrons affects NaCl excretion. Antidiuretic hormone (ADH) increases the GFR of the juxtamedullary nephrons.

Due to autoregulation of renal blood flow, only slight changes in renal plasma flow (RPF) and glomerular filtration rate (GFR) occur (even in a denervated kidney) when the systemic blood pressure fluctuates between 80 and about 180 mmHg (\rightarrow C). Resistance in the interlobular arteries and afferent arterioles located upstream to the cortical glomeruli is automatically adjusted when the mean blood pressure changes $(\rightarrow B, C)$. If the blood pressure falls below about 80 mmHg, however, renal circulation and filtration will ultimately fail $(\rightarrow C)$. RBF and GFR can also be regulated independently by making isolated changes in the (serial) resistances of the afferent and efferent arterioles (\rightarrow p. 152).

Non-invasive **determination of RBF** is possible if the **renal plasma flow** (**RPF**) is known (normally about **0.6 L/min**). RPF is obtained by measuring the amount balance (Fick's principle) of an intravenously injected test substance (e.g., p-aminohippurate, **PAH**) that is almost completely eliminated in the urine during one renal pass (PAH is filtered and highly secreted, \rightarrow p. 156ff.). The eliminated amount of PAH is calculated as the arterial inflow of PAH into the kidney minus the venous flow of PAH out of the kidney per unit time. Since

Amount/time = $(\text{volume/time}) \cdot \text{concentration}$ [7.1] $(\text{RPF} \cdot \text{Pa}_{\text{PAH}}) - (\text{RPF} \cdot \text{Prv}_{\text{PAH}}) = \dot{V}_{\text{U}} \cdot U_{\text{PAH}}$ [7.2]

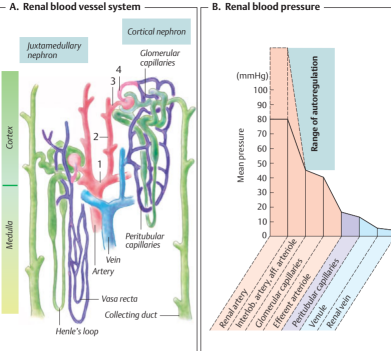
RPF = $\dot{V}_U \cdot U_{PAH}/(Pa_{PAH} - PrV_{PAH})$. [7.3] where Pa_{PAH} is the arterial PAH conc., PrV_{PAH} is the renal venous PAH conc., U_{PAH} is the urinary PAH conc., and \dot{V}_U is the urine output/time. PrV_{PAH} makes up only about 10% of the Pa_{PAH} and normally is not measured directly, but is estimated by dividing PAH clearance (= $\dot{V}_U \cdot U_{PAH}/Pa_{PAH}$; $\rightarrow p. 152$) by a factor of 0.9. Therefore.

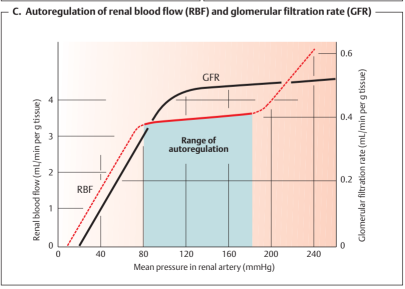
RPF = $\dot{V}_U \cdot U_{PAH}/(0.9 \cdot P_{aPAH})$. [7.4] This equation is only valid when the Pa_{PAH} is not too high. Otherwise, PAH secretion will be saturated and PAH clearance will be much smaller than RPF (\rightarrow p. 161 A).

RBF is derived by inserting the known hematocrit (HCT) value (\rightarrow p. 88) into the following equation:

RBF = RPF/(1 - HCT) [7.5]

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Glomerular Filtration and Clearance

The **glomerular filtration rate (GFR)** is the total volume of fluid filtered by the glomeruli per unit time. It is normally about **120 mL/min** per 1.73 m² of body surface area, equivalent to around 180 L/day. Accordingly, the volume of exchangeable extracellular fluid of the whole body (ca. 17 L) enters the renal tubules about 10 times a day. About 99% of the GFR returns to the extracellular compartment by tubular reabsorption. The mean *fractional excretion of H₂O* is therefore about 1% of the GFR, and absolute H₂O excretion (= **urine output/time** = \mathring{V}_U) is about 1 to 2 L per day. (The filtration of dissolved substances is described on p. 154).

The GFR makes up about 20% of renal plasma flow, RPF (\rightarrow p. 150). The **filtration fraction** (FF) is defined as the ratio of GFR/RPF. The filtration fraction is increased by *atriopeptin*, a peptide hormone that increases efferent arteriolar resistance (R_e) while lowering afferent arteriolar resistance (R_a). This raises the effective filtration pressure in the glomerular capillaries without significantly changing the overall resistance in the renal circulation.

The effective filtration pressure (P_{eff}) is the driving "force" for filtration. P_{eff} is the glomerular capillary pressure ($P_{cap} \approx 48 \text{ mmHg}$) minus the pressure in Bowman's capsule ($P_{Bow} \approx 13 \text{ mmHg}$) and the oncotic pressure in plasma ($\pi_{cap} = 25 \text{ to } 35 \text{ mmHg}$):

$$P_{eff} = P_{cap} - P_{Bow} - \pi_{cap}$$
 [7.6] P_{eff} at the arterial end of the capillaries equals $48-13-25=10$ mmHg. Because of the high filtration fraction, the plasma protein concentration and, therefore, π_{cap} values along the glomerular capillaries increase (\rightarrow p. 378) and P_{eff} decreases. (The mean effective filtration pressure, \bar{P}_{eff} , is therefore used in Eq. 7.7.) Thus, filtration ceases (near distal end of capillary) when π_{cap} rises to about 35 mmHg, decreasing P_{eff} to zero (filtration equilibrium).

GFR is the product of \overline{P}_{eff} (mean for all glomeruli), the glomerular filtration area A (dependent on the number of intact glomeruli), and the water permeability k of the glomerular filter. The *ultrafiltration coefficient K_f* is used to represent $A \cdot k$. This yields

$$\mathbf{GFR} = \overline{\mathbf{P}}_{\mathbf{eff}} \cdot \mathbf{K}_{\mathbf{f}}. \tag{7.7}$$

Indicators present in the plasma are used to **measure GFR.** They must have the following properties:

- They must be freely filterable
- Their filtered amount must not change due to resorption or secretion in the tubule
- They must not be metabolized in the kidney
- They must not alter renal function

Inulin, which must be infused intravenously, fulfills these requirements. **Endogenous creatinine** (normally present in blood) can also be used with certain limitations.

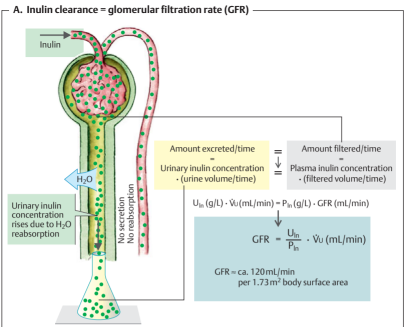
The amount of indicator filtered over time $(\rightarrow A)$ is calculated as the plasma concentration of the indicator $(P_{In}, in g/L \text{ or mol/L})$ times the GFR in L/min. The same amount of indicator/time appears in the urine (conditions 2 and 3; see above) and is calculated as \dot{V}_U (in L/min), times the indicator conc. in urine $(U_{In}, in g/L \text{ or mol/L}, resp.)$, i.e. $P_{In} \cdot GFR = \dot{V}_U \cdot U_{In}$, or:

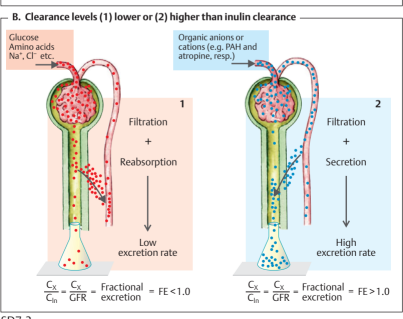
$$GFR = \frac{\dot{\mathbf{V}}_{U} \cdot \mathbf{U}_{ln}}{\mathbf{P}_{ln}} [L/min] (\rightarrow \mathbf{A}).$$
 [7.8]

The expression on the right of Eq. 7.8 represents **clearance**, regardless of which substance is being investigated. Therefore, the *inulin or creatinine clearance represents the GFR*. (Although the plasma concentration of creatinine, P_{cr} , rises as the GFR falls, P_{cr} alone is a quite unreliable measure of GFR.)

Clearance can also be regarded as the completely indicator-free (or cleared) plasma volume flowing through the kidney per unit time. **Fractional excretion (FE)** is the ratio of clearance of a given substance X to inulin clearance (C_X/C_{In}) and defines which fraction of the filtered quantity of X was excreted (cf. p. 154). FE < 1 if the substance is removed from the tubule by reabsorption (e.g. Na⁺, Cl⁻, amino acids, glucose, etc.; \rightarrow **B1**), and FE > 1 if the substance is subject to filtration plus tubular secretion (\rightarrow **B2**). For PAH (\rightarrow p. 150), tubular secretion is so effective that FE_{PAH} \approx 5 (500%).

The absolute rate of reabsorption or secretion of a freely filterable substance X (mol/min) is calculated as the difference between the filtered amount/time (GFR- P_X) and the excreted amount/time (\dot{V}_U - U_X), where a positive result means net reabsorption and a negative net secretion. (For inulin, the result would be zero.)





Transport Processes at the Nephron

Filtration of solutes. The glomerular filtrate also contains small dissolved molecules of plasma (ultrafiltrate) (\rightarrow p. 152). The glomerular sieving coefficient GSC of a substance (= concentration in filtrate/concentration in plasma water) is a measure of the permeability of the glomerular filter for this substance (\rightarrow p. 148). Molecules with a radius of r < 1.8 nm (molecular mass < ca. 10000 Da) can freely pass through the filter (GSC \approx 1.0), while those with a radius of r > 4.4 nm (molecular mass > 80 000 DA, e.g., globulins) normally cannot pass through it (GSC = 0). Only a portion of molecules where $1.8 \, \text{nm} < r < 4.4 \, \text{nm}$ applies are able to pass through the filter (GSC ranges between 1 and 0). Negatively charged particles (e.g., albumin: $r = 3.4 \,\text{nm}$; GSC ≈ 0.0003) are less permeable than neutral substances of equal radius because negative charges on the wall of the glomerular filter repel the ions. When small molecules are bound to plasma proteins (protein binding), the bound fraction is practically non-filterable (\rightarrow p. 24).

Molecules entrapped in the glomerular filter are believed to be eliminated by phagocytic mesangial macrophages (\rightarrow p.94ff.) and glomerular podocytes.

Tubular epithelium. The epithelial cells lining the renal tubule and collecting duct are polar cells. As such, their luminal (or apical) membrane on the urine side differs significantly from that of the basolateral membrane on the blood side. The luminal membrane of the proximal tubule has a high brush border consisting of microvilli that greatly increase the surface area (especially in the convoluted proximal tubule). The basolateral membrane of this tubule segment has deep folds (basal labyrinth) that are in close contact with the intracellular mitochondria (→ p.9B), which produce the ATP needed for Na⁺-K⁺-ATPase (\rightarrow p. 26) located in the basolateral membrane (of all epithelial cells). The large surface areas (about 100 m²) of the proximal tubule cells of both kidneys are needed to reabsorb the lion's share of filtered solutes within the contact time of a couple of seconds. Postproximal tubule cells do not need a brush border since the amount of substances reabsorbed decreases sharply from

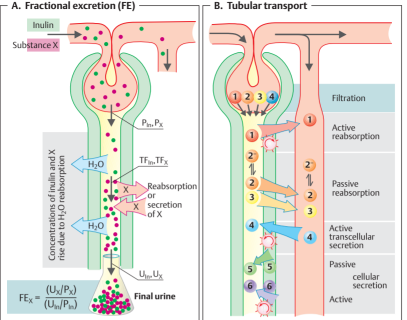
the proximal to the distal segments of the tubules.

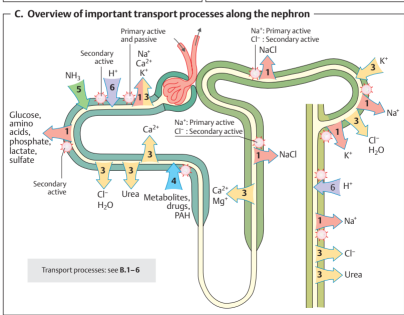
Whereas permeability of the two membranes in series is decisive for transcellular transport (reabsorption, secretion), the tightness of tight junctions (\rightarrow p. 18) determines the paracellular permeability of the epithelium for water and solutes that cross the epithelium by paracellular transport. The tight junctions in the proximal tubule are relatively permeable to water and small ions which, together with the large surface area of the cell membranes. makes the epithelium well equipped for paraand transcellular mass transport ($\rightarrow \mathbf{D}$, column 2). The thin limbs of Henle's loop are relatively "leaky", while the thick ascending limb and the rest of the tubule and collecting duct are "moderately tight" epithelia. The tighter epithelia can develop much higher transepithelial chemical and electrical gradients than "leaky" epithelia.

Measurement of reabsorption, secretion and excretion. Whether and to which degree a substance filtered by the glomerulus is reabsorbed or secreted at the tubule and collecting duct cannot be determined based on its urinary concentration alone as concentrations rise due to the reabsorption of water $(\rightarrow p. 164)$. The urinary/plasma inulin (or creatinine) concentration ratio, Uin/Pin is a measure of the degree of water reabsorption. These substances can be used as indicators because they are neither reabsorbed nor secreted $(\rightarrow p. 152)$. Thus, changes in indicator concentration along the length of the tubule occur due to the H_2O reabsorption alone $(\rightarrow A)$. If U_{in}/P_{in} = 200, the inulin concentration in the final urine is 200 times higher than in the original filtrate. This implies that fractional excretion of H_2O (FE_{H₂O}) is 1/200 or 0.005 or 0.5% of the GFR. Determination of the concentration of a (freely filterable and perhaps additionally secreted) substance X in the same plasma and urine samples for which Uin/Pin was measured will yield U_x/P_x. Considering U_{in}/P_{in}, the fractional excretion of X, FE_X can be calculated as follows (\rightarrow **A** and **D**, in % in column 5):

 $\begin{array}{lll} \textbf{FE}_X = \textbf{(U}_X/P_X)/\textbf{(U}_{In}/P_{In}) & [7.9] \\ \text{Eq. } 7.9 & \text{can also be derived from } C_X/C_{in} \\ (\rightarrow \text{p. } 152) & \text{when simplified for } V_U. & \text{The } \textbf{fractional reabsorption} \text{ of } X \text{ (FR}_X) \text{ is calculated as} \\ \end{array}$

 $FR_X = 1 - FE_X \tag{7.10}$





Reabsorption in different segments of the tubule. The concentration of a substance X (TF_X) and inulin (TF_{in}) in tubular fluid can be measured via micropuncture (\rightarrow **A**). The values can be used to calculate the non-reabsorbed fraction (*fractional delivery*, FD) of a freely filtered substance X as follows:

 ${\rm FD} = ({\rm TF_X/P_X})/({\rm TF_{in}/P_{in}}),$

where P_X and P_{in} are the respective concentrations in plasma (more precisely: in plasma water).

Fractional reabsorption (FR) up to the sampling site can then be derived from $1 - FD \rightarrow D$, columns 2 and 3, in %).

Reabsorption and secretion of various substances (see pp. 16-30, transport mechanisms). Apart from H2O, many inorganic ions (e.g., Na+, Cl-, K+, Ca2+, and Mg2+) and organic substances (e.g., HCO₃-, D-glucose, L-amino acids, urate, lactate, vitamin C, peptides and proteins: \rightarrow C. D. p. 158ff.) are also subject to tubular **reabsorption** (\rightarrow **B1-3**). Endogenous products of metabolism (e.g., urate, glucuronides, hippurates, and sulfates) and foreign substances (e.g., penicillin, diuretics, and PAH; \rightarrow p. 150) enter the tubular urine by way of transcellular secretion (\rightarrow B4, C). Many substances, such as ammonia (NH₃) and H⁺ are first produced by tubule cells before they enter the tubule by cellular secretion. NH₃ enters the tubule lumen by passive transport $(\rightarrow B5)$, while H+ ions are secreted by active transport $(\rightarrow$ **B6** and p. 174 ff.).

 Na^+/K^+ transport by Na^+-K^+ -ATPase ($\rightarrow p, 26$) in the basolateral membrane of the tubule and collecting duct serves as the "motor" for most of these transport processes. By primary active transport (fueled directly by ATP consumption), Na+-K+-ATPase pumps Na+ out of the cell into the blood while pumping K+ in the opposite direction (subscript "i" = intracellular and "o" = extracellular). This creates two driving "forces" essential for the transport of numerous substances (including Na⁺ and K⁺): first, a chemical Na⁺ gradient ($[Na^+]o > [Na^+]i$) and (because $[K^+]i > [K^+]o$), second, a membrane potential (inside the cell is negative relative to the outside) which represents an electrical gradient and can drive ion transport $(\rightarrow pp. 32ff. and 44).$

Transcellular transport implies that two membranes must be crossed, usually by two different mechanisms. If a given substance (D-glucose, PAH, etc.) is actively transported across an epithelial barrier (i.e., against an electrochemical gradient; → see p. 26ff.), at least one of the two serial membrane transport steps must also be active.

Interaction of transporters. Active and passive transport processes are usually closely interrelated. The active absorption of a solute such as Na⁺ or D-glucose, for example, results in the development of an osmotic gradient $(\rightarrow p, 24)$, leading to the passive absorption of water. When water is absorbed, certain solutes are carried along with it (solvent drag; \rightarrow p. 24), while other substrates within the tubule become more concentrated. The latter solutes (e.g., Cl- and urea) then return to the blood along their concentration gradients by passive reabsorption. Electrogenic ion transport and ion-coupled transport (\rightarrow p. 28) can depolarize or hyperpolarize only the luminal or only the basolateral membrane of the tubule cells. This causes a transepithelial potential which serves as the driving "force" for paracellular ion transport in some cases.

Since non-ionized forms of weak electrolytes are more lipid-soluble than ionized forms, they are better able to penetrate the membrane (non-ionic diffusion; \rightarrow B2). Thus, the pH of the urine has a greater influence on passive reabsorption by non-ionic diffusion. Molecular size also influences diffusion: the smaller a molecule, the larger its diffusion coefficient (\rightarrow p. 20ff.).

	1	2	3	4	5	6
	_	Fractio				
Substance	Concentration in plasma water (P) [mmol/L]	% in proximal tubule (TF/P)	% in loop of Henle (TF/P)	Total %	Fractional excretion (FE) [% of filtered amount]	P= Plasma concentration in tubular urine ↑ raises FE ↓ lowers FE
H ₂ O		65%	10%	93%-99.5%	0.5%-7%	ADH: ↓
Na ⁺	153	65% (1.0)	25% (0.4)	95%-99.5%	0.5%-5%	Aldosterone: ADH: ANP:
K+	4,6	65% (1.0)	10%-20%	Secretion possible	2%-150%	Aldosterone: †
Ca ²⁺	Free: 1.6	60% (1.1)	30%	95%-99%	1%-5%	PTH: Acidosis:
Mg ²⁺	Free: 0.6	15% (2.5)	ca. 70%	80%-95%	5%-20%	P rise:
Cl	112	55% (1.3)	ca. 20%	95%-99.5%	0.5%-5%	
HCO ₃	24	93% (0.2)		98%-99%	1%-2%	Alkalosis:
Phosphate	2.2	65% (1.0)	15%	80%-97%	3%-20%	P rise: PTH: Ca ²⁺ falls: Acidosis:
D-Glucose	5	96% (0.1)	4%	≈100%	≈0%	Sharp P rise:
Urea	5	50% (1.4)	Secretion	ca. 60%	ca. 40%	Diuresis:
Creatinine	0.1	0% (2.9)	0%	0%	100%	
PAH (I.V.)	C _{test}	Secretion	Secretion	Secretion	≈500%	Sharp Prise: ↓

Reabsorption of Organic Substances

The **filtered load of a substance** is the product of its plasma concentration and GFR. Since the GFR is high (ca. 180 L/day), enormous quantities of substances enter the primary urine each day (e.g., 160 g/day of D-glucose).

Fractional excretion (FE, \rightarrow p. 154) of **D-glucose** is very low (FE \approx 0.4%). This virtually complete reabsorption is achieved by secondary active transport (Na^{+} -glucose symport) at the luminal cell membrane (\rightarrow **B** and p. 29 B1). About 95% of this activity occurs in the proximal tubule. If the plasma glucose conc. exceeds 10–15 mmol/L, as in diabetes mellitus (normally 5 mmol/L), glucosuria develops, and urinary glucose conc. rises (\rightarrow **A**). Glucose reabsorption therefore exhibits saturation kinetics (Michaelis-Menten kinetics; \rightarrow p. 28). The above example illustrates prerenal glucosuria. Renal glucosuria can also occur when one of the tubular glucose carriers is defective.

Low-affinity carriers in the luminal cell membrane of the pars convoluta (sodium-glucose transporter type 2 = SGLT2) and high-affinity carriers (SGLT1) in the pars recta are responsible for D-glucose reabsorption. The co-transport of D-glucose and Na⁺ occurs in each case, namely at a ratio of 1:1 with SGLT2 and 1:2 with SGLT1. The energy required for this form of secondary active glucose transport is supplied by the electrochemical Na⁺ gradient directed towards the cell interior. Because of the co-transport of two Na⁺ ions, the gradient for SGLT1 is twice as large as that for SGLT2. A uniporter (GLUT2 = glucose transporter type 2) on the blood side facilitates the passive transport of accumulated intracellular glucose out of the cell (facilitated diffusion, \rightarrow p. 22). **D-galactose** also makes use of the SGLT1 carrier, while D-fructose is passively absorbed by tubule cells (GLUT5).

The plasma contains over 25 **amino acids**, and about 70 g of amino acids are filtered each day. Like D-glucose, most L-amino acids are reabsorbed at proximal tubule cells by Na⁺-coupled secondary active transport (\rightarrow B and p. 29 B3). At least 7 different amino acid transporters are in the proximal tubule, and the specificities of some overlap. J_{max} and K_{M} (\rightarrow p. 28) and, therefore, saturability and reabsorption capacities vary according to the type of amino acid and carrier involved. Fractional excretion of most amino acids \approx to 1% (ranging from 0.1% for L-valine to 6% for L-histidine).

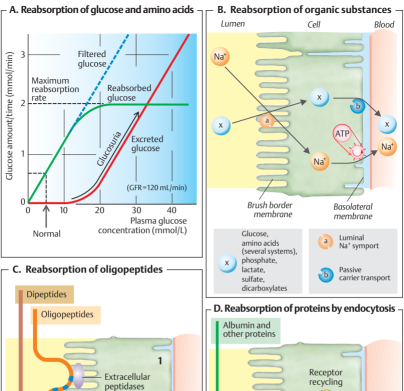
Increased urinary excretion of amino acids (hyperaminoaciduria) can occur. Prerenal hyperaminoaciduria occurs when plasma amino acid concentrations are elevated (and reabsorption becomes saturated, as in A), whereas renal hyperaminoaciduria occurs due to deficient transport. Such a dysfunction may be specific (e.g., in cystinuria, where only L-cystine, L-arginine and L-lysine are hyperexcreted) or unspecific (e.g., in Fanconi's syndrome, where not only amino acids but also glucose, phosphate, bicarbonate etc. are hyperexcreted).

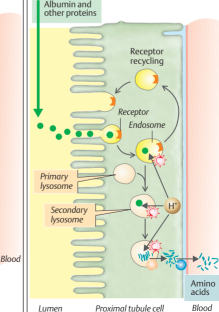
Certain substances (lactate, sulfate, phosphate, dicarboxylates, etc.) are also reabsorbed at the proximal tubule by way of Na * symport, whereas urea is subject to passive back diffusion (\rightarrow p. 166). **Urate** and **oxalate** are both reabsorbed and secreted (\rightarrow p. 160), with the predominant process being reabsorption for urate (FE \approx 0.1) and secretion for oxalate (FE > 1). If the urinary conc. of these poorly soluble substances rises above normal, they will start to precipitate (increasing the risk of **urinary calculus formation**). Likewise, the excessive urinary excretion of **cystine** can lead to cystine calculi.

Oligopeptides such as glutathione and angiotensin II are broken down so quickly by *luminal peptidases* in the brush border that they can be reabsorbed as free amino acids (\rightarrow C1). **Dipeptides** resistant to luminal hydrolysis (e.g., carnosine) must be reabsorbed as intact molecules. A symport carrier (*PepT2*) driven by the inwardly directed H⁺ gradient (\rightarrow p. 174) transports the molecules into the cells *tertiary active* H⁺ symport; \rightarrow p. 26, 29 B4). The dipeptides are then hydrolyzed within the cell (\rightarrow C2). The PepT2 carrier is also used by certain drugs and toxins.

Proteins. Although *albumin* has a low sieving coefficient of 0.0003 (→ p. 154, 2400 mg/day are filtered at a plasma conc. of 45 g/L (180 L/day · 45 g/L · 0.0003 = 2400 mg/day). Only 2 to 35 mg of albumin are excreted each day (FE ≈ 1%). In the proximal tubule, albumin, lysozyme, α₁-microglobulin, β₂-microglobulin and other proteins are reabsorbed by *receptormediated endocytosis* (→ p. 28) and are "digested" by lysosomes (→ **D**). Since this type of reabsorption is nearly saturated at normal filtered loads of proteins, an elevated plasma protein conc. or increased protein sieving coefficient will lead to *proteinuria*.

25-OH-cholecalciferol, which is bound to **DBP** (vitamin D-binding protein) in plasma and glomerular filtrate, is reabsorbed in combination with DBP by receptor-mediated endocytosis (\rightarrow p. 292).





2

Reabsorbed as free amino acids

Lumen

Intracellular

peptidases

Cell

Na⁺ symport carrier

b) H⁺ symport carrier

Excretion of Organic Substances

Food provides necessary nutrients, but also contains inert and harmful substances. The body can usually sort out these substances already at the time of intake, either based on their smell or taste or, if already eaten, with the help of specific digestive enzymes and intestinal absorptive mechanisms (e.g., D-glucose and L-amino acids are absorbed, but L-glucose and D-amino acids are not). Similar distinctions are made in **hepatic excretion** (\Rightarrow bile \Rightarrow stools): useful bile salts are almost completely reabsorbed from the gut by way of specific carriers, while waste products such as bilirubin are mainly eliminated in the feces. Likewise, the kidney reabsorbs hardly any useless or harmful substances (including end-products such as creatinine). Valuable substances (e.g., D-glucose and L-amino acids), on the other hand, are reabsorbed via specific transporters and thus spared from excretion (\rightarrow p. 158).

The liver and kidney are also able to modify endogenous waste products and foreign compounds (xenobiotics) so that they are "detoxified" if toxic and made ready for rapid elimination. In unchanged form or after the enzymatic addition of an OH or COOH group, the substances then combine with glucuronic acid, sulfate, acetate or glutathione to form conjugates. The conjugated substances are then secreted into the bile and proximal tubule lumen (with or without further metabolic processing).

Tubular Secretion

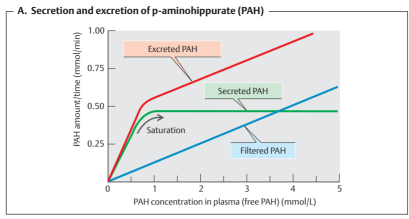
The proximal tubule utilizes active transport mechanisms to secrete numerous waste products and xenobiotics. This is done by way of carriers for organic anions (OA-) and organic cations (OC+). The secretion of these substances makes it possible to raise their clearance level above that of inulin and, therefore, to raise their fraction excretion (FE) above 1.0 = 100% (\rightarrow p. 152) in order to eliminate them more effectively ($\rightarrow A$; compare red and blue curves). Secretion is carrier-mediated and is therefore subject to saturation kinetics. Unlike reabsorbed substances such as D-glucose $(\rightarrow p. 159 \text{ A})$, the fractional excretion (FE) of organic anions and cations decreases when their plasma concentrations rise ($\rightarrow A$; PAH secretion curve reaches plateau, and slope of PAH excretion curve decreases). Some organic anions (e.g., urate and oxalate) and cations (e.g., choline) are both secreted and reabsorbed (bidirectional transport), which results in net reabsorption (urate, choline) or net secretion (oxalate).

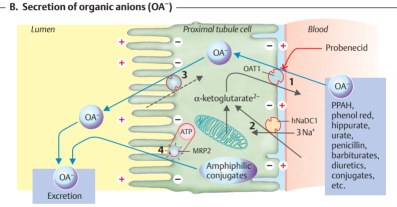
The secreted **organic anions** (OA^-) include indicators such as PAH (p-aminohippurate; \rightarrow p. 150) and phenol red; endogenous substances such as oxalate, urate, hippurate; drugs such as penicillin G, barbiturates, and numerous diuretics (\rightarrow p. 172); and conjugated substances (see above) containing glucuronate, sulfate or glutathione. Because of its high affinity for the transport system, *probenecid* is a potent inhibitor of OA^- secretion.

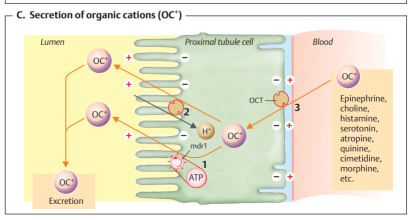
The active step of OA^- secretion ($\rightarrow B$) occurs across the basolateral membrane of proximal tubule cells and accumulates organic anions in the cell whereby the inside-negative membrane potential has to be overcome. The membrane has a broad specificity carrier (OAT1 = organic anion transporter type 1) that transports OA- from the blood into the tubule cells in exchange for a dicarboxylate, such as succinate2or α -ketoglutarate²⁻; \rightarrow **B1**). The latter substance arises from the glutamine metabolism of the cell (→ p. 177 D2): the human Na⁺-dicarboxylate transporter hNADC-1 also conveys dicarboxylates (in combination with 3 Na⁺) into the cell by secondary active transport (\rightarrow **B2**). The transport of OA⁻ is therefore called tertiary active transport. The efflux of OA- into the lumen is passive (facilitated diffusion; \rightarrow **B3**). An ATP-dependent conjugate pump (MRP2 = multi-drug resistance protein type 2) in the luminal membrane is also used for secretion of amphiphilic conjugates, such as glutathione-linked lipophilic toxins (\rightarrow **B4**).

The **organic cations** (\mathbf{OC}^+) secreted include endogenous substances (epinephrine, choline, histamine, serotonin, etc.) and drugs (atropine, quinidine, morphine, etc.).

The active step of $\mathbf{OC^+}$ secretion occurs across the *luminal* membrane of proximal tubule cells (luminal accumulation occurs after overcoming the negative membrane potential inside the cell). The membrane contains (a) direct ATP-driven carriers for organic cations (mdr1; primary active $\mathbf{OC^+}$ transport; \rightarrow $\mathbf{C1}$) and (b) a multispecific $\mathbf{OC^+}$ /H⁺ antiporter (tertiary active transport; \rightarrow $\mathbf{C2}$). The $\mathbf{OC^+}$ diffuse passively from the blood into the cell by way of a polyspecific organic cation transporter (\mathbf{OCT} : \rightarrow $\mathbf{C3}$).







Reabsorption of Na⁺ and Cl⁻

About 99% of the *filtered load of Na*⁺ is reabsorbed (ca. 27 000 mmol/day), i.e., the fractional excretion of Na⁺ (FE_{Na}) is about 1%. The precise value of FE_{Na} needed (range 0.5 to 5%) is regulated by aldosterone, atriopeptin (ANF) and other hormones (\rightarrow p. 170).

Sites of Na⁺ reabsorption. The reabsorption of Na⁺ occurs in all parts of the renal tubule and collecting duct. About 65% of the filtered Na⁺ is reabsorbed in the proximal tubule, while the luminal Na ⁺ conc. remains constant (\rightarrow p. 166). Another 25% is reabsorbed in the loop of Henle, where luminal Na⁺ conc. drops sharply; \rightarrow p. 157 D, columns 2 and 3). The distal convoluted tubule and collecting duct also reabsorb Na⁺. The latter serves as the site of hormonal *fine adjustment* of Na⁺ excretion.

Mechanisms of Na⁺ **reabsorption.** Na⁺-K⁺-ATPase pumps Na⁺ ions out of the cell while conveying K⁺ ions into the cell (\rightarrow **A** and p. 156), thereby producing a chemical Na⁺ gradient (\rightarrow **A2**). Back diffusion of K⁺ (\rightarrow **A3**) also leads to the development of a membrane potential (\rightarrow **A4**). Both combined result in a high electrochemical Na⁺ gradient that provides the driving "force" for passive Na⁺ influx, the features of which vary in the individual nephron segments (\rightarrow **B**).

- ♦ In the **proximal tubule**, Na⁺ ions diffuse passively from the tubule lumen into the cells via (a) the electroneutral Na⁺/H⁺ exchanger type 3 (NHE3), an *Na*⁺/H⁺-antiport carrier for electroneutral exchange of Na⁺ for H⁺ (→ **B1**, p. 29 B4 and p. 174) and (b) various *Na*⁺ symport carriers for reabsorption of D-glucose etc. (→ **B1** and p. 158). Since most of these symport carriers are electrogenic, the luminal cell membrane is depolarized, and an early proximal *lumen-negative transepithelial potential* (L**N**TP) develops.
- ♦ In the thick ascending limb (TAL) of the loop of Henle (\rightarrow B6), Na⁺ is reabsorbed via the bumetanide-sensitive co-transporter BSC, a Na⁺-K⁺-2 Cl⁻ symport carrier (\rightarrow p. 172). Although BSC is primarily electroneutral, the absorbed K⁺ recirculate back to the lumen through K⁺ channels. This hyperpolarizes the luminal membrane, resulting in the development of a lumen-positive transepithelial potential (LPTP).

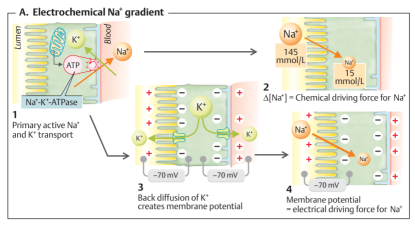
- In the distal convoluted tubule, DCT (→ B8), Na⁺ is reabsorbed via the thiazide-sensitive cotransporter TSC, an electroneutral Na⁺-Cl⁻ symport carrier (→ p. 172).
- ♦ In principal cells of the connecting tubule and collecting duct (\rightarrow **B9**), Na⁺ exits the lumen via Na^+ channels activated by aldosterone and antidiuretic hormone (ADH) and inhibited by prostaglandins and ANF (\rightarrow p. 170).

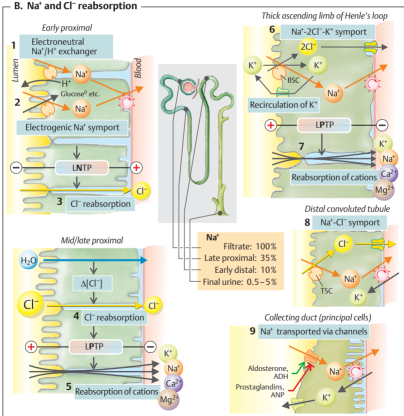
Since these four passive Na⁺ transport steps in the luminal membrane are serially connected to active Na⁺ transport in the basolateral membrane (Na⁺-K⁺-ATPase), the associated **transepithelial Na⁺ reabsorption** is also *active*. This makes up about ¹/₃ of the Na⁺ reabsorption in the proximal tubule, and 1 ATP molecule is consumed for each 3 Na⁺ ions absorbed (→ p. 26). The other ²/₃ of proximal sodium reabsorption is passive and **paracellular**.

Two **driving "forces"** are responsible for this: (1) the LPTP in the mid and late proximal tubule (\rightarrow **B5**) and in the loop of Henle (\rightarrow **B7**) drives Na" and other cations onto the blood side of the epithelium. (2) **Solvent drag** (\rightarrow p. 24): When water is reabsorbed, solutes for which the reflection coefficient < 1 (including Na*) are "dragged along" due to friction forces (like a piece of wood drifts with flowing water). Since driving forces (1) and (2) are indirect products of Na*-K*-ATPase, the **energy balance** rises to about 9 Na* per ATP molecule in the proximal tubule (and to about 5 Na* per ATP molecule in the rest of the kidney).

On the basolateral side, Na $^+$ ions exit the proximal tubule cell via Na $^+$ -K $^+$ -ATPase and an Na $^+$ -3 HCO $_3^-$ symport carrier (\rightarrow p. 174). In the latter case, Na $^+$ exits the cell via tertiary active transport as secondary active secretion of H $^+$ (on the opposite cell side) results in intracellular accumulation of HCO $_3^-$.

The **fractional CI**⁻ **excretion** (**FE**_{CI}) ranges from 0.5% to 5%. About 50% of all **CI**⁻ **reabsorption** occurs in the *proximal tubule*. The early proximal **LNTP** drives CI⁻ through paracellular spaces out of the lumen (\rightarrow **B3**). The reabsorption of CI⁻ lags behind that of Na⁺ and H₂O, so the luminal CI⁻ conc. rises. As a result, CI⁻ starts to diffuse down its chemical gradient paracellularly along the mid and late proximal tubule (\rightarrow **B4**), thereby producing a **LPTP** (reversal of potential, \rightarrow **B5**). At the TAL and the DCT, CI⁻ enters the cells by secondary active transport and exits passively through ADH-activated basolateral CI⁻ channels (\rightarrow **B6**, **8**).





Reabsorption of Water, Formation of Concentrated Urine

The glomeruli filter around 180 L of plasma water each day (= GFR; \rightarrow p. 152). By comparison, the normal **urine output** (\dot{V}_U) is relatively small (0.5 to 2 L/day). Normal fluctuations are called antidiuresis (low Vu) and diuresis (high \dot{V}_{U} : \rightarrow p. 172). Urine output above the range of normal is called polyuria. Below normal output is defined as oliguria (< 0.5 L/day) or anuria (< 0.1 L/day). The **osmolality** (\rightarrow p. 377) of plasma and glomerular filtrate is about 290 mOsm/kg H₂O (= P_{osm}); that of the final urine (U_{0sm}) ranges from 50 (hypotonic urine in extreme water diuresis) to about 1200 mOsm/ kg H₂O (hypertonic urine in maximally concentrated urine). Since water diuresis permits the excretion of large volumes of H₂O without the simultaneous loss of NaCl and other solutes, this is known as "free water excretion", or "free water clearance" (C_{H_2O}). This allows the kidney to normalize decreases in plasma osmolality, for example (\rightarrow p. 170). The C_{H_2O} represents to the volume of water that could be theoretically extracted in order for the urine to reach the same osmolality as the plasma:

$$C_{H_2O} = \dot{V}_U (1 - [U_{osm}/P_{osm}]).$$
 [7.11]

Countercurrent Systems

A **simple exchange** system (\rightarrow **A1**) can consist of two tubes in which *parallel streams* of water flow, one cold (0 °C) and one hot (100 °C). Due to the exchange of heat between them, the water leaving the ends of both tubes will be about 50 °C, that is, the initially steep temperature gradient of 100 °C will be offset.

In **countercurrent exchange** of heat (\rightarrow **A2**), the fluid within the tubes flows in *opposite directions*. Since a temperature gradient is present in all parts of the tube, heat is exchanged along the entire length. Molecules can also be exchanged, provided the wall of the tube is permeable to them and that a concentration gradient exists for the substance.

If the countercurrent exchange of heat occurs in a **hairpin-shaped loop**, the bend of which is in contact with an environment with a temperature different from that inside the tube (ice, \rightarrow **A3**), the fluid exiting the loop will be only slightly colder than that entering it, because heat always passes from the warmer limb of the loop to the colder limb.

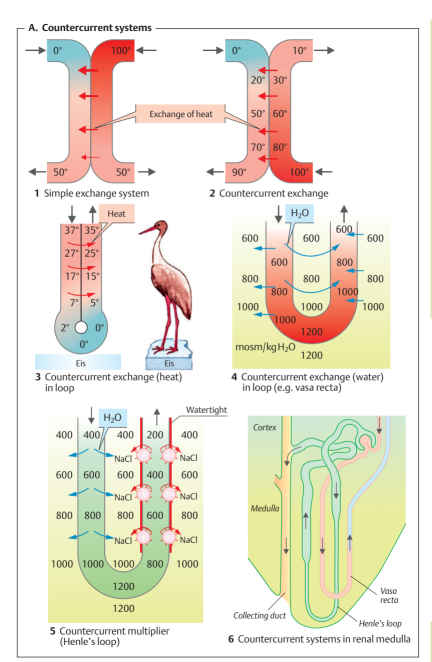
Countercurrent exchange of water in the vasa recta of the renal medulla (\$\infty\$ A6 and p. 150) occurs if the medulla becomes increasingly hy-

pertonic towards the papillae (see below) and if the vasa recta are permeable to water. Part of the water diffuses by osmosis from the descending vasa recta to the ascending ones. thereby "bypassing" the inner medulla (\rightarrow **A4**). Due to the extraction of water, the concentration of all other blood components increases as the blood approaches the papilla. The plasma osmolality in the vasa recta is therefore continuously adjusted to the osmolality of the surrounding interstitium, which rises towards the papilla. The hematocrit in the vasa recta also rises. Conversely, substances entering the blood in the renal medulla diffuse from the ascending to the descending vasa recta, provided the walls of both vessels are permeable to them (e.g., urea; \rightarrow **C**). The countercurrent exchange in the vasa recta permits the necessary supply of blood to the renal medulla without significantly altering the high osmolality of the renal medulla and hence impairing the urine concentration capacity of the kidney.

In a **countercurrent multiplier** such as the **loop of Henle**, a concentration gradient between the two limbs is maintained by the expenditure of energy (\rightarrow A5). The countercurrent flow amplifies the relatively small gradient at all points between the limbs (*local gradient* of about 200 mOsm/kg H₂O) to a relatively large gradient along the limb of the loop (about 1000 mOsm/kg H₂O). The longer the loop and the higher the one-step gradient, the steeper the multiplied gradient. In addition, it is inversely proportional to (the square of) the flow rate in the loop.

Reabsorption of Water

Approximately 65% of the GFR is reabsorbed at the **proximal convoluted tubule**, **PCT** (\rightarrow **B** and p. 157 D). The driving "force" for this is the reabsorption of solutes, especially Na⁺ and Cl⁻. This slightly dilutes the urine in the tubule, but H₂O immediately follows this small osmotic gradient because the PCT is "leaky" (\rightarrow p. 154). The reabsorption of water can occur by a paracellular route (through leaky tight junctions) or transcellular route, i.e., through water channels (aquaporin type 1 = AQP1) in the two cell membranes. The urine in PCT therefore remains (virtually) isotonic. Oncotic pressure (\rightarrow p. 378) in the peritubular capillaries pro-



vides an additional driving force for water reabsorption. The more water filtered at the glomerulus, the higher this oncotic pressure. Thus, the reabsorption of water at the proximal tubule is, to a certain extent, adjusted in accordance with the GFR (glomerulotubular balance).

Because the descending limb of the loop of Henle has aquaporins (AOP1) that make it permeable to water, the urine in it is largely in osmotic balance with the hypertonic interstitium, the content of which becomes increasingly hypertonic as it approaches the papillae $(\rightarrow A5)$. The urine therefore becomes increasingly concentrated as it flows in this direction. In the thin descending limb, which is only sparingly permeable to salt, this increases the conc. of Na+ and Cl-. Most water drawn into the interstitium is carried off by the vasa recta $(\rightarrow B)$. Since the thin and thick ascending limbs of the loop of Henle are largely impermeable to water, Na+ and Cl- passively diffuses (thin limb) and is actively transported (thick limb) out into the interstitium $(\rightarrow B)$. Since water cannot escape, the urine leaving the loop of Henle is hypotonic.

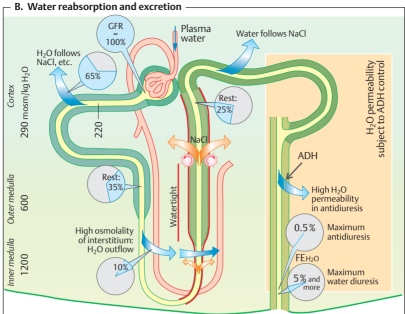
Active reabsorption of Na $^+$ and Cl $^-$ from the thick ascending limb of the loop of Henle (TAL; \rightarrow p. 162) creates a **local gradient** (ca. 200 mOsm/kg H $_2$ O; \rightarrow A5) at all points between the TAL on the one side and the descending limb and the medullary interstitium on the other. Since the high osmolality of fluid in the medullary interstice is the reason why water is extracted from the collecting duct (see below), active NaCl transport is the ATP-consuming "motor" for the kidney's urine-concentrating mechanism and is up-regulated by sustained stimulation of ADH secretion.

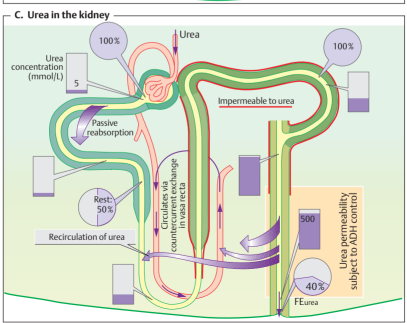
Along the course of the **distal convoluted tubule** and, at the latest, at the **connecting tubule**, which contains aquaporins and ADH receptors of type V_2 (explained below), the fluid in the tubule will again become isotonic (in osmotic balance with the isotonic interstice of the renal cortex) if ADH is present (\rightarrow p. 168), i.e., when *antidiuresis* occurs. Although Na $^+$ and Cl $^-$ are still reabsorbed here (\rightarrow p. 162), the osmolality does not change significantly because H_2O is reabsorbed (ca. 5% of the GFR) into the interstitial space due to osmotic forces and *urea* increasingly determines the osmolality of the tubular fluid.

Final adjustment of the excreted urine volume occurs in the collecting duct. In the presence of antidiuretic hormone, ADH (which binds to basolateral V₂ receptors, named after vasopressin, the synonym for ADH), aquaporins (AQP2) in the (otherwise water-impermeable) luminal membrane of principal cells are used to extract enough water from the urine passing through the increasingly hypertonic renal medulla. Thereby, the Uosm rises about four times higher than the $P_{osm}(U_{osm}/P_{osm} \approx 4)$, corresponding to maximum antidiuresis. The absence of ADH results in water diuresis, where U_{osm}/P_{osm} can drop to < 0.3. The U_{osm} can even fall below the osmolality at the end of TAL, since reabsorption of Na+ and Cl- is continued in the distal convoluted tubule and collecting duct (\rightarrow p. 162) but water can hardly follow.

Urea also plays an important role in the formation of concentrated urine. A protein-rich diet leads to increased urea production, thus increased the urine-concentrating capacity of the kidney. About 50% of the filtered urea leaves the proximal tubule by diffusion $(\rightarrow C)$. Since the ascending limb of the loop of Henle, the distal convoluted tubule, and the cortical and outer medullary sections of the collecting duct are only sparingly permeable to urea, its conc. increases downstream in these parts of the nephron (\rightarrow C). ADH can (via V₂ receptors) introduce urea carriers (urea transporter type 1, UT1) in the luminal membrane, thereby making the inner medullary collecting duct permeable to urea. Urea now diffuses back into the interstitium (where urea is responsible for half of the high osmolality there) via UT1 and is then transported by UT2 carriers back into the descending limb of the loop of Henle, comprising the recirculation of urea $(\rightarrow C)$. The non-reabsorbed fraction of urea is excreted: FE_{urea} ≈ 40%. Urea excretion increases in water diuresis and decreases in antidiuresis, presumably due to up-regulation of the UT2 carrier.

Urine concentration disorders primarily occur due to (a) excessive medullary blood flow (washing out Na⁺, Cl[−] and urea); (b) osmotic diuresis; (c) loop diuretics (→p. 172); (d) deficient secretion or effectiveness of ADH, as seen in *central* or *peripheral diabetes insipidus*, respectively.





Body Fluid Homeostasis

Life cannot exist without water. Water is the initial and final product of countless biochemical reactions. It serves as a solvent, transport vehicle, heat buffer, and coolant, and has a variety of other functions. Water is present in cells as intracellular fluid, and surrounds them as extracellular fluid. It provides a constant environment (internal milieu) for cells of the body, similar to that of the primordial sea surrounding the first unicellular organisms (\rightarrow p. 2).

The volume of fluid circulating in the body remains relatively constant when the **water balance** (\rightarrow **A**) is properly regulated. The average **fluid intake** of ca. 2.5 L per day is supplied by *beverages*, *solid foods*, and *metabolic oxidation* (\rightarrow p. 229 C). The fluid intake must be high enough to counteract **water losses** due to *urination*, *respiration*, *perspiration*, and *defecation* (\rightarrow p. 265 C). The mean daily H₂O turnover is 2.5 L/70 kg (1/30 th the body weight [BW]) in adults and 0.7 liters/7 kg (1/10th the BW) in infants. The water balance of infants is therefore more susceptible to disturbance.

Significant **rises in the H₂O turnover** can occur, but must be adequately compensated for if the body is to function properly (regulation, \rightarrow p. 170). Respiratory H₂O losses occur, for example, due to *hyperventilation* at high altitudes (\rightarrow pp. 106 and 136), and *perspiration losses* (\rightarrow p. 222) occur due to exertion at high temperatures (e.g., hiking in the sun or hot work environment as in an irronworks). Both can lead to the loss of several liters of water per hour, which must be compensated for by increasing the intake of fluids (and salt) accordingly. Conversely, an increased intake of fluids will lead to an increased volume of urine being excreted (\rightarrow p. 170).

Water loss (hypovolemia) results in the stimulation of **thirst**, a sensation controlled by the so-called *thirst center in the hypothalamus*. Thirst is triggered by significant rises in the osmolality of body fluids and angiotensin II concentration of cerebrospinal fluid (\rightarrow p. 170).

Body water content. The fraction of total body water (TBW) to body weight (BW = 1.0) ranges from **0.46** (46%) **to 0.75** depending on a person's age and $sex (\rightarrow B)$. The TBW content in

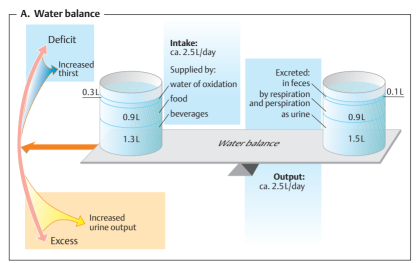
infants is 0.75 compared to only 0.64 (0.53) in young men (women) and 0.53 (0.46) in elderly men (women). Gender-related differences (and interindividual differences) are mainly due to differences in a person's total body fat content. The average fraction of water in most body tissues (in young adults) is 0.73 compared to a fraction of only about 0.2 in fat $(\rightarrow B)$.

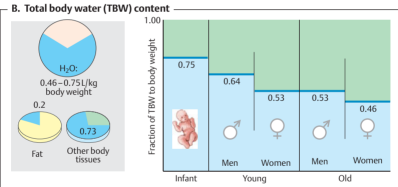
Fluid compartments. In a person with an average TBW of ca. 0.6, about 3/5 (0.35 BW) of the TBW is intracellular fluid (ICF), and the other 2/5 (0.25 BW) is extracellular. Extracellular fluid (ECF) is located between cells (interstice, 0.19), in blood (plasma water (0.045) and in "transcellular" compartments (0.015) such as the CSF and intestinal lumen (\rightarrow C). The protein concentration of the plasma is significantly different from that of the rest of the ECF. Moreover, there are fundamental differences in the ionic composition of the ECF and the ICF $(\rightarrow p. 93 B)$. Since most of the body's supply of Na⁺ ions are located in extracellular compartments, the total Na+ content of the body determines its ECF volume (\rightarrow p. 170).

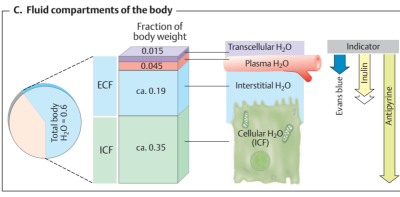
Measurement of fluid compartments. In clinical medicine, the body's fluid compartments are usually measured by *indicator dilution techniques*. Provided the indicator substance, S, injected into the bloodstream spreads to the target compartment only (\rightarrow C), its volume V can be calculated from:

V[L] = injected amount of

 $S [mol]/C_S [mol/L]$ [7.12] where C_S is the concentration of S after it spreads throughout the target compartment (measured in collected blood specimens). The ECF volume is generally measured using inulin as the indicator (does not enter cells), and the TBW volume is determined using antipyrine. The ICF volume is approximately equal to the antipyrine distribution volume minus the inulin distribution volume. Evans blue, a substance entirely bound by plasma proteins, can be used to measure the plasma volume. Once this value is known, the blood volume can be calculated as the plasma volume divided by [1 - hematocrit] (\rightarrow p. 88), and the interstitial volume is calculated as the ECF volume minus the plasma volume.







Salt and Water Regulation

Osmoregulation. The osmolality of most body fluids is about 290 mOsm/kg H₂O. Any increase in the osmolality of extracellular fluid (ECF) due, for example, to NaCl absorption or water loss, results in an outflow of water from the intracellular space, because the intracellular fluid (ICF) and ECF are in osmotic balance (\rightarrow p. 173; B2, B6). The osmolality of the ECF must be tightly regulated to protect cells from large volume fluctuations. Osmoregulation is controlled by osmosensors (or osmoreceptors) found mainly in the hypothalamus, hormones (e.g., antidiuretic hormone = ADH = adiuretin = vasopressin) and the kidney, the target organ of ADH (\rightarrow p. 166).

Water deficit (→ A1). Net water losses (hypovolemia) due, for example, to sweating, urination or respiration, make the ECF hypertonic. Osmolality rises of 1% or more $(\geq 3 \text{ mOsm/kg H}_2\text{O})$ are sufficient to stimulate the secretion of ADH from the posterior lobe of the pituitary (\rightarrow p. 280). ADH decreases urinary H_2O excretion (\rightarrow p. 166). The likewise hypertonic cerebrospinal fluid (CSF) stimulates central osmosensors in the hypothalamus, which trigger hyperosmotic thirst. The perception of thirst results in an urge to replenish the body's water reserves. Peripheral osmosensors in the portal vein region and vagal afferent neurons warn the hypothalamus of water shifts in the gastrointestinal tract.

Water excess (\rightarrow A2). The absorption of hypotonic fluid reduces the osmolality of ECF. This signal inhibits the secretion of ADH, resulting in water diuresis (\rightarrow p. 166) and normalization of plasma osmolality within less than 1 hour.

Water intoxication occurs when excessive volumes of water are absorbed too quickly, leading to symptoms of nausea, vomiting and shock. The condition is caused by an undue drop in the plasma osmolality before adequate inhibition of ADH secretion has occurred.

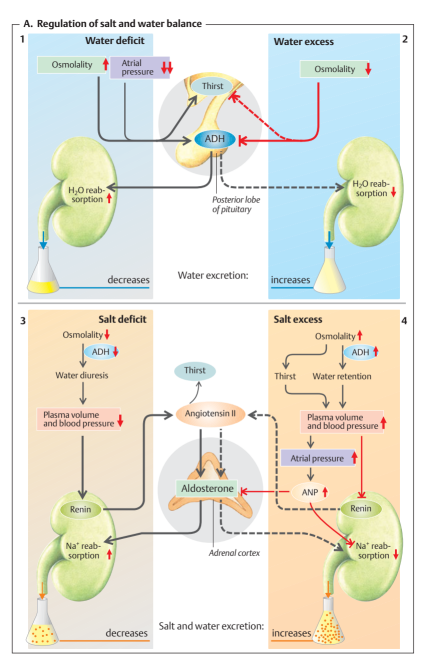
Volume regulation. Around 8–15 g of NaCl are absorbed each day. The kidneys have to excrete the same amount over time to maintain Na⁺ and ECF homeostasis (\rightarrow p. 168). Since Na⁺ is the major extracellular ion (Cl⁻ balance is maintained secondarily), *changes in total body*

Na⁺ *content lead to changes in ECF volume.* It is regulated mainly by the following **factors**:

- ♦ Renin–angiotensin system (RAS) (\rightarrow p. 184). Its activation promotes the retention of Na⁺ via angiotensin II (AT II; lowers GFR), aldosterone (\rightarrow A4) and ADH.
- Atriopeptin (atrial natriuretic peptide; ANP) is a peptide hormone secreted by specific cells of the cardiac atrium in response to rises in ECF volume and hence atrial pressure. ANP promotes the renal excretion of Na⁺ by raising the filtration fraction (→ p. 152) and inhibits Na⁺ reabsorption from the collecting duct.
- ◆ ADH. ADH secretion is stimulated by (a) increased plasma and CSF osmolality; (b) the Gauer-Henry reflex, which occurs when stretch receptors in the atrium warn the hypothalamus of a decrease (>10%) in ECF volume (~atrial pressure); (c) angiotensin II (→ p. 184).
- ◆ Pressure diuresis (→ p. 172), caused by an elevated arterial blood pressure, e.g. due to an elevated ECF volume, results in increased excretion of Na⁺ and water, thereby lowering ECF volume and hence blood pressure. This control circuit is thought to be the major mechanism for long term blood pressure regulation.

Salt deficit (\rightarrow A3). When hyponatremia occurs in the presence of a primarily normal H₂O content of the body, blood osmolality and therefore ADH secretion decrease, thereby increasing transiently the excretion of H₂O. The ECF volume, plasma volume, and blood pressure consequently decrease (\rightarrow A4). This, in turn, activates the RAS, which triggers hypovolemic thirst by secreting AT II and induces Na⁺ retention by secreting aldosterone. The retention of Na⁺ increases plasma osmolality leading to secretion of ADH and, ultimately, to the retention of water. The additional intake of fluids in response to thirst also helps to normalize the ECF volume.

Salt excess (\rightarrow A4). An abnormally high NaCl content of the body in the presence of a normal H_2O volume leads to increased plasma osmolality (thirst) and ADH secretion. Thus, the ECF volume rises and RAS activity is curbed. The additional secretion of ANP, perhaps together with a natriuretic hormone with a longer half-life than ANP (ouabain?), leads to increased excretion of NaCl and H_2O and, consequently, to normalization of the ECF volume.



Diuresis and Diuretics

Increases in urine excretion above 1 mL/min (diuresis) can have the following causes:

- ♦ Water diuresis: Decreases in plasma osmolality and/or an increased blood volume lead to the reduction of ADH levels and, thus, to the excretion of "free water" (\rightarrow p. 164).
- ◆ Osmotic diuresis results from the presence of non-reabsorbable, osmotically active substances (e.g., mannitol) in the renal tubules. These substances retain H₂O in the tubule lumen, which is subsequently excreted. Osmotic diuresis can also occur when the concentration of an reabsorbable substance (e.g., glucose) exceeds its tubular reabsorption capacity resulting, for example, in hyperglycemia (→ p. 158). The glucosuria occurring in diabetes mellitus is therefore accompanied by diuresis and a secondary increase in thirst. Hyperbicarbonaturia can lead to osmotic diuresis to the same reason (→ p. 176).
- ◆ Pressure diuresis occurs when osmolality in the renal medulla decreases in the presence of increased renal medullary blood flow due, in most cases, to hypertension (→ p. 170).
- ◆ **Diuretics** (→ **A**) are drugs that induce diuresis. Most of them (except osmotic diuretics like mannitol) work primarily by inhibiting NaCl reabsorption (*saluretics*) and, secondarily, by decreasing water reabsorption. The goal of therapeutic diuresis, e.g., in treating edema and hypertension, is to reduce the ECF volume.

Although diuretics basically inhibit NaCl transport throughout the entire body, they have a large degree of **renal "specificity"** because they act from the tubular lumen, where they become highly concentrated due to **tubular secretion** (\rightarrow p. 160) and tubular water reabsorption. Therefore, dosages that do not induce unwanted systemic effects are therapeutically effective in the tubule lumen.

Diuretics of the **carbonic anhydrase inhibitor type** (e.g., acetazolamide, benzolamide) decrease Na^*/H^* exchange and HCO_3^- reabsorption in the proximal tubule (\rightarrow p. 174ff.). The overall extent of diuresis achieved is small because more distal segments of the tubule reabsorb the NaCl not reabsorbed upstream and because the GFR decreases due to tubuloglomerular feedback, TGF (\rightarrow p. 184). In addition, increased HCO_3^- excretion also leads to non-respiratory (metabolic) acidosis. Therefore, this type of diuretic is used only in patients with concomitant alkalosis.

diuretics (e.g., furosemide Loop and bumetanide) are highly effective. They inhibit the bumetanide-sensitive co-transporter BSC $(\rightarrow p. 162 B6)$, a Na⁺-2Cl⁻-K⁺ symport carrier, in the thick ascending limb (TAL) of the loop of Henle. This not only decreases NaCl reabsorption there, but also stalls the "motor" on the concentration mechanism (\rightarrow p. 166). Since the lumen-positive transepithelial potential (LPTP) in the TAL also falls (\rightarrow p. 162 B7), paracellular reabsorption of Na+, Ca2+ and Mg2+ is also inhibited. Because increasing amounts of non-reabsorbed Na+ now arrive at the collecting duct (\rightarrow p. 181 B3), K⁺ secretion increases and the simultaneous loss of H+ leads to hypokalemia and hypokalemic alkalosis.

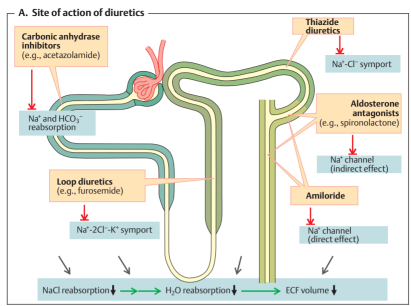
Loop diuretics inhibit BSC at the **macula densa**, thereby "tricking" the juxtaglomerular apparatus (JGA) into believing that no more NaCl is present in the tubular lumen. The GFR then rises as a result of the corresponding tubuloglomerular feedback (\rightarrow p. 184), which further promotes diuresis.

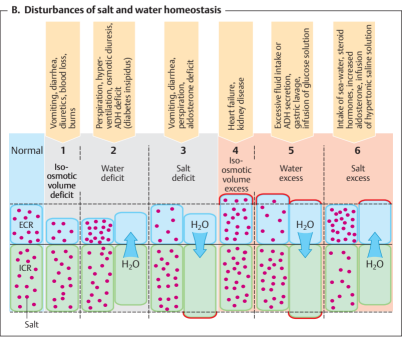
Thiazide diuretics inhibit NaCl resorption in the distal tubule (TSC, \rightarrow p. 162 B8). Like loop diuretics, they increase Na $^+$ reabsorption downstream, resulting in losses of K $^+$ and H $^+$.

Potassium-sparing diuretics. *Amiloride* block Na $^+$ channels in the principal cells of the connecting tubule and collecting duct, leading to a *reduction of K* $^+$ *excretion. Aldosterone antagonists* (e.g., spironolactone), which block the cytoplasmic aldosterone receptor, also have a potassium-sparing effect.

Disturbances of Salt and Water Homeostasis

When osmolality remains normal, disturbances of salt and water homeostasis (\rightarrow **B** and p. 170) only affect the ECF volume (\rightarrow **B1** and 4). When the osmolality of the ECF increases (hyperosmolality) or decreases osmolality), water in the extra- and intracellular compartments is redistributed (\rightarrow **B2. 3. 5.** 6). The main causes of these disturbances are listed in B (orange background). The effects of these disturbances are hypovolemia in cases 1, 2 and 3, intracellular edema (e.g., swelling of the brain) in disturbances 3 and 5, and extracellular edema (pulmonary edema) in disturbances 4, 5 and 6.





The Kidney and Acid-Base Balance

Main functions of renal H^+ secretion ($\rightarrow A$):

- reabsorption of filtered bicarbonate (\rightarrow **B**),
- excretion of H^+ ions measurable as titratable acidity (\rightarrow C), and
- nonionic transport of NH₄⁺, i.e. in the form of NH₃ (→ D1, 2).
- 1. Very large quantities of H⁺ ions are secreted into the lumen of the proximal tubule $(\rightarrow A1)$ by (a) primary active transport via H+-ATPase and (b) by secondary active transport via an electroneutral Na+/H+-antiporter (NHE3 carrier, \rightarrow p. 162). The *luminal pH* then decreases from 7.4 (filtrate) to about 6.6. One OH- ion remains in the cell for each H⁺ ion secreted: OH⁻ reacts with CO₂ to form HCO₃- (accelerated by carbonic anhydrase-II, see below). HCO3leaves the cell for the blood, where it binds one H⁺ ion. Thus, each H⁺ ion secreted into the lumen (and excreted) results in the elimination of one H⁺ ion from the body, except the secreted H⁺ is accompanied by a secreted NH₃ (see below).

2. In the **connecting tubule** and **collecting duct** $(\rightarrow$ **A2**) *type A intercalated cells* secrete H⁺ ions via H⁺/K⁺-ATPase and H⁺-ATPase, allowing the luminal pH to drop as far as 4.5. In metabolic alkalosis, *type B intercalated cells* can secrete HCO₃⁻ $(\rightarrow$ **A3**).

Carbonic anhydrase (CA) is important in all cases where H^+ ions exit from one side of a cell and/or HCO_3^- exits from the other, e.g., in renal tubule cells, which contain CA^{II} in the cytosol and CA^{IV} on the outside of the luminal membrane; \rightarrow A, B, D), as well as in the stomach, small intestine, pancreatic duct and erythrocytes, etc. CA catalyzes the gross reaction

 $H_2O + CO_2 \qquad H^+ + HCO_3^-$.

Carbonic acid (H_2CO_3) is often considered to be the intermediate product of this reaction, but OH⁻ (not H_2O) probably combines with CA. Therefore, the reactions H_2O OH⁻ + H⁺ and OH⁻ + CO₂ HCO₃- underlie the aforementioned gross reaction.

Reabsorption of HCO₃[−] (→ B). The amount of HCO₃[−] filtered each day is 40 times the quantity present in the blood. HCO₃[−] must therefore be reabsorbed to maintain acid–base balance (→ p. 183ff.). The H⁺ ions secreted into the lumen of the proximal convoluted tubule react with about 90% of the filtered HCO₃[−] to form

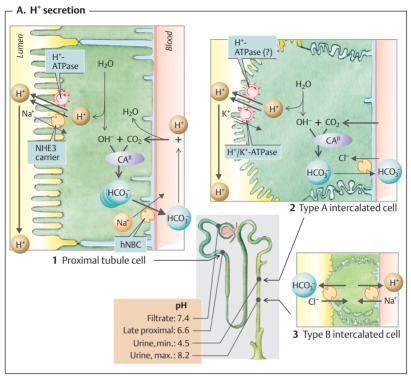
 CO_2 and $H_2O (\rightarrow B)$. CA^{IV} anchored in the membrane catalyzes this reaction. CO2 readily diffuses into the cell, perhaps via aquaporin 1 $(\rightarrow p. 166)$. CA^{II} then catalyzes the transformation of CO₂ + H₂O to H⁺ + HCO₃⁻ within the cell $(\rightarrow B)$. The H⁺ ions are again secreted, while HCO₃⁻ exits through the basolateral membrane of the cell via an electrogenic carrier (hNBC = human Na^+ -bicarbonate co-transporter: $\rightarrow B$). The hNBC co-transports 1 Na+ with 3 HCO₃-(and/or with 1 $HCO_3^- + 1 CO_3^{2-}$?) Thus, $HCO_3^$ is transported through the luminal membrane in the form of CO_2 (driving force: ΔP_{CO_2}), and exits the cell across the basolateral membrane as HCO₃- (main driving force: membrane potential).

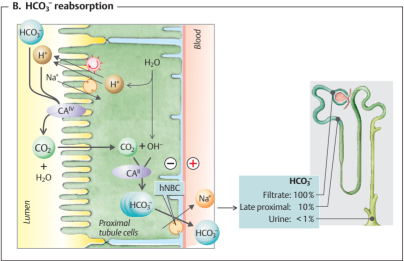
Hypokalemia leads to a rise in membrane potential (Nernst equation, \rightarrow p. 32) and thus to a rise in basolateral HCO_3^- transport. This results in increased H^+ secretion and, ultimately, in **hypokalemic alkalosis**.

Urinary acid excretion. If the dietary protein intake is 70 g per day (→ p. 226), a daily load of about 190 mmol of H⁺ occurs after the amino acids of the protein have been metabolized. HCl (from arginine, lysine and histidine), H₂SO₄ (from methionine and cystine), H₃PO₄, and lactic acid are the main sources of H⁺ ions. They are "fixed" acids which, unlike CO₂, are not eliminated by respiration. Since about 130 mmol H⁺/day are used to break down organic anions (glutamate⁻, aspartate⁻, lactate⁻, etc.), the **net** H⁺ **production** is about **60** (40–80) **mmol/day**. Although the H⁺ ions are buffered at their production site, they must be excreted to regenerate the buffers.

In extreme cases, the **urinary pH** can rise to about pH 8 (high HCO_3^- excretion) or fall to about pH 4.5 (maximum H^+ conc. is 0.03 mmol/L). At a daily urine output of 1.5 L, the kidneys will excrete < 1% of the produced H^+ ions in their free form.

Titratable acids (80% phosphate, 20% uric acid, citric acid, etc.) comprise a significant fraction (10–30 mmol/day) of H⁺ excretion (→C1). This amount of H⁺ ions can be determined by titrating the urine with NaOH back to the plasma pH value, which is normally pH 7.4 (→C2). Around 80% of **phosphate** (pK_a = 6.8) in the blood occurs in the form of HPO₄²⁻, whereas about all phosphate in acidic urine occurs as H₂PO₄⁻ (→p, 380), i.e., the secreted





 H^+ ions are buffered by filtered HPO_4^{2-} . Non-reabsorbed phosphate (5–20% of the filtered quantity, \rightarrow p. 178) is therefore loaded with H^+ ions, about half of it in the proximal tubule (pH 7.4 \Rightarrow ca. 6.6), and the rest in the collecting duct (pH 6.6 \Rightarrow 4.5) (\rightarrow C1). When *acidosis* occurs, increased quantities of phosphate are mobilized from the bone and excreted. The resulting increase in H^+ excretion precedes the increased NH_4^+ production associated with acidosis (see below).

Excretion of ammonium ions (NH₄ $^+$ NH₃ + H $^+$), about 25–50 mmol/day on average diet, is *equivalent* to H $^+$ disposal and is therefore an *indirect* form of H $^+$ excretion (\rightarrow **D**). NH₄ $^+$ is *not* a titratable form of acidity. Unlike HPO₄²⁻ + H $^+$

 $H_2PO_4^-$, the reaction $NH_3 + H^+$ NH_4^+ does not function in the body as a buffer because of its high pK_a value of ca. 9.2. Nevertheless, for every NH_4^+ excreted by the kidney, one HCO_3^- is spared by the liver. This is equivalent to one H^+ disposed since the spared HCO_3^- ion can buffer a H^+ ion. With an average dietary intake of protein, the amino acid metabolism produces roughly equimolar amounts of HCO_3^- and NH_4^+ (ca. 700-1000 mmol/day). The liver utilizes about 95% of these two products to produce $\mathbf{urea} (\to \mathbf{D1})$:

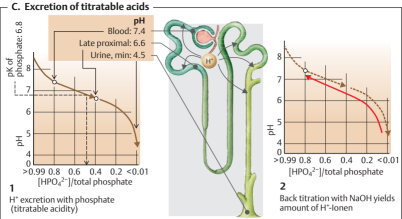
Thus, one HCO₃⁻ less is consumed for each NH₄⁺ that passes from the liver to the kidney and is eliminated in the urine. Before exporting NH₄⁺ to the kidney, the **liver** incorporates it into glutamate yielding **glutamine**; only a small portion reaches the kidney as free NH₄⁺. High levels of NH₄⁺ NH₃ are toxic.

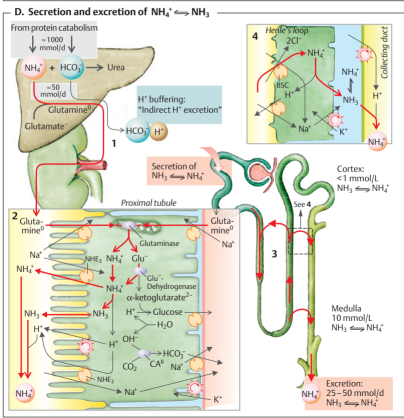
In the **kidney**, *glutamine* enters proximal tubule cells by Na⁺ symport and is cleaved by mitochondrial *glutaminase*, yielding NH₄⁺ and glutamate⁻ (Glu⁻). Glu⁻ is further metabolized by *glutamate dehydrogenase* to yield α -ketoglutarate²⁻, producing a second NH₄⁺ ion (\rightarrow **D2**). The NH₄⁺ can reach the tubule lumen on two ways: (1) it dissociates within the cell to yield NH₃ and H⁺, allowing NH₃ to diffuse (non-ionically, \rightarrow p. 22) into the lumen, where it re-joins the separately secreted H⁺ ions; (2) the NHE3 carrier secretes NH₄⁺ (instead of H⁺). Once NH₄⁺ has arrived at *the thick ascending limb* of the loop of Henle (\rightarrow **D4**), the BSC car-

rier (\rightarrow p. 162) reabsorbs NH₄⁺ (instead of K⁺) so that it remains in the renal medulla. Recirculation of NH₄⁺ through the loop of Henle yields a very high conc. of NH₄+ NH3 + H+ towards the papilla (\rightarrow **D3**). While the H⁺ ions are then actively pumped into the lumen of the collecting duct (\rightarrow **A2, D4**), the NH₃ molecules arrive there by non-ionic diffusion (\rightarrow **D4**). The NH₃ gradient required to drive this diffusion can develop because the especially low luminal pH value (about 4.5) leads to a much smaller NH3 conc. in the lumen than in the medullary interstitium where the pH is about two pH units higher and the NH3 conc. is consequently about 100-times higher than in the lumen.

Disturbances of acid-base metabolism (see also p. 142ff.). When chronic non-respiratory acidosis of non-renal origin occurs, NH4+ excretion rises to about 3 times the normal level within 1 to 2 days due to a parallel increase in hepatic glutamine production (at the expense of urea formation) and renal alutaminase activity. Non-respiratory alkalosis only decreases the renal NH₄⁺ production and H⁺ secretion. This occurs in conjunction with an increase in filtered HCO₃- (increased plasma concentration, → p. 144), resulting in a sharp rise in HCO₃⁻ excretion and, consequently, in osmotic diuresis (\rightarrow p. 172). To compensate for respiratory disturbances (→p. 144), it is important that increased (or decreased) plasma Pco, levels result in increased (or decreased) H⁺ secretion and, thus, in increased (or decreased) HCO₃⁻ resorption.

The kidney can also be the primary site of an acidbase disturbance (renal acidosis), with the defect being either generalized or isolated. In a generalized defect, as observed in renal failure, acidosis occurs because of reduced H* excretion. In an isolated defect with disturbance of proximal H* secretion, large portions of filtered HCO₃⁻ are not reabsorbed, leading to proximal renal tubular acidosis. When impaired renal H* secretion occurs in the collecting duct, the urine can no longer be acidified (pH > 6 despite acidosis) and the excretion of titratable acids and NH₄* is consequently impaired (distal renal tubular acidosis).





Reabsorption and Excretion of Phosphate, Ca²⁺ and Mg²⁺

Phosphate metabolism. The plasma phosphate conc. normally ranges from 0.8-1.4 mmol/L, and a corresponding amount of ca. 150-250 mmol/day of inorganic phosphate P_i (HPO₄²-H₂PO₄⁻) is filtered each day, a large part of which is reabsorbed. The fractional excretion (\rightarrow **A1**), which ranges between 5 and 20%, functions to balance Pi, H+, and Ca^{2+} . P_i excretion rises in the presence of a P_i excess (elevated Pi levels in plasma) and falls during a Pi deficit. Acidosis also results in phosphaturia and increased H+ excretion (titratable acidity, \rightarrow p. 174ff.). This also occurs in phosphaturia of other causes. Hypocalcemia and parathyrin also induce a rise in Pi excretion $(\rightarrow$ A3 and p. 290f.).

 P_1 is reabsorbed at the proximal tubule (\rightarrow A2,3). Its luminal membrane contains the type 3 Na^+ - P_1 symport carrier (NaPi-3). The carrier accepts $H_2PO_4^-$ and HPO_4^{2-} and cotransports it with Na $^+$ by secondary active transport (\rightarrow p. 26ff.).

Regulation of P_i reabsorption. P_i deficits, alkalosis, hypercalcemia, and low PTH levels result in the increased incorporation of NaPi-3 transporters into the luminal membrane, whereas P_i excesses, acidosis, hypocalcemia and increased PTH secretion results in internalization (down-regulation) and subsequent lysosomal degradation of NaPi-3 (\rightarrow A3).

Calcium metabolism (see also p. 36). Unlike the Na* metabolism, the calcium metabolism is regulated mainly by absorption of Ca^{2+} in the gut (\rightarrow p. 290ff.) and, secondarily, by renal excretory function. *Total plasma calcium* (bound calcium + ionized Ca^{2+}) is a mean 2.5 mmol/L. About 1.3 mmol/L of this is present as *free*, *ionized* Ca^{2+} , 0.2 mmol/L forms *complexes* with phosphate, citrate, etc., and the rest of 1 mmol/L is *bound to plasma proteins* and, thus, not subject to glomerular filtration (\rightarrow p. 154). *Fractional excretion of* Ca^{2+} (FE_{Ca}) in the urine is 0.5%–3% (\rightarrow A1).

 Ca^{2+} reabsorption occurs practically throughout the entire nephron (\rightarrow A1,2). The reabsorption of filtered Ca^{2+} occurs to about 60% in the proximal tubule and about 30% in the thick ascending limb (TAL) of the loop of

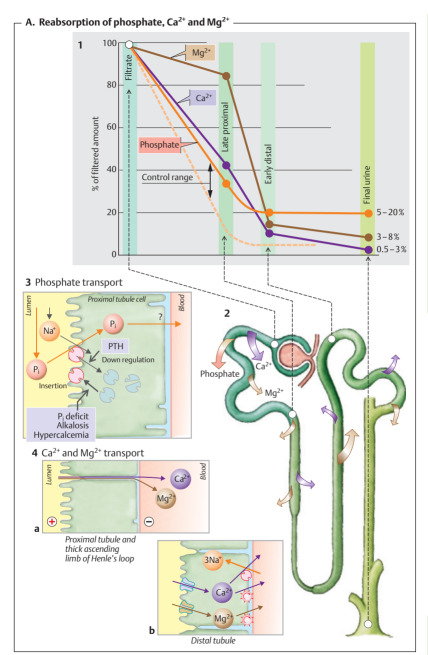
Henle and is paracellular, i.e., passive (\rightarrow A4a and p. 163 B5, B7). The lumen-positive transepithelial potential (LPTP) provides most of the driving force for this activity. Since Ca2+ reabsorption in TAL depends on NaCl reabsorption, loop diuretics (\rightarrow p. 172) inhibit Ca²⁺ reabsorption there. **PTH** promotes Ca²⁺ reabsorption in TAL as well as in the distal convoluted tubule, where Ca2+ is reabsorbed by transcellular active transport (\rightarrow **A4b**). Thereby, Ca²⁺ influx into the cell is passive and occurs via luminal Ca²⁺ channels, and Ca²⁺ efflux is active and occurs via Ca2+-ATPase (primary active Ca2+ transport) and via the 3 Na⁺/1 Ca²⁺ antiporter (secondary active Ca²⁺ transport). Acidosis inhibits Ca2+ reabsorption via unclear mechanisms.

Urinary calculi usually consist of *calcium phosphate* or *calcium oxalate*. When Ca²⁺, Pi or oxalate levels are increased, the solubility product will be exceeded but calcium complex formers (e.g., citrate) and inhibitors of crystallization (e.g., nephrocalcin) normally permit a certain degree of supersaturation. Stone formation can occur if there is a deficit of these substances or if extremely high urinary concentrations of Ca²⁺, Pi and oxalate are present (applies to all three in pronounced antidiuresis).

Magnesium metabolism and **reabsorption**. Since part of the magnesium in plasma (0.7-1.2 mmol/L) is protein-bound, the Mg²⁺ conc. in the filtrate is only 80% of the plasma magnesium conc. Fractional excretion of Mg²⁺, FE_{Mg}, is 3-8% (\rightarrow **A1,2**). Unlike Ca²⁺, however, only about 15% of the filtered Mg²⁺ ions leave the proximal tubule. About 70% of the Mg²⁺ is subject to paracellular reabsorption in the TAL (\rightarrow **A4** and p. 163 B5, B7). Another 10% of the Mg²⁺ is subject to transcellular reabsorption in the distal tubule (\rightarrow **A4b**), probably like Ca²⁺ (see above).

Mg²⁺ excretion is stimulated by hypermagnesemia, hypercalcemia, hypervolemia and loop diuretics, and is inhibited by Mg²⁺ deficit, Ca²⁺ deficit, volume deficit, PTH and other hormones that mainly act in the TAL.

The kidney has **sensors** for divalent cations like Ca^{2^+} and Mg^{2^+} (\rightarrow p. 36). When activated, the sensors inhibit NaCl reabsorption in the TAL which, like loop diuretics, reduces the driving force for paracellular cation resorption, thereby diminishing the normally pronounced Mg^{2^+} reabsorption there.



Potassium Balance

The dietary intake of K⁺ is about 100 mmol/day (minimum requirement: 25 mmol/day), About 90% of intake is excreted in the urine, and 10% is excreted in the feces. The plasma K⁺ conc. normally ranges from 3.5 to 4.8 mmol/L, while intracellular K+ conc. can be more than 30 times as high (due to the activity of Na+-K+-ATPase: \rightarrow **A**). Therefore, about 98% of the ca. 3000 mmol of K⁺ ions in the body are present in the cells. Although the extracellular K⁺ conc. comprises only about 2% of total body K+, it is still very important because (a) it is needed for regulation of K+ homeostasis and (b) relatively small changes in cellular K+ (influx or efflux) can lead to tremendous changes in the plasma K+ conc. (with an associated risk of cardiac arrhythmias). Regulation of K+ homeostasis therefore implies distribution of K+ through intracellular and extracellular compartments and adjustment of K+ excretion according to K+ intake.

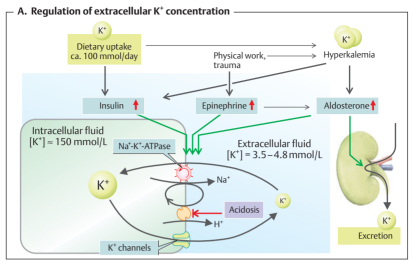
Acute regulation of the extracellular K+ conc. is achieved by internal shifting of K+ between the extracellular fluid and intracellular fluid (\rightarrow A). This relatively rapid process prevents or mitigates dangerous rises in extracellular K+ (hyperkalemia) in cases where large quantities of K⁺ are present due to high dietary intake or internal K+ liberation (e.g., in sudden hemolysis). The associated K+ shifting is mainly subject to hormonal control. The insulin secreted after a meal stimulates Na+-K+-ATPase and distributes the K⁺ supplied in the animal and vegetable cells of the food to the cells of the body. This is also the case in diet-independent hyperkalemia, which stimulates insulin secretion per se. Epinephrine likewise increases cellular K⁺ uptake, which is particularly important in muscle work and traumatwo situations that lead to a rise in plasma K⁺. In both cases, the increased epinephrine levels allow the re-uptake of K⁺ in this and other cells. Aldosterone also increases the intracellular K + conc. (see below).

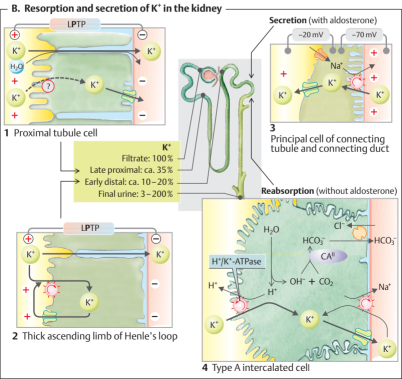
Changes in pH affect the intra- and extracellular distribution of $K^+(\rightarrow A)$. This is mainly because the ubiquitous Na^+/H^+ antiporter works faster in alkalosis and more slowly in acidosis $(\rightarrow A)$. In acidosis, Na^+ influx therefore decreases, Na^+-K^+ -ATPase slows down, and the

extracellular K^+ concentration rises (especially in non-respiratory acidosis, i.e., by $0.6\,\mathrm{mmol/L}$ per 0.1 unit change in pH). Alkalosis results in hypokalemia.

Chronic regulation of K+ homeostasis is mainly achieved by the **kidney** (\rightarrow **B**). K⁺ is subject to free glomerular filtration, and most of the filtered K+ is normally reabsorbed (net reabsorption). The excreted amount can, in some cases, exceed the filtered amount (net secretion, see below). About 65% of the filtered K⁺ is reabsorbed before reaching the end of the proximal tubule, regardless of the K+ supply. This is comparable to the percentage of Na⁺ and H_2O reabsorbed (\rightarrow **B1** and p. 157, column 2). This type of K+ transport is mainly paracellular and therefore passive. Solvent drag $(\rightarrow p.24)$ and the lumen-positive transepithelial potential, LPTP (\rightarrow B1 and p. 162), in the mid and late proximal segments of the tubule provide the driving forces for it. In the loop of Henle, another 15% of the filtered K+ is reabsorbed by trans- and paracellular routes $(\rightarrow B2)$. The amount of K⁺ excreted is determined in the connecting tubule and collecting duct, Larger or smaller quantities of K⁺ are then either reabsorbed or secreted according to need. In extreme cases, the fractional excretion of K+ (FEK) can rise to more than 100% in response to a high K+ intake, or drop to about 3–5% when there is a K^+ deficit ($\rightarrow B$).

Cellular mechanisms of renal K+ transport. The connecting tubule and collecting duct contain **principal cells** $(\rightarrow B3)$ that reabsorb Na⁺ and secrete K+. Accumulated intracellular K+ can exit the cell through K+ channels on either side of the cell. The electrochemical K+ gradient across the membrane in question is decisive for the efflux of K+. The luminal membrane of principal cells also contains Na+ channels through which Na⁺ enters the cell (\rightarrow p. 162). This depolarizes the luminal membrane. which reaches a potential of about -20 mV, while the basolateral membrane maintains its normal potential of ca. $-70 \,\mathrm{mV} \ (\rightarrow B3)$. The driving force for K^+ efflux $(E_m - E_K, \rightarrow p. 32)$ is therefore higher on the luminal side than on the opposite side. Hence, K+ preferentially exits the cell toward the lumen (secretion). This is mainly why K⁺ secretion is coupled with Na⁺ reabsorption, i.e., the more Na+ reabsorbed by the principle cell, the more K⁺ secreted.





Another apparent reason is that the reabsorption-related increase in intracellular Na⁺ concentration decreases the driving force for the 3 Na⁺/Ca²⁺ exchange at the basolateral cell membrane, resulting in a rise in the **cytosolic Ca²⁺** concentration. This rise acts as a signal for more frequent opening of luminal K⁺ channels.

Type A **intercalated cells** (\rightarrow **B4**) can active reabsorb K⁺ in addition to secreting H⁺ ions. Like the parietal cells of the stomach, their luminal membrane contains a H^+/K^+ -ATPase for this purpose.

Factors that affect K^+ excretion (\rightarrow C):

- 1. An increased K^+ intake raises the intracellular and plasma K^+ concentrations, which thereby increases the chemical driving force for K^+ secretion.
- **2. Blood pH**: The intracellular K⁺ conc. in renal cells *rises in alkalosis* and *falls in acute acidosis*. This leads to a simultaneous fall in K⁺ excretion, which again rises in chronic acidosis. The reasons for this are that (a) acidosis-related inhibition of Na⁺-K⁺-ATPase reduces proximal Na⁺ reabsorption, resulting in increased distal urinary flow (see no. 3), and (b) the resulting hyperkalemia stimulates aldosterone secretion (see no. 4).
- **3.** If there is **increased urinary flow** in the connecting tubule and collecting duct (e.g., due to a high Na $^+$ intake, osmotic diuresis, or other factors that inhibit Na $^+$ reabsorption upstream), larger quantities of K $^+$ will be excreted. This explains the *potassium-losing effect of certain diuretics* (\rightarrow p. 173). The reason for this is, presumably, that K $^+$ secretion is limited at a certain luminal K $^+$ concentration. Hence, the larger the volume/time, the more K $^+$ taken away over time.
- **4. Aldosterone** leads to retention of Na^+ , an increase in extracellular volume (\rightarrow p. 184), a moderate increase in H $^+$ secretion (cellular pH rises), and *increased K^+ excretion*. It also increases the number of Na^+ -K $^+$ -ATPase molecules in the target cells and leads to a chronic increase in mitochondrial density in K $^+$ adaptation, for example (see below).

Cellular mechanisms of aldosterone effects. The increase in Na⁺ reabsorption is achieved by increased production of transport proteins, called aldosterone-induced proteins (AIPs). This is a genome-mediated effect that begins approx. 30 min to 1 hour after al-

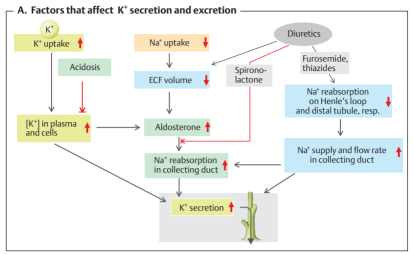
dosterone administration or secretion. The maximum effects are observed after several hours. Aldosterone increases Na* reabsorption, thereby depolarizing the luminal cell membrane (\rightarrow B3). Consequently, it increases the driving force for K* secretion and increases K* conductance by increasing the pH of the cell. Both effects lead to increased K* secretion. Aldosterone also has a very rapid (few seconds to minutes) non-genomic effect on the cell membrane, the physiological significance of which has yet to be explained.

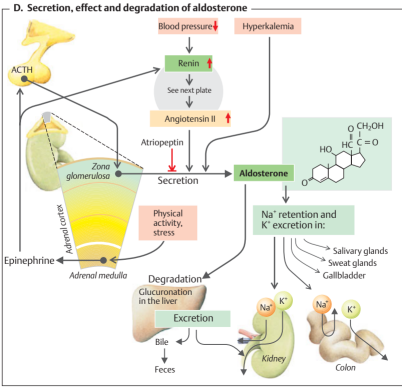
The capacity of the K^* excretory mechanism increases in response to long-term increases in the K^* supply (K^* adaptation). Even when renal function is impaired, this largely maintains the K^* balance in the remaining, intact parts of the tubular apparatus. The colon can then take over more than $^1/_3$ of the K^* excretion.

Mineralocortico(stero)ids. Aldosterone is the main mineralocorticoid hormone synthesized and secreted by the *zona glomerulosa of the adrenal cortex* (→ \mathbf{D} and p.294ff.). As with other steroid hormones, aldosterone is not stored, but is synthesized as needed. The principal function of aldosterone is to regulate Na⁺ and K⁺ transport in the kidney, gut, and other organs (→ \mathbf{D}). Aldosterone secretion increases in response to (a) drops in blood volume and blood pressure (mediated by angiotensin II; → \mathbf{p} . 184) and (b) hyperkalemia (→ \mathbf{D}). Aldosterone synthesis is inhibited by atriopeptin (→ \mathbf{p} , 171 A4).

Normal **cortisol** concentrations are *not* effective at the aldosterone receptor only because cortisol is converted to cortisone by an 11β -hydroxysteroid oxidoreductase in aldosterone's target cells.

Hyperaldosteronism can be either primary (aldosterone-secreting tumors of adrenal cortex, as observed in Conn's syndrome) or secondary (in volume depletion, \rightarrow p. 184). Na⁺ retention resulting in high ECF volumes and hypertension as well as a simultaneous K^+ losses and hypokalemic alkalosis are the consequences. When more than about 90% of the adrenal cortex is destroyed, e.g. by autoimmune adrenalitis, metastatic cancer or tuberculosis, primary chronic adrenocortical insufficiency develops (Addison's disease). The aldosterone deficit leads to a sharp increase in Na+ excretion, resulting in hypovolemia, hypotension and K⁺ retention (hyperkalemia). As glucocorticoid deficiency also develops, complications can be life-threatening, especially under severe stress (infections, trauma). If only one gland is destroyed, ACTH causes hypertrophy of the other (\rightarrow p. 297 A).





Tubuloglomerular Feedback, Renin–Angiotensin System

The **juxtaglomerular apparatus** (**JGA**) consists of (a) juxtaglomerular cells of the *afferent arteriole* (including renin-containing and sympathetically innervated granulated cells) and *efferent arteriole*, (b) *macula densa cells* of the thick ascending limb of the loop of Henle and (c) juxtaglomerular mesangial cells (*polkissen*, \rightarrow **A**) of a given nephron (\rightarrow **A**).

JGA functions: (1) local transmission of tubuloglomerular feedback (TGF) at its own nephron via angiotensin II (ATII) and (2) systemic production of angiotensin II as part of the renin–angiotensin system (RAS).

Tubuloglomerular feedback (TGF). Since the daily GFR is 10 times larger than the total ECF volume $(\rightarrow p. 168)$, the excretion of salt and water must be precisely adjusted according to uptake. Acute changes in the GFR of the individual nephron (iGFR) and the amount of NaCl filtered per unit time can occur for several reasons. An excessive iGFR is associated with the risk that the distal mechanisms for NaCl reabsorption will be overloaded and that too much NaCl and H2O will be lost in the urine. A too low iGFR means that too much NaCl and H2O will be retained. The extent of NaCl and H₂O reabsorption in the proximal tubule determines how quickly the tubular urine will flow through the loop of Henle. When less is absorbed upstream, the urine flows more quickly through the thick ascending limb of the loop, resulting in a lower extent of urine dilution $(\rightarrow p. 162)$ and a higher NaCl concentration at the macula densa. [NaCl]_{MD}. If the [NaCl]_{MD} becomes too high, the afferent arteriole will constrict to curb the GFR of the affected nephron within 10 s or vice versa (negative feedback). It is unclear how the [NaCl]_{MD} results in the signal to constrict, but type 1 A angiotensin II (AT_{1A}) receptors play a key role.

If, however, the [NaCl]_{MD} changes due to **chronic shifts in total body NaCl** and an associated change in ECF volume, rigid coupling of the iGFR with the [NaCl]_{MD} through TGF would be fatal. Since long-term increases in the ECF volume reduce proximal NaCl reabsorption, the [NaCl]_{MD} would increase, resulting in a decrease in the GFR and a further increase in the ECF volume. The reverse occurs in ECF volume deficit. To prevent these effects, the [NaCl]_{MD}/iGFR response curve is shifted in the appropriate direction by certain substances. Nitric oxide (NO) shifts the curve when there is an ECF excess (increased iGFR at same [NaCl]_{MD}), and (only locally effective) angiotensin II shifts it in the other direction when there is an ECF deficit.

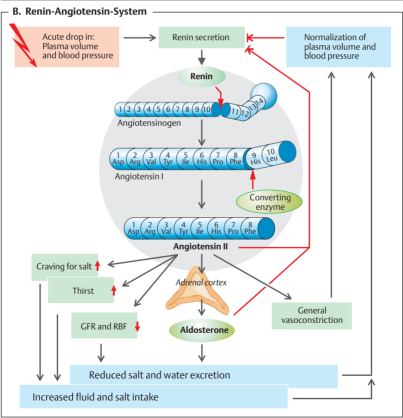
Renin-angiotensin system (RAS). If the mean renal blood pressure acutely drops below 90 mmHg or so, renal baroreceptors will trigger the release of renin, thereby increasing the systemic plasma renin conc. Renin is a peptidase that catalyzes the cleavage of angiotensin I from the renin substrate angiotensinogen (released from the liver). Angiotensin-converting enzyme (ACE) produced in the lung, etc. cleaves two amino acids from angiotensin I to produce angiotensin I approx. 30-60 minutes after the drop in blood pressure (\rightarrow B).

Control of the RAS $(\rightarrow$ **B**). The blood pressure *threshold* for renin release is raised by α_1 -adrenoceptors, and *basal renin secretion* is increased by β_1 -adrenoceptors. *Angiotensin II* and *aldosterone* are the most important effectors of the RAS. Angiotensin II stimulates the release of aldosterone by adrenal cortex (see below). Both hormones directly (fast action) or indirectly (delayed action) lead to a renewed increase in arterial blood pressure $(\rightarrow$ **B**), and renin release therefore decreases to normal levels. Moreover, both hormones *inhibit renin release* (negative feedback).

If the mean blood pressure is decreased in only one kidney (e.g., due to stenosis of the affected renal artery), the affected kidney will also start to release more renin which, in this case, will lead to **renal hypertension** in the rest of the circulation.

Angiotensin II effects: Beside altering the structure of the myocardium and blood vessels (mainly via AT_2 receptors), angiotensin II has the following fast or delayed effects mediated by AT_1 receptors ($\rightarrow A$).

- Vessels: Angiotensin II has potent vasoconstrictive and hypertensive action, which (via endothelin) takes effect in the arterioles (fast action).
- CNS: Angiotensin II takes effect in the hypothalamus, resulting in vasoconstriction through the circulatory "center" (rapid action). It also increases ADH secretion in the hypothalamus, which stimulates thirst and a craving for salt (delayed action).
- ♠ Kidney: Angiotensin II plays a major role in regulating renal circulation and GFR by constricting of the afferent and/or efferent arteriole (delayed action; cf. autoregulation, → p. 150). It directly stimulates Na* reabsorption in the proximal tubule (delayed action).
- Adrenal gland: Angiotensin II stimulates aldosterone synthesis in the adrenal cortex (delayed action; \rightarrow p. 182) and leads to the release of epinephrine in the adrenal medulla (fast action).



Cardiovascular System

Overview

Blood is pumped from the left ventricle of the heart to capillaries in the periphery via the arterial vessels of the *systemic* (or greater) *circulation* and returns via the veins to the right heart. It is then expelled from the right ventricle to the lungs via the *pulmonary* (or lesser) *circulation* and returns to the left heart $(\rightarrow A)$.

The total **Blood volume** is roughly 4-5 L $(\approx 7\%)$ of the fat-free body mass: \rightarrow table on p.88). Around 80% of the blood circulates through the veins, right heart and pulmonary vessels, which are jointly referred to as the low pressure system ($\rightarrow A$, left). These highly distensible capacitance vessels function as a blood reservoir in which blood is stored and released as needed via venous vasoconstriction (\rightarrow e.g., p. 218). When the blood volume increasesdue, for example, to a blood transfusion—over 99% of the transfused volume remains in the low-pressure system (high capacitance), while only < 1% circulates in the arterial high-pressure system (low capacitance). Conversely, a decrease will be reflected almost entirely by a decrease in the blood stores in the lowpressure system. Central venous pressure (measured in or near to the right atrium; normally 4-12 cm H₂O) is therefore a good indicator of blood volume (and ECF volume) in individuals with a normally functioning heart and lungs.

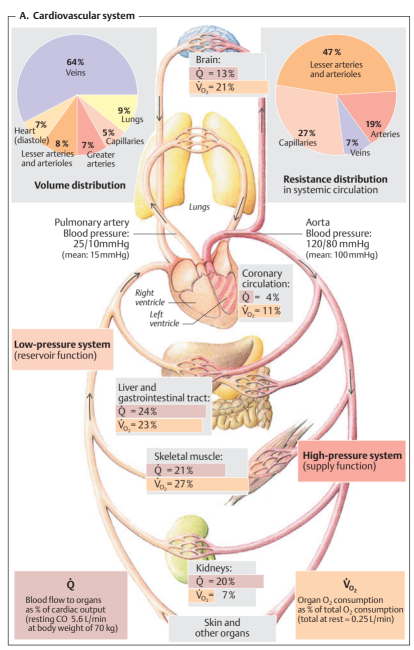
Cardiac output (CO). The cardiac output is calculated as heart rate (HR) times stroke volume (SV). Under normal resting conditions, the CO is approx. $70 \, [\text{min}^{-1}] \times 0.08 \, [\text{L}] = 5.6 \, \text{L/min}$ or, more precisely, a mean $3.4 \, \text{L/min}$ per m^2 body surface area. An increase in HR (up to about $180 \, \text{min}^{-1}$) and/or SV can increase the CO to $15-20 \, \text{L/min}$.

The **distribution of blood** to the organs arranged in *parallel* in the **systemic circulation** (\rightarrow **A**, \dot{Q} values) is determined by their *functional priority* (vital organs) and by the *current needs* of the body (see also p.213 A). Maintaining adequate cerebral perfusion (approx. 13% of the resting CO) is the top priority, not only because the *brain* is a major vital organ, but also because it is very susceptible to

hypoxic damage (\rightarrow p. 130). Myocardial perfusion via coronary arteries (approx. 4% of the CO at rest) must also be maintained, because any disruption of cardiac pumping function will endanger the entire circulation. About 20 to 25% of the CO is distributed to the kidneys. This fraction is very large relative to the kidney weight (only 0.5% of body mass). Renal blood flow is primarily used to maintain renal excretory and control functions. Thus, renal blood flow may be reduced transiently in favor of cardiac and cerebral perfusion, e.g., to ward off impending shock (\rightarrow p. 218). During strenuous physical exercise, the CO increases and is alloted mainly to the skeletal muscle. During digestion, the gastrointestinal tract also receives a relatively high fraction of the CO. Naturally, both of these organ groups cannot receive the maximum blood supply at the same time (\rightarrow p. 75 A). Blood flow to the skin (approx. 10% of the resting CO) mainly serves the purpose of heat disposal (\rightarrow p. 222ff.). The cutaneous blood flow rises in response to increased heat production (physical work) and/ or high external temperatures and decreases (pallor) in favor of vital organs in certain situations (e.g., shock; \rightarrow p. 218).

The total CO flows through the **pulmonary circulation** as it and the systemic circulation are arranged in series (\rightarrow **A**). Oxygen-depleted (venous) blood is carried via the pulmonary arteries to the lungs, where it is *oxygenated* or "arterialized." A relatively small quantity of additional oxygenated blood from the systemic circulation reaches the lung tissue via the bronchial arteries. All blood in the pulmonary circulation drains via the pulmonary veins.

Peripheral resistance. Flow resistance in the pulmonary circulation is only about 10% of the *total peripheral resistance (TPR)* in the systemic circulation. Consequently, the mean pressure in the right ventricle (approx. 15 mmHg = $2 \, \text{kPa}$) is considerably lower than in the left ventricle ($100 \, \text{mmHg} = 13.3 \, \text{kPa}$). Since the resistance in the lesser arteries and arterioles amounts to nearly 50% of $\text{TPR} (\rightarrow \textbf{A}, \text{top right})$, they are called *resistance vessels*.



Blood Vessels and Blood Flow

In the systemic circulation, blood is ejected from the left ventricle to the aorta and returns to the right atrium via the venae cavae (\rightarrow A). As a result, the **mean blood pressure** (\rightarrow p. 206) drops from around 100 mmHg in the aorta to 2–4 mmHg in the venae cavae (\rightarrow A2), resulting in a pressure difference (Δ P) of about 97 mmHg (pulmonary circulation; \rightarrow p. 122). According to **Ohm's Law**,

 $\Delta P = \tilde{Q} \cdot R \text{ (mmHg)}$ [8.1] where Q is the blood flow $(L \cdot min^{-1})$ and R is the flow resistance $(mmHg \cdot min \cdot L^{-1})$. Equation 8.1 can be used to calculate blood flow in a given organ (R = organ resistance) as well as in the entire cardiovascular system, where Q is the cardiac output $(CO; \rightarrow p. 186)$ and R is the total peripheral flow resistance (TPR). The TPR at rest is about $18 \text{ mmHg} \cdot min \cdot L^{-1}$.

The aorta and greater arteries distribute the blood to the periphery. In doing so, they act as a hydraulic filter because (due to their high compliance, $\Delta V/\Delta P_{tm}$) they convert the intermittent flow generated by the heart to a nearly steady flow through the capillaries. The high systolic pressures generated during the ejection phase cause the walls of these blood vessels to stretch, and part of the ejected blood is "stored" in the dilated lumen (windkessel). Elastic recoil of the vessel walls after aortic valve closure maintains blood flow during diastole. Arterial vessel compliance decreases with age.

Flow velocity (\dot{V}) and flow rate (\dot{Q}) of the blood. Assuming an aortic cross-sectional area (CSA) of 5.3 cm² and a total CSA of 20 cm² of all downstream arteries (\rightarrow A5), the mean \dot{V} (during systole and diastole) at rest can be calculated from a resting CO of 5.6 L/min: It equals 18 cm/s in the aorta and 5 cm/s in the arteries (\rightarrow A3). As the aorta receives blood only during the ejection phase (\rightarrow p. 90), the maximum resting values for \dot{V} and \dot{Q} in the aortic root are much higher during this phase (\dot{V} = 95 cm/s, \dot{Q} = 500 mL/s).

In the Hagen-Poiseuille equation,

 $\mathbf{R} = \mathbf{8} \cdot \mathbf{l} \cdot \mathbf{\eta} / (\pi \cdot \mathbf{r^4})$ [8.2] the flow resistance (R) in a tube of known length (I) is dependent on the viscosity ($\mathbf{\eta}$) of the fluid in the tube (\rightarrow p.92) and the fourth

power of the inner radius of the tube (r⁴). Decreasing the radius by only about 16% will therefore suffice to double the resistance.

The **lesser arteries** and **arterioles** account for nearly 50% of the TPR (*resistance vessels*; \rightarrow **A1** and p. 187 A) since their small radii have a much stronger effect on total TPR ($R \sim 1/r^4$) than their large total CSA ($R \sim r^2$). Thus, the blood pressure in these vessels drops significantly. Any change in the radius of the small arteries or arterioles therefore has a radical effect on the TPR (\rightarrow p. 212ff.). Their width and that of the **precapillary sphincter** determines the amount of blood distributed to the capillary beds (exchange area).

Although the **capillaries** have even smaller radii (and thus much higher individual resistances than the arterioles), their total contribution to the TPR is only about 27% because their total CSA is so large (\rightarrow A1 and p. 187 A). The *exchange of fluid and solutes* takes place across the walls of capillaries and **postcapillary venules**. Both vessel types are particularly suirable for the task because (a) their \dot{V} is very small (0.02–0.1 cm/s; \rightarrow A3) (due to the large total CSA), (b) their total surface area is very large (approx. 1000 m²), (3) and their walls can be very thin as their inner radius (4.5 μ m) is extremely small (*Laplace's law*, see below).

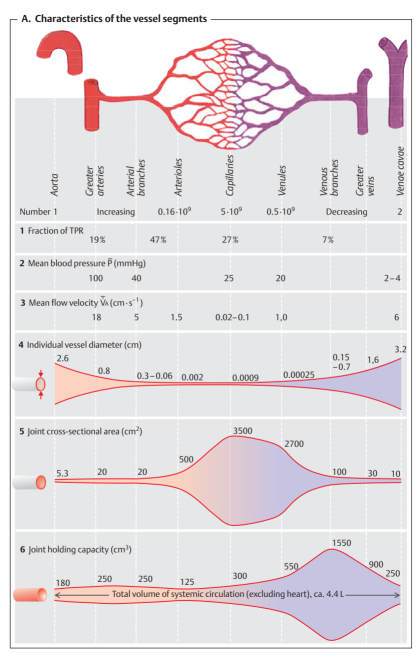
Transmural pressure P_{tm} [N/m²], that is, the pressure difference across the wall of a hollow organ (=internal pressure minus external pressure), causes the wall to stretch. Its materials must therefore be able to withstand this stretch. The resulting tangential *mural tension* T [N/m] is a function of the inner radius r [m] of the organ. According to **Laplace's law** for cylindrical (or spherical) hollow bodies,

$$P_{tm} = T/r \text{ (or } P_{tm} = 2T/r, \text{ resp.)}$$
 [8.3a/b]

Here, T is the total mural tension, regardless how thick the wall is. A thick wall can naturally withstand a given P_{tm} more easily than a thin one. In order to determine the **tension** exerted **per unit CSA of the wall** (i.e., the stress requirements of the wall material in N/m²), the thickness of the wall (w) must be considered. Equation 8.3 a/b is therefore transformed to

 $P_{tm} = T \cdot w/r$ (or $P_{tm} = 2 T \cdot w/r$, resp.) [8.4a/b]

The blood collects in the **veins**, which can accommodate large volumes of fluid (\rightarrow **A6**). These *capacitance vessels* serve as a *blood reservoir* (\rightarrow p. 186).



Cardiac Cycle

The resting heart rate is 60–80 beats per minute. A **cardiac cycle** (\rightarrow **A**) therefore takes roughly 1 s. It can be divided into four distinct phases: (I) contraction phase and (II) ejection phase, both occurring in *systole*; (III) relaxation phase and filling phase (IV), both occurring in *diastole*. At the end of phase IV, the atria contract (phase IVc). Electrical excitation of the atria and ventricles precedes their contraction.

The **cardiac valves** determine the direction of blood flow within the heart, e.g., from the atria to the ventricles (phase IV) or from the ventricles to the aorta or pulmonary artery (phase II). All cardiac valves are closed during phases I and III (\rightarrow **A**, top). Opening and closing of the valves is controlled by the pressures exerted on the two sides of the valves.

Cardiac cycle. Near the end of ventricular diastole, the sinoatrial (SA) node emits an **electrical impulse**, marking to the beginning of the P wave of the ECG (phase IVc, \rightarrow A1 and p. 196ff.). This results in *atrial contraction* (\rightarrow A4) and is followed by *ventricular excitation* (QRS complex of the ECG). The ventricular pressure then starts to rise (\rightarrow A2, blue line) until it exceeds the atrial pressure, causing the atrioventricular valves (mitral and tricuspid valves) to close. This marks the *end of diastole*. The mean *end-diastolic volume* (**EDV**) in the ventricle is now about 120 mL (\rightarrow A4) or, more precisely, 70 mL/m² body surface area.

The **isovolumetric contraction** phase now begins (phase I, ca. 50 ms). With all valves are closed, the ventricles now contract, producing the **first heart sound** (\rightarrow **A6**), and the ventricular pressure increases rapidly. The slope of this ascending pressure curve indicates the maximum rate of pressure developed (**maximum dP/dt**). The semilunar valves (aortic and pulmonary valves) now open because the pressure in the left ventricle (\rightarrow **A2**, blue line) exceeds that in the aorta (black broken curve) at about 80 mmHg, and the pressure in the right ventricle exceeds that in the pulmonary artery at about 10 mmHg.

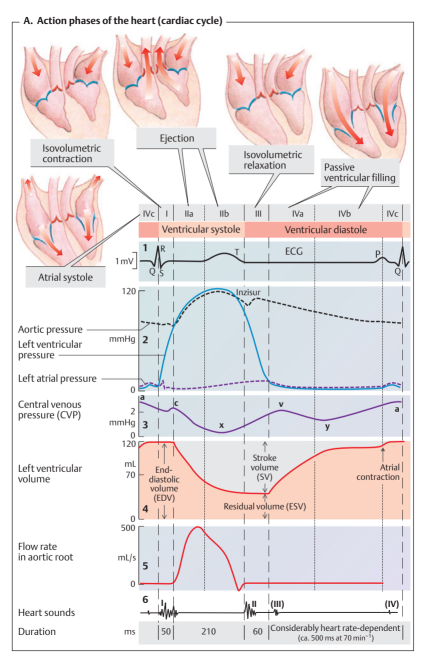
The **ejection** phase (now begins phase II; ca. 210 ms at rest). During this period, the pressure in the left ventricle and aorta reaches a maximum of ca. 120 mmHg (systolic pressure). In the early phase of ejection (IIa or *rapid ejec*-

tion phase), a large portion of the stroke volume (SV) is rapidly expelled (\rightarrow **A4**) and the blood flow rate reaches a maximum (\rightarrow A5). Myocardial excitation subsequently decreases (T wave of the ECG, \rightarrow A1) and ventricular pressure decreases (the remaining SV fraction is slowly ejected, phase IIb) until it falls below that of the aorta or pulmonary artery, respectively. This leads to closing of the semilunar valves, producing the second heart sound $(\rightarrow A6)$. The mean SV at rest is about 80 mL or. more precisely, 47 mL/m² body surface area. The corresponding mean ejection fraction (SV) EDV) at rest is about 0.67. The end-systolic volume (ESV) remaining in the ventricles at this point is about $40 \, \text{mL} (\rightarrow \text{A4})$.

The first phase of ventricular diastole or **isovolumetric relaxation** now begins (phase III; ca. 60 ms). The atria have meanwhile refilled, mainly due to the suction effect created by the lowering of the valve plane during ejection. As a result, the central venous pressure (CVP) decreases (\rightarrow **A3**, falls from c to x). The ventricular pressure now drops rapidly, causing the atrioventricular valves to open again when it falls short of atrial pressure.

The **filling phase** now begins (phase IV; ca. 500 ms at rest). The blood passes rapidly from the atria into the ventricles, resulting in a drop in CVP (\rightarrow A3, point y). Since the ventricles are 80% full by the first quarter of diastole, this is referred to as *rapid ventricular filling* (phase IVa; \rightarrow A4). *Ventricular filling slows down* (phase IVb), and the *atrial systole* (phase IVc) and the awave of CVP follows (\rightarrow A2,3). At a normal heart rate, the atrial contraction contributes about 15% to ventricular filling. When the heart rate increases, the duration of the cardiac cycle decreases mainly at the expense of diastole, and the contribution of atrial contraction to ventricular filling increases.

The heart beats produce a **pulse wave** (pressure wave) that travels through the arteries at a specific *pulse wave velocity* (PWV): the PWV of the aorta is $3-5\,\text{m/s}$, and that of the radial artery is $5-12\,\text{m/s}$. PWV is much higher than the blood flow velocity (\dot{V}), which peaks at 1 m/s in the aorta and increases proportionally to (a) decreases in the compliance of aortic and arterial walls and (b) increases in blood pressure.



Cardiac Impulse Generation and Conduction

The heart contains muscle cells that generate (pacemaker system), conduct (conduction system) and respond to electrical impulses (working myocardium). Cardiac impulses are generated within the heart (automaticity). The frequency and regularity of pacemaking activity are also intrinsic to the heart (rhythmicity). Myocardial tissue comprises a functional (not truly anatomical) syncytium because the cells are connected by gap junctions (\rightarrow p. 16ff.). This also includes the atrioventricular junction (\rightarrow p. 195 A). Thus, an impulse arising in any part of the heart leads to complete contraction of both ventricles and atria or to none at all (allor-none response).

Cardiac contraction is normally stimulated by impulses from the **sinoatrial node** (SA node), which is therefore called the *primary pacemaker*. The impulses are conducted (\rightarrow **A**) through the atria to the *atrioventricular node* (**AV node**). The *bundle of His* is the beginning of the **specialized conduction system**, including also the left and right (Tawara's) *bundle branches* and the *Purkinje fibers*, which further transmit the impulses to the ventricular myocardium, where they travel from inside to outside and from apex to base of the heart. This electrical activity can be tracked in vivo (\rightarrow **C**) by electrocardiography (\rightarrow p. 196ff.).

Pacemaker potential (\rightarrow **B1**, top). The cell potential in the SA node is a *pacemaker potential*. These cells do not have a constant resting potential. Instead, they slowly depolarize immediately after each repolarization, the most negative value of which is the *maximum diastolic potential* (MDP, ca. -70 mV). The slow *diastolic depolarization* or *prepotential* (PP) prevails until the *threshold potential* (TP) has again been reached. Thus triggering another action potential (AP).

The pacemaker potential (\rightarrow **B1**, bottom) is subject to various underlying **changes in ion conductance** (g) and **ionic flow** (I) through the plasma membrane (\rightarrow p. 32 ff.). Starting at the MDP, a hyperpolarization-triggered increase in non-selective conductance and influx (I_{fi} = "funny") of cations into the cells lead to slow depolarization (PP). When the TP is reached, g_{Ca} increases quickly, and the slope of the

pacemaker potential rises. This upslope is caused by increased influx of Ca^{2+} (I_{Ca}). When the potential rises to the positive range, g_K increases sharply, resulting in the efflux of K^+ (I_K), and the pacemaker cells repolarize to the MDP.

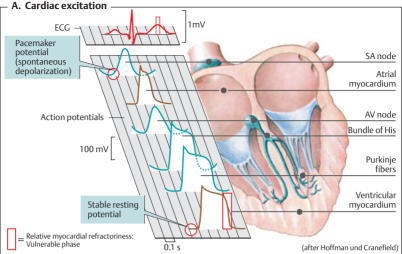
Each action potential in the SA node normally generates one heart beat. The **heart rate** is therefore determined by the rate of impulse generation by the pacemaker. The impulse generation frequency *decreases* (\rightarrow **B3**, broken curves) when (a) the PP slope decreases (\rightarrow **B3a**), (b) the TP becomes less negative (\rightarrow **B3b**), (c) the MDP becomes more negative, resulting in the start of spontaneous depolarization at lower levels (\rightarrow **B3c**), or (d) repolarization after an action potential occurs more slowly (slope flatter).

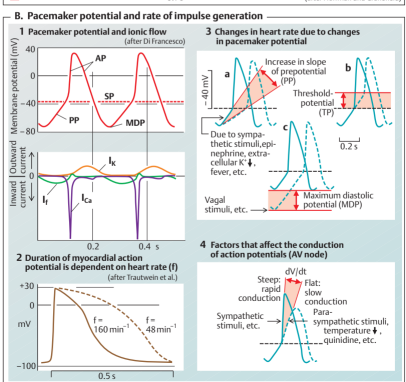
The first three conditions extend the time required to reach the threshold potential.

All components of the conduction system can depolarize spontaneously, but the SA node is the natural or nomotopic pacemaker in cardiac excitation (sinus rhythm normally has a rate of 60 to 100 min-1). The intrinsic rhythms of the other pacemakers are slower than the sinus rhythm ($\rightarrow C$, table) because the slope of their PPs and repolarizations are "flatter" (see above). APs arising in the SA node therefore arrive at subordinate ("lower") levels of the conduction system before spontaneous depolarization has reached the intrinsic TP there. The intrinsic rhythm of the lower components come into play (ectopic pacemakers) when (a) their own frequency is enhanced, (b) faster pacemakers are depressed, or (c) the conduction from the SA node is interrupted (\rightarrow p. 200). The heart beats at the AV rhythm (40 to 55 min⁻¹)/or even slower (25 to 40 min⁻¹) when controlled by tertiary (ventricular) pacemakers.

Overdrive suppression. The automaticity of lower pacemaker cells (e.g., AV node or Purkinje cells) is suppressed temporarily after they have been driven by a high frequency. This leads to increased Na⁺ influx and therefore to increased activity of the Na⁺-K⁺-ATPase. Because it is electrogenic (→ p. 28), the cells become hyperpolarized and it takes longer to reach threshold than without prior high-frequency overdrive (→ B3c).

The cells of the **working myocardium** contain voltage-gated *fast Na*⁺ *channels* that permit the brief but rapid influx of Na⁺ at the beginning of





an AP. The slope of their APs therefore rises more sharply than that of a pacemaker potential (\rightarrow **A**). A **resting potential** prevails between APs, i.e. spontaneous depolarization normally does not occur in the working myocardium. The long-lasting myocardial AP has a characteristic **plateau** (\rightarrow p. 59 A). Thus, the first-stimulated parts of the myocardium are still in a refractory state when the AP reaches the last-stimulated parts of the myocardium. This prevents the cyclic re-entry of APs in the myocardium. This holds true, regardless of whether the heart rate is very fast or very slow since the duration of an AP varies according to heart rate (\rightarrow **B2**).

Role of Ca2+. The incoming AP opens voltagegated Ca2+ channels (associated with dihydropyridine receptors) on the sarcolemma of myocardial cells, starting an influx of Ca2+ from the ECF (\rightarrow p. 63/B3). This produces a local increase in cytosolic Ca²⁺ (Ca²⁺ "spark") which, in turn, triggers the opening of ligand-gated, ryanodine-sensitive Ca2+ channels in the sarcoplasmic reticulum (Ca2+ store). The influx of Ca2+ into the cytosol results in electromechanical coupling (\rightarrow p. 62) and myocardial contraction. The cytosolic Ca²⁺ is also determined by active transport of Ca2+ ions back (a) into the Ca2+ stores via a Ca2+-ATPase, called SERCA, which is stimulated by phospholamban, and (b) to the ECF. This is achieved with the aid of a Ca²⁺-ATPase and a 3 Na⁺/Ca²⁺ exchange carrier that is driven by the electrochemical Na+ gradient established by Na+-K+-ATPase.

Although the heart beats autonomously, efferent cardiac nerves are mainly responsible for modulating heart action according to changing needs. The autonomic nervous system (and epinephrine in plasma) can alter the following aspects of heart action: (a) rate of impulse generation by the pacemaker and, thus the heart rate (chronotropism); (b) velocity of impulse conduction, especially in the AV node (dromotropism); and (c) contractility of the heart, i.e., the force of cardiac muscle contraction at a given initial fiber length (inotropism).

These changes in heart action are induced by **acetylcholine** (ACh; \rightarrow p.82) released by parasympathetic fibers of the vagus nerve (binds with M_2 cholinoceptors on pacemaker cells), by **norepinephrine** (NE) released by sympathetic nerve fibers, and by plasma **epi**-

nephrine (E). NE and E bind with β_1 -adrenoceptors (\rightarrow p. 84ff.). The firing frequency of the SA node is increased by NE and E (positive chronotropism) and decreased by ACh (negative chronotropism) because these substances alter the slopes of the PP and the MDP in the SA cells (\rightarrow **B3a** and **c**). Under the influence of ACh, the slope of the PP becomes flatter and the MDP becomes more negative gk rises. In versely, slope and amplitude of PP rises under the influence of E or sympathetic stimuli (higher I_f) due to a rise in cation (Na⁺) conductance and, under certain conditions, a decrease in the g_K Only NE and E have chronotropic effects in the lesser components of the impulse conduction system. This is decisive when the AV node or tertiary pacemakers take over.

ACh (left branch of vagus nerve) decreases the velocity of **impulse conduction in the AV node**, whereas NE and E increase it due to their negative and positive dromotropic effects, respectively. This is mainly achieved through changes in the amplitude and slope of the upstroke of the AP (\rightarrow **B3c** and **B4**), g_K and g_{Ca} .

In positive inotropism, NE and E have a direct effect on the working myocardium. The resulting increase in contractility is based on an increased influx of Ca2+ ions from the ECF triggered by β_1 -adrenoceptors, resulting in an increased cytosolic Ca2+. This Ca2+ influx can be inhibited by administering Ca2+ channel blockers (Ca2+ antagonists). Other factors that increase cardiac contractility are an increase in AP duration, resulting in a longer duration of Ca2+ influx, and inhibition of Na+-K+-ATPase (e.g., by the cardiac glycosides digitalis and strophanthin). The consequences are: flatter Na⁺ gradient across the cell membrane ⇒ decreased driving force for 3 Na+/Ca2+ exchange carriers ⇒ decreased Ca2+ efflux ⇒ increased cytosolic Ca2+ conc.

When the heart rate is low, Ca^{2+} influx over time is also low (fewer APs per unit time), allowing plenty of time for the efflux of Ca^{2+} between APs. The mean cytosolic Ca^{2+} conc. is therefore reduced and contractility is low. Only by this indirect mechanism are parasympathetic neurons able to elicit a negative inotropic effect (*frequency inotropism*). NE and E can exert their positive inotropic effects either indirectly by increasing the high heart or directly via β_1 -adrenoceptors of the working

Electrocardiogram (ECG)

The ECG records **potential differences** (few **m/V**) caused by cardiac excitation. This provides information on heart position, relative chamber size, heart rhythm, impulse origin/propagation and rhythm/conduction disturbances, extent and location of myocardial ischemia, changes in electrolyte concentrations, and drug effects on the heart. However, it does *not* provide data on cardiac contraction or pumping function.

ECG potential differences arise at the interface between stimulated and non-stimulated myocardium. Totally stimulated or unstimulated myocardial tissue does not generate any visible potentials The migration of the excitatory front through the heart muscle gives rise to numerous potentials that vary in magnitude and direction.

These vectors can be depicted as arrows, where the length of the arrow represents the magnitude of the potential and the direction of the arrow indicates the direction of the potential (arrowhead is +). As in a force parallelogram, the **integral vector** (summation vector) is the sum of the numerous individual vectors at that moment (\rightarrow **A**, red arrow).

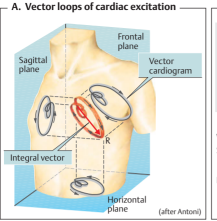
The magnitude and direction of the integral vector change during the cardiac cycle, producing the typical *vector loop* seen on a **vector-cardiogram**. (In **A**, the maximum or chief vector is depicted by the arrow, called the "electrical axis" of the heart, see below).

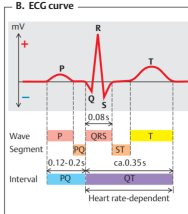
Limb and chest leads of the ECG make it possible to visualize the course of the integral vector over time, projected at the plane determined by the leads (scalar ECG). Leads parallel to the integral vector show full deflection (R wave $\approx 1-2$ mV), while those perpendicular to it show no deflection. Einthoven leads I. II. and III are bipolar limb leads positioned in the frontal plane. Lead I records potentials between the left and right arm, lead II those between the right arm and left leg, and lead III those between the left arm and left $leg \rightarrow C1$. Goldberger leads are unipolar augmented limb leads in the frontal plane. One lead (right arm, aVR, left arm aVL, or left leg, aVF; \rightarrow D2) acts as the different electrode, while the other two limbs are connected and serve as the indifferent (reference) electrode (\rightarrow D1). Wilson leads (V_1-V_6) are unipolar chest leads positioned on

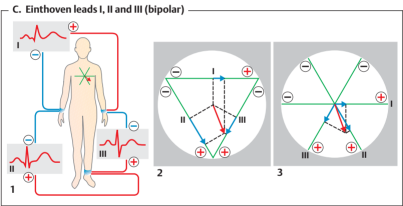
the left side of the thorax in a nearly horizontal plane $(\rightarrow F)$. When used in combination with the aforementioned leads in the frontal plane, they provide a three-dimensional view of the integral vector. To make recordings with the chest leads (different electrode), the three limb leads are connected to form an indifferent electrode with high resistances (5 k Ω). The chest leads mainly detect potential vectors directed towards the back. These vectors are hardly detectable in the frontal plane. Since the mean QRS vector (see below) is usually directed downwards and towards the left back region, the QRS vectors recorded by leads V₁-V₃ are usually negative, while those detected by V5 and V_6 are positive.

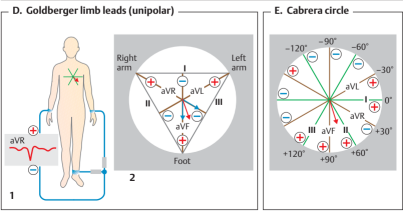
Intraesophageal leads and additional leads positioned in the region of the right chest $(V_{r3}-V_{r6})$ and left back (V_7-V_9) are useful in certain cases $(\rightarrow$ **F2**).

An ECG depicts electrical activity as waves, segments, and intervals (\rightarrow **B** and p. 195 C). By convention, upward deflection of the waves is defined as positive (+), and downward deflection as negative (-). The electrical activity associated with atrial depolarization is defined as the **Pwave** ($< 0.3 \, \text{mV}$, $< 0.1 \, \text{s}$). Repolarization of the atria normally cannot be visualized on the ECG since it tends to be masked by the QRS complex. The QRS complex (< 0.1 s) consists of one, two or three components: Q wave (mV < 1/4 of R, < 0.04 s), **R wave** and/or **S wave** $(R+S > 0.6 \,\mathrm{mV})$. The potential of the mean ORS vector is the sum of the amplitudes of the Q, R and S waves (taking their positive and negative polarities into account). The voltage of the mean QRS vector is higher (in most leads) than that of the P wave because the muscle mass of the ventricles is much larger than that of the atria. The R wave is defined as the first positive deflection of the ORS complex, which means that R waves from different leads may not be synchronous. The QRS complex represents the depolarization of the ventricles, and the T wave represents their repolarization. Although opposing processes, the T wave usually points in the same direction as the R wave (+ in most leads). This means that depolarization and repolarization do not travel in the same direction $(\rightarrow p. 195 C, QRS \text{ and } T$: vector arrows point in the same direction despite reversed polarity during repolarization). The PO (or PR) segment









(complete atrial excitation) and the **ST segment** (complete ventricular excitation) lie approx. on the **isoelectric line** (0 mV). The **PQ** (or PR) **interval** (< 0.2 s) is measured from the beginning of the P wave to the beginning of the Q wave (or to the R wave if Q wave is absent) and corresponds to the time required for *atrioventricular conduction* (\rightarrow **B**). The **QT interval** is measured from the start of the Q wave to the end of the T wave. It represents the overall time required for depolarization and repolarization of the ventricles and is dependent on the heart rate (0.35 to 0.40 s at a heart rate of 75 min⁻¹).

Figure **E** illustrates the six frontal leads (Einthoven and Goldberger leads) on the *Cabrera circle*. Synchronous measurement of the amplitude of Q, R and S from two or more leads can be used to determine any integral vector in the frontal plane (\rightarrow **G**). The direction of the largest mean QRS vector is called the **QRS axis** (\rightarrow **C3** and **G**, red arrows). If the excitation spreads normally, the QRS axis roughly corresponds to the anatomic longitudinal axis of the heart.

The mean QRS axis ("electrical axis") of the heart, which normally lies between +90 degrees to -30 degrees in adults (\rightarrow **G**, **H**). Right type ($\alpha = +120^{\circ}$ to $+90^{\circ}$) is not unusual in children, but is often a sign of abnormality in adults. Mean QRS axes ranging from +90 degrees to +60 degrees are described as the vertical type (\rightarrow G1), and those ranging from +60 degrees to +30 degrees are classified as the intermediate type (\rightarrow **G2**). Left type occurs when $\alpha = +30$ degrees to -30 degrees (\rightarrow **G3**). Abnormal deviation: Right axis deviation (>+120°) can develop due right ventricular hypertrophy, while left axis deviation (more negative than -30°) can occur due to left ventricular hypertrophy.

An extensive **myocardial infarction (MI)** can shift the electrical axis of the heart. Marked Q wave abnormality (\rightarrow 11) is typical in transmural myocardial infarction (involving entire thickness of ventricular wall): Q wave duration >0.04 s and Q wave amplitude >25% of total amplitude of the QRS complex. These changes appear within 24 hours of MI and are caused by failure of the dead myocardium to conduct electrical impulses. Preponderance of the excitatory vector in the healthy contralateral side of the heart therefore occurs while the affected part of the

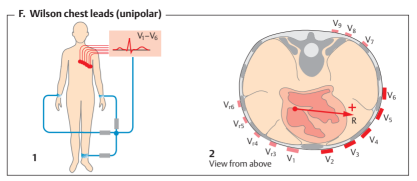
myocardium should be depolarizing (first 0.04s of QRS). The so-called "0.04-sec vector" is therefore said to point away from the infarction. Anterior MI is detected as highly negative O waves (with smaller R waves) mainly in leads V5, V6, I and aVL. Q wave abnormalities can persist for years after MI (\rightarrow 12/3), so they may not necessarily be indicative of an acute infarction. ST elevation points to ischemic but not (yet) necrotic parts of the myocardium. This can be observed: (1) in myocardial ischemia (angina pectoris), (2) in the initial phase of transmural MI. (3) in nontransmural MI, and (4) along the margins of a transmural MI that occurred a few hours to a few days prior $(\rightarrow 14)$. The ST seament normalizes within 1 to 2 days of MI, but the Twave remains inverted for a couple of weeks (\rightarrow 15 and 2).

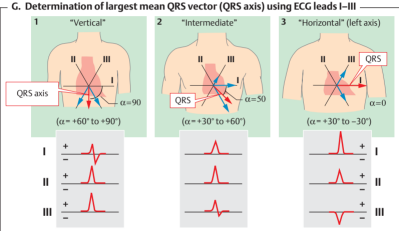
Excitation in Electrolyte Disturbances

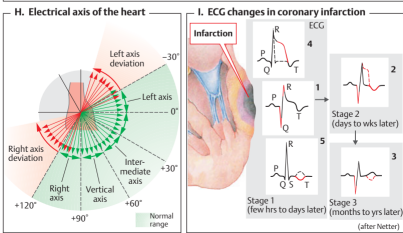
Hyperkalemia. Mild hyperkalemia causes various changes, like elevation of the MDP $(\rightarrow p. 192)$ in the SA node. It can sometimes have positive chronotropic effects (\rightarrow p. 193 B3c). In severe hyperkalemia, the more positive MDP leads to the inactivation of Na+ channels $(\rightarrow p.46)$ and to a reduction in the slope and amplitude of APs in the AV node (negative dromotropic effect; \rightarrow p. 193 B4). Moreover, the K⁺ conductance (gk) rises, and the PP slope becomes flatter due to a negative chronotropic effect (→ p. 193 B3a). Faster myocardial repolarization decreases the cytosolic Ca2+ conc. In extreme cases, the pacemaker is also brought to a standstill (cardiac paralysis). Hypokalemia (moderate) has chronotropic and inotropic effects (→ p. 193 B3a), whereas hypercalcemia is thought to raise the g_K and thereby shortens the duration of the myocardial AP.

ECG. Changes in serum K⁺ and Ca²⁺ induce characteristic changes in myocardial excitation.

- Hyperkalemia (> 6.5 mmol/L): tall, peaked T waves and conduction disturbances associated with an increased PQ interval and a widened QRS. Cardiac arrest can occur in extreme cases.
- Hypokalemia (< 2.5 mmol/L): ST depression, biphasic T wave (first positive, then negative) followed by a positive U wave.
- Hypercalcemia (> 2.75 mmol/L total calcium): shortened QT interval due to a shortened ST segment.
- ◆ *Hypocalcemia* (< 2.25 mmol/L total calcium): prolonged QT interval.







Cardiac Arrhythmias

Arrhythmias are pathological changes in cardiac *impulse generation or conduction* that can be visualized by ECG. **Disturbances of impulse generation** change the sinus rhythm. *Sinus tachycardia* (→ **A2**): The sinus rhythm rises to 100 min⁻¹ or higher e.g., due to physical exertion, anxiety, fever (rise of about 10 beats/min for each 1 °C) or hyperthyroidism. *Sinus bradycardia*: The heart rate falls below 60 min⁻¹ (e.g., due to hypothyroidism). In both cases the rhythm is regular whereas in *sinus arrhythmias* the rate varies. In adolescents, sinus arrhythmias can be physiological and respiration-dependent (heart rate increases during inspiration and decreases during expiration).

Ectopic pacemakers. Foci in the atrium, AV node or ventricles can initiate abnormal ectopic (heterotopic) impulses, even when normal (nomotopic) stimulus generation by the SA node is taking place $(\rightarrow A)$. The rapid discharge of impulses from an atrial focus can induce atrial tachycardia (serrated baseline instead of normal P waves), which triggers a ventricular response rate of up to 200 min⁻¹. Fortunately, only every second or third stimulus is transmitted to the ventricles because part of the impulses arrive at the Purkinje fibers (longest APs) during their refractory period. Thus, Purkinje fibers act as impulse frequency filters. Elevated atrial contraction rates of up to 350 min-1 are defined as atrial flutter, and all higher rates are defined as atrial fibrillation (up to 500 min⁻¹). Ventricular stimulation is then totally irregular (absolute arrhythmia). Ventricular tachycardia is a rapid train of impulses originating from a ventricular (ectopic) focus, starting with an extrasystole (ES) (\rightarrow B3; second ES). The heart therefore fails to fill adequately, and the stroke volume decreases. This can lead to ventricular fibrillation (extremely frequent and uncoordinated contractions; \rightarrow **B4**). Because of failure of the ventricle to transport blood, ventricular fibrillation can be fatal.

Ventricular fibrillation mainly occurs when an ectopic focus fires during the **relative re-fractory period** of the previous AP (called the "vulnerable phase" synchronous with T wave on the ECG; \rightarrow p. 193 A). The APs triggered during this period have *smaller slopes*, *lower*

propagation velocities, and shorter durations. This leads to re-excitation of myocardial areas that have already been stimulated (*re-entry cycles*). Ventricular fibrillation can be caused by electrical accidents and can usually be corrected by timely electrical *defibrillation*.

Extrasystoles (ES). The spread of impulses arising from an supraventricular (atrial or nodal) ectopic focus to the ventricles can disturb their sinus rhythm. leading to a supraventricular arrhythmia. When atrial extrasystoles occur, the Pwave on the ECG is distorted while the QRS complex remains normal. Nodal extrasystoles lead to retrograde stimulation of the atria, which is why the P wave is negative and is either masked by the QRS complex or appears shortly thereafter (\rightarrow **B1** right). Since the SA node often is discharged by a supraventricular extrasystole, the interval between the R wave of the extrasvstole (R_{ES}) and the next normal R wave increases by the amount of time needed for the stimulus to travel from the focus to the SA node. This is called the postextrasystole pause. The RR intervals are as follows: R_{FS}R > RR and (RR_{FS} + R_{FS}R) < 2 RR (\rightarrow **B1**).

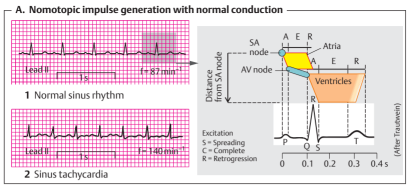
Ventricular (or infranodal) **ES** (\rightarrow **B2**, **B3**) distorts the QRS complex of the ES. If the sinus rate is slow enough, the ES will cause a ventricular contraction between two normal heart beats; this is called an *interpolated* (or interposed) ES (\rightarrow **B2**). If the sinus rate is high, the next sinus stimulus reaches the ventricles while they are still refractory from the ectopic excitation. Ventricular contraction is therefore blocked until the next sinus stimulus arrives, resulting in a *compensatory pause*, where RR_{ES} + R_{ES}R = 2 RR.

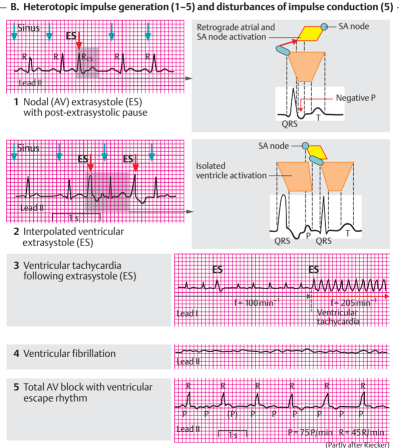
Disturbances of impulse conduction: AV block. First-degree AV block: prolonged but otherwise normal impulse conduction in the AV node (PQ interval > 0.2 sec); second-degree AV block: only every second (2:1 block) or third (3:1 block) impulse is conducted. Third-degree AV block: no impulses are conducted; sudden cardiac arrest may occur (Adam-Stokes attack or syncope). Ventricular atopic pacemakers then take over (ventricular bradycardia with normal atrial excitation rate), resulting in partial or total disjunction of QRS complexes and P waves (\rightarrow **B5**). The heart rate drops to 40 to 55 min-1 when the AV node acts as the pacemaker (\rightarrow **B5**), and to a mere 25 to 40 min⁻¹ when tertiary (ventricular) pacemakers take over. Artificial pacemakers are then used.

Bundle branch block: disturbance of conduction in a branch of the bundle of His. Severe QRS changes occur because the affected side of the myocardium is activated by the healthy side via abnormal pathways.

200

201





Ventricular Pressure-Volume Relationships

The relationship between the volume (length) and pressure (tension) of a ventricle illustrates the interdependence between muscle length and force in the specific case of the heart (\rightarrow p. 66ff.). The **work diagram** of the heart can be constructed by plotting the changes in *ventricular pressure over volume* during one complete cardiac cycle (\rightarrow A1, points A-D-S-V-A, pressure values are those for the left ventricle).

The following *pressure–volume curves* can be used to construct a **work diagram of the ventricles**:

- Passive (or resting) pressure-volume curve: Indicates the pressures that result passively (without muscle contraction) at various ventricular volume loads (→ A1, 2; blue curve).
- ◆ Isovolumic peak curve (→ A1, 2, green curves): Based on experimental measurements made using an isolated heart. Data are generated for various volume loads by measuring the peak ventricular pressure at a constant ventricular volume during contraction. The contraction is therefore isovolumetric (isovolumic), i.e., ejection does not take place (→ A2, vertical arrows).
- ◆ Isotonic (or isobaric) peak curve (→ A1, 2, violet curves). Also based on experimental measurements taken at various volume loads under isotonic (isobaric) conditions, i.e., the ejection is controlled in such a way that the ventricular pressure remains constant while the volume decreases (→ A2, horizontal arrows).
- Afterloaded peak curve: (A1, 2, orange curves). Systole (→ p. 190) consists of an isovolumic contraction phase (→ A1, A−D and p. 191A, phase I) followed by an auxotonic ejection phase (volume decreases while pressure continues to rise) (→ A1, D−S and p. 191 A, phase II). This type of mixed contraction is called an afterloaded contraction (see also p. 67 B). At a given volume load (preload) (→ A1, point A), the afterloaded peak value changes (→ A1, point S) depending on the aortic end-diastolic pressure (→ A1, point D). All the afterloaded peak values are represented on the curve, which appears as a (nearly) straight line connecting the isovolumic and isotonic peaks for each respective volume load (point A) (→ A1, points T and M).

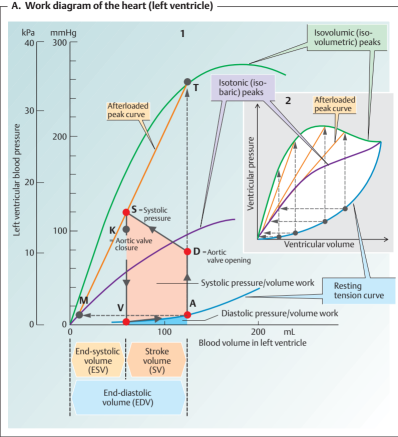
Ventricular work diagram. The pressure-volume relationships observed during the cardiac cycle (\rightarrow p. 190) can be plotted as a work diagram, e.g., for the left ventricle (\rightarrow A1): The *end-diastolic volume* (**EDV**) is 125 mL (\rightarrow A1, point A). During the *isovolumetric contraction*

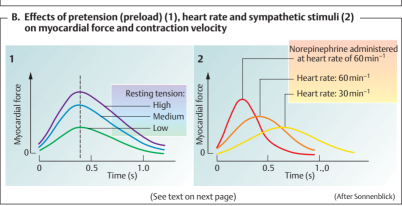
phase, the pressure in the left ventricle rises (all valves closed) until the diastolic aortic pressure (80 mmHg in this case) is reached $(\rightarrow A1$, point D). The aortic valve then opens. During the ejection phase, the ventricular volume is reduced by the stroke volume (SV) while the pressure initially continues to rise $(\rightarrow p. 188, Laplace's law, Eq. 8.4b: P_{tm} \uparrow be$ cause $r \downarrow and w \uparrow$). Once maximum (systolic) pressure is reached (\rightarrow A1, point S), the volume will remain virtually constant, but the pressure will drop slightly until it falls below the aortic pressure, causing the aortic valve to close (\rightarrow A1, point K). During the isovolumetric relaxation phase, the pressure decreases to (almost) $0 (\rightarrow A1$, point V). The ventricles now contain only the end-systolic volume (ESV), which equals 60 mL in the illustrated example. The ventricular pressure rises slightly during the filling phase (passive pressure-volume curve).

Cardiac Work and Cardiac Power

Since work ($I = N \cdot m$) equals pressure ($N \cdot m^{-2}$ = Pa) times volume (m3), the area within the working diagram (\rightarrow A1, pink area) represents the pressure/volume (P/V) work achieved by the left ventricle during systole (13,333 Pa- $0.00008 \text{ m}^3 = 1.07 \text{ I}$; right ventricle: 0.16 I). In systole, the bulk of cardiac work is achieved by active contraction of the myocardium, while a much smaller portion is attributable to passive elastic recoil of the ventricle, which stretches while filling. This represents diastolic filling work (\rightarrow A1, blue area under the blue curve), which is shared by the ventricular myocardium (indirectly), the atrial myocardium, and the respiratory and skeletal muscles $(\rightarrow p. 204, venous return).$

Total cardiac work. In addition to the cardiac work performed by the left and right ventricles in systole (ca. 1.2 J at rest), the heart has to generate 20% more energy (0.24 J) for the pulse wave (\rightarrow p. 188, windkessel). Only a small amount of energy is required to accelerate the blood at rest (1% of total cardiac work), but the energy requirement rises with the heart rate. The total **cardiac power** (=work/time, \rightarrow p. 374) at rest (70 min⁻¹ = 1.17 s⁻¹) is approximately 1.45 [\cdot 1.17 s⁻¹ = 1.7 W.





Frank–Starling mechanism (FSM): The heart *autonomously* responds to changes in ventricular volume load or aortic pressure load by adjusting the stroke volume (SV) in accordance with the myocardial *preload* (resting tension; → p. 66 ff.). The FSM also functions to maintain an equal SV in both ventricles to prevent congestion in the pulmonary or systemic circulation.

Preload change. When the volume load (preload) *increases*, the start of isovolumic contraction shifts to the right along the passive P–V curve (\rightarrow **A1**, from point A to point A₁). This increases end-diastolic volume (EDV), stroke volume (SV), cardiac work and end-systolic volume (ESV) (\rightarrow **A**).

Afterload change. When the aortic pressure load (afterload) *increases*, the aortic valve will not open until the pressure in the left ventricle has risen accordingly (\rightarrow **A2**, point D_t). Thus, the SV in the short transitional phase (SV_t) will decrease, and ESV will rise (ESV_t). Consequently, the start of the isovolumic contraction shifts to the right along the passive P–V curve (\rightarrow **A2**, point A₂). SV will then normalize (SV₂) despite the increased aortic pressure (D₂), resulting in a relatively large increase in ESV (ESV₂).

Preload or afterload-independent changes in myocardial contraction force are referred to as contractility or inotropism. It increases in response to norepinephrine (NE) and epinephrine (E) as well as to increases in heart rate (β_1 -adrenoceptor-mediated, positive inotropic effect and frequency inotropism, respectively; \rightarrow p. 194). This causes a number of effects, particularly, an increase in isovolumic pressure peaks (\rightarrow A3, green curves). The heart can therefore pump against increased pressure levels (\rightarrow A3, point D₃) and/or eject larger SVs (at the expense of the ESV) (\rightarrow A3, SV₄).

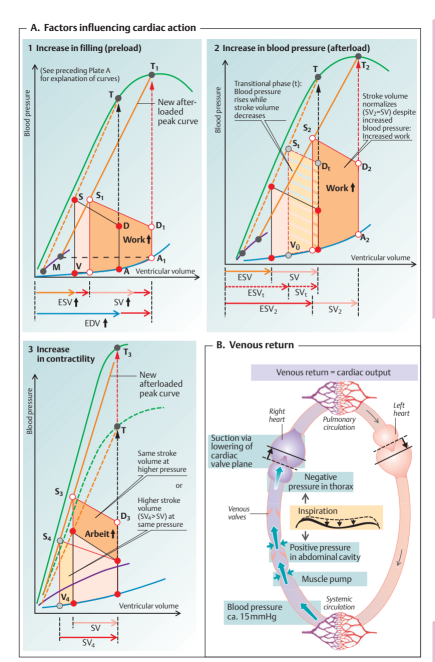
While changes in the preload only affect the **force** of contraction (\rightarrow p. 203 B1), changes in contractility also affect the **velocity** of contraction (\rightarrow p. 203/B2). The steepest increase in isovolumic pressure per unit time (*maximum dP/dt*) is therefore used as a measure of contractility in clinical practice. dP/dt is increased E and NE and decreased by bradycardia (\rightarrow p. 203 B2) or heart failure.

Venous Return

Blood from the capillaries is collected in the veins and returned to the heart. The **driving forces** for this venous return (\rightarrow **B**) are: (a) *vis a tergo*, i.e., the postcapillary blood pressure (BP) (ca. 15 mmHg); (b) the suction that arises due to lowering of the cardiac valve plane in systole; (c) the pressure exerted on the veins during skeletal muscle contraction (*muscle pump*); the valves of veins prevent the blood from flowing in the wrong direction, (d) the increased abdominal pressure together with the lowered intrathoracic pressure during inspiration (P_{pl} ; \rightarrow p. 108), which leads to thoracic venous dilatation and suction (\rightarrow p. 206).

Orthostatic reflex. When rising from a supine to a standing position (orthostatic change), the blood vessels in the legs are subjected to additional hydrostatic pressure from the blood column. The resulting vasodilation raises blood volume in the leg veins (by ca. 0.4 L). Since this blood is taken from the central blood volume, i.e., mainly from pulmonary vessels, venous return to the left atrium decreases, resulting in a decrease in stroke volume and cardiac output. A reflexive increase (orthostatic reflex) in heart rate and peripheral resistance therefore occurs to prevent an excessive drop in arterial BP (\rightarrow pp. 7 E and 212 ff.); orthostatic collapse can occur. The drop in central blood volume is more pronounced when standing than when walking due to muscle pump activity. Conversely, pressure in veins above the heart level, e.g., in the cerebral veins, decreases when a person stands still for prolonged periods of time. Since the venous pressure just below the diaphragm remains constant despite changes in body position, it is referred to as a hydrostatic indifference point.

The **central venous pressure** (**CVP**) is measured at the right atrium (normal range: $0-12 \text{ cm H}_2\text{O}$ or 0-9 mmHg). Since it is mainly dependent on the blood volume, the CVP is used to monitor the blood volume in clinical medicine (e.g., during a transfusion). Elevated CVP (> $20 \text{ cm H}_2\text{O}$ or 15 mmHg) may be pathological (e.g., due to heart failure or other disease associated with cardiac pump dysfunction), or physiological (e.g., in pregnancy).



Arterial Blood Pressure

The term blood pressure (BP) per se refers to the arterial BP in the systemic circulation. The maximum BP occurs in the aorta during the systolic ejection phase; this is the systolic pressure (Ps); the minimum aortic pressure is reached during the isovolumic contraction phase (while the aortic valves are closed) and is referred to as the diastolic pressure (Pd) $(\rightarrow A1$ and p. 191, phase I in A2). The systolicdiastolic pressure difference (Ps-Pd) represents the blood pressure amplitude, also called pulse pressure (PP), and is a function of the stroke volume (SV) and arterial compliance (C = dV/dP, \rightarrow p. 188). When C decreases at a constant SV, the systolic pressure P_s will rise more sharply than the diastolic pressure Pd. i.e., the PP will increase (common in the elderly; described below). The same holds true when the SV increases at a constant C.

If the **total peripheral resistance** (TPR, \rightarrow p. 188) **increases** while the SV ejection time remains constant, then P_s and the P_d will increase by the same amount (no change in PP). However, increases in the TPR normally lead to retardation of SV ejection and a decrease in the ratio of arterial volume rise to peripheral drainage during the ejection phase. Consequently, P_s rises less sharply than P_d and PP decreases.

Normal range. In individuals up to 45 years of age, P_d normally range from 60 to 90 mmHg and P_s from 100 to 140 mmHg at rest (while sitting or reclining). A P_s of up to 150 mmHg is considered to be normal in 45 to 60-year-old adults, and a P_s of up to 160 mmHg is normal in individuals over 60 (\rightarrow C). Optimal BP regulation (\rightarrow p.212) is essential for proper tissue perfusion.

Abnormally low BP (**hypotension**) can lead to *shock* (\rightarrow p. 218), *anoxia* (\rightarrow p. 130) and tissue destruction. Chronically elevated BP (**hypertension**; \rightarrow p. 216) also causes damage because important vessels (especially those of the heart, brain, kidneys and retina) are injured.

The **mean BP** (= the average measured over time) is the decisive factor of peripheral perfusion (\rightarrow p. 188).

The mean BP can be determined by continuous BP measurement using an arterial catheter, etc. $(\rightarrow A)$.

By attenuating the pressure signal, only the mean BP is recorded.

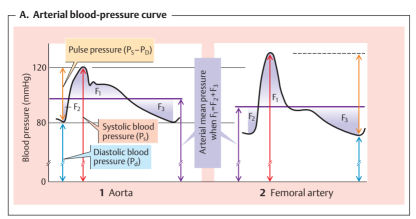
Although the mean BP falls slightly as the blood travels from the aorta to the arteries, the P_s in the greater arteries (e.g., femoral artery) is usually higher than in the aorta (A1 v. A2) because their compliance is lower than that of the aorta (see pulse wave velocity, p. 190).

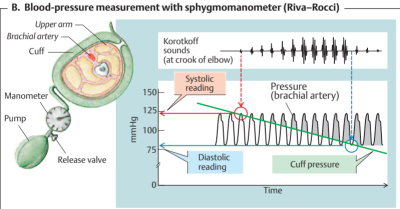
Direct invasive BP measurements show that the BP curve in arteries distal to the heart is not synchronous with that of the aorta due to the time delay required for passage of the pulse wave $(3-10 \text{ m/s}; \rightarrow \text{p. }190)$; its shape is also different $(\rightarrow \text{A1/A2})$.

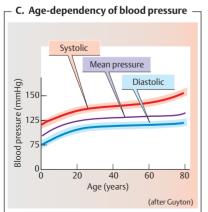
The BP is routinely measured externally (at the level of the heart) according to the Riva-Rocci method by sphygmomanometer (\rightarrow B). An inflatable cuff is snugly wrapped around the arm and a stethoscope is placed over the brachial artery at the crook of the elbow. While reading the manometer, the cuff is inflated to a pressure higher than the expected P_s (the radial pulse disappears). The air in the cuff is then slowly released (2-4 mmHg/s). The first sounds synchronous with the pulse (Korotkoff sounds) indicate that the cuff pressure has fallen below the P_s . This value is read from the manometer. These sounds first become increasingly louder, then more quiet and muffled and eventually disappear when the cuff pressure falls below the P_d (second reading).

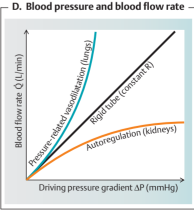
Reasons for false BP readings. When re-measuring the blood pressure, the cuff pressure must be completely released for 1 to 2 min. Otherwise venous pooling can mimic elevated $P_{\rm d}$. The cuff of the sphygmomanometer should be 20% broader than the diameter of the patient's upper arm. Falsely high $P_{\rm d}$ readings can also occur if the cuff is too loose or too small relative to the arm diameter (e.g., in obese or very muscular patients) or if measurement has to be made at the thigh.

The **blood pressure in the pulmonary artery** is much lower than the aortic pressure (\rightarrow p. 186). The pulmonary vessels have relatively thin walls and their environment (air-filled lung tissue) is highly compliant. Increased cardiac output from the right ventricle therefore leads to expansion and thus to decreased resistance of the pulmonary vessels (\rightarrow **D**). This prevents excessive rises in pulmonary artery pressure during physical exertion when cardiac output rises. The pulmonary vessels also function to buffer short-term fluctuations in blood volume (\rightarrow p. 204).









207

Endothelial Exchange Processes

Nutrients and waste products are exchanged across the walls of the capillaries and postpapillary venules (exchange vessels; \rightarrow p. 188). Their endothelia contain small (ca. 2–5 nm) or large (20–80 nm, especially in the kidneys and liver) functional pores: permeable, intercellular fissures or endothelial fenestrae, respectively. The degree of endothelial permeability varies greatly from one organ to another. Virtually all endothelia allow water and inorganic ions to pass, but most are largely impermeable to blood cells and large protein molecules. Transcytosis and carriers (\rightarrow p. 26f.) allow for passage of certain larger molecules.

Filtration and reabsorption. About 20 L/day of fluid is filtered (excluding the kidneys) into the interstitium from the body's exchange vessels. About 18 L/day of this fluid is thought to be reabsorbed by the venous limb of these vessels. The remaining 2 L/day or so make up the lymph flow and thereby return to the bloodstream $(\rightarrow A)$. The filtration or reabsorption rate Q_f is a factor of the endothelial filtration coefficient Kf (= water permeability k · exchange area A) and the effective filtration pressure $P_{eff}(Q_f = K_f \cdot P_{eff})$. Peff is calculated as the hydrostatic pressure difference ΔP minus the oncotic pressure difference $\Delta\pi$ across the capillary wall (Star*ling's relationship*; \rightarrow **A**), where Δ P = capillary pressure (P_{cap}) minus interstitial pressure (P_{int}, normally ≈ 0 mmHg). At the level of the heart, ΔP at the arterial end of the systemic capillaries is about 30 mmHg and decreases to about 22 mmHg at the venous end. Since $\Delta\pi$ (ca. 24 mmHg; \rightarrow **A**) counteracts Δ P, the initially high filtration rate ($P_{eff} = +6 \text{ mmHg}$) is thought to change into reabsorption whenever Peff becomes negative. (Since ΔP is only 10 mmHg in the lungs, the pulmonary P_{eff} is very low). $\Delta \pi$ occurs because the concentration of proteins (especially albumin) in the plasma is much higher than their interstitial concentration. The closer the reflection coefficient of the plasma proteins (σ_{prot}) to 1.0, the higher $\Delta \pi$ and, consequently, the lower the permeability of the membrane to these proteins (\rightarrow p. 377).

According to Starling's relationship, water reabsorption should occur as long as $P_{\rm eff}$ is negative. However, recent data suggest that a negative $P_{\rm eff}$ re-

sults in only transient reabsorption. After several minutes it stops because the interstitial oncotic pressure rises due to "self-regulation". Thus, a major part of the 18 L/d expected to be reabsorbed from the exchange vessels (see above) might actually be reabsorbed in the lymph nodes. Rhythmic contraction of the arterioles (**vasomotion**) may also play a role by decreasing P_{eff} and thus by allowing intermittent capillary reabsorption.

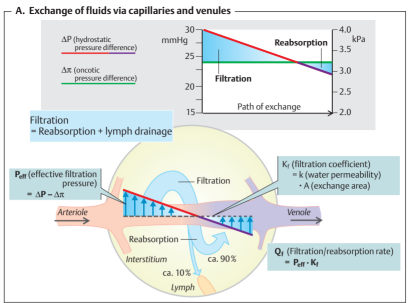
In parts of the body below the heart, the effects of **hydrostatic pressure** from the blood column increase the pressure in the capillary lumen (in the feet ≈ 90 mmHg). The filtration rate in these regions therefore rise, especially when standing still. This is counteracted by two "self-regulatory" mechanisms: (1) the outflow of water results in an increase in the luminal protein concentration (and thus $\Delta\pi$) along the capillaries (normally the case in glomerular capillaries, $\rightarrow p$. 152); (2) increased filtration results in an increase in $P_{\rm int}$ and a consequent decrease in ΔP .

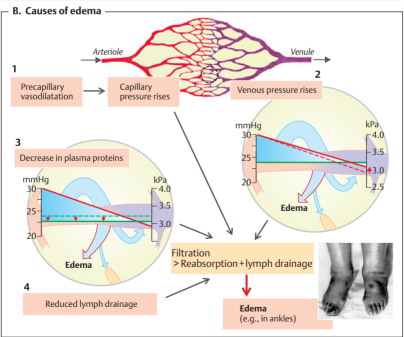
Edema. Fluid will accumulate in the interstitial space (extracellular edema), portal venous system (ascites), and pulmonary interstice (pulmonary edema) if the volume of filtered fluid is higher than the amount returned to the blood.

Causes of edema $(\rightarrow B)$:

- Increased capillary pressure (\rightarrow **B1**) due to precapillary vasodilatation ($P_{cap} \uparrow$), especially when the capillary permeability to proteins also increases ($\sigma_{prot} \downarrow$ and $\Delta \pi \downarrow$) due, for example, to infection or anaphylaxis (histamine etc.). Hypertension in the portal vein leads to ascites.
- ♦ Increased venous pressure ($P_{cap} \uparrow$, → **B2**) due, for example, to venous thrombosis or cardiac insufficiency (cardiac edema).
- Decreased concentration of plasma proteins, especially albumin, leading to a drop in $\Delta\pi$ (\rightarrow **B3** and p. 379 A) due, for example, to loss of proteins (proteinuria), decreased hepatic protein synthesis (e.g., in liver cirrhosis), or to increased breakdown of plasma proteins to meet energy requirements (hunger edema).
- Decreased lymph drainage due, e.g., to lymph tract compression (tumors), severance (surgery), obliteration (radiation therapy) or obstruction (bilharziosis) can lead to localized edema (→ B4).
- ♦ Increased hydrostatic pressure promotes edema formation in lower regions of the body (e.g., in the ankles; \rightarrow **B**).

Diffusion. Although dissolved particles are dragged through capillary walls along with filtered and reabsorbed water (solvent drag; \rightarrow p. 24), diffusion plays a much greater role in the exchange of solutes. *Net diffusion* of a substance (e.g., O_2 , CO_2) occurs if its plasma and interstitial conc. are different.





Myocardial Oxygen Supply

Coronary arteries. The blood flow to the myocardium is supplied by the two coronary arteries that arise from the aortic root. The right coronary artery (approx. 1/7th of the blood) usually supplies the greater portion of the right ventricle, while the left coronary artery (6/7th of the blood) supplies the left ventricle (\rightarrow A). The contribution of both arteries to blood flow in the septum and posterior wall of the left ventricle varies.

Coronary blood flow (Qcor) is phasic, i.e., the amount of blood in the coronary arteries fluctuates during the cardiac cycle due to extremely high rises in extravascular tissue pressure during systole (\rightarrow **B**, **C**). The blood flow in the epicardial coronary artery branches and subepicardial vessels remains largely unaffected by these pressure fluctuations. However, the subendocardial vessels of the left ventricle are compressed during systole when the extravascular pressure in that region (~ pressure in left ventricle, P_{IV}) exceeds the pressure in the lumen of the vessels ($\rightarrow C$). Consequently, the left ventricle is mainly supplied during diastole (\rightarrow **B** middle). The fluctuations in right ventricular blood flow are much less distinct because right ventricular pressure (P_{RV}) is lower $(\rightarrow B, C)$.

Myocardial O2 consumption (\dot{V} o₂) is defined as \dot{Q}_{cor} times the arteriovenous O2 concentration difference, (C_a – C_v)o₂. The myocardial (C_a – C_v)o₂ is relatively high (0.12 L/L blood), and oxygen extraction at rest ($[C_a$ – $C_v]$ o₂/ C_a o₂ = 0.12/0.21) is almost 60% and, thus, not able to rise much further. Therefore, an increase in \dot{Q}_{cor} is practically the only way to increase myocardial \dot{V} o₂ when the O₂ demand rises (\rightarrow **D**, right side).

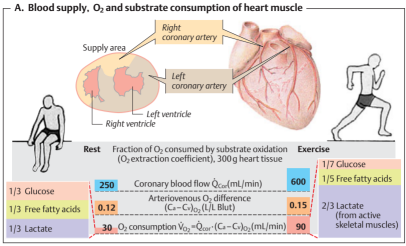
Adaptation of the myocardial O_2 supply according to need is therefore primarily achieved by adjusting vascular resistance (\rightarrow D, left side). The (distal) coronary vessel resistance can normally be reduced to about $^{1}/_{4}$ the resting value (coronary reserve). The coronary blood flow Q_{cor} (approx. 250 mL/min at rest) can therefore be increased as much as 4–5 fold. In other words, approx. 4 to 5 times more O_2 can be supplied during maximum physical exertion.

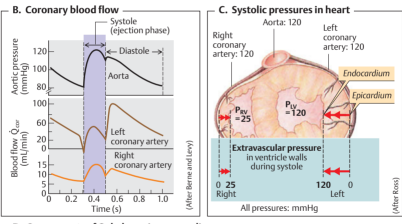
Arteriosclerosis (atherosclerosis) of the coronary arteries leads to luminal narrowing and a resultant decrease in poststenotic pressure. Dilatation of the distal vessels then occurs as an autoregulatory response (see below). Depending on the extent of the stenosis, it may be necessary to use a fraction of the coronary reserve, even during rest. As a result, lower or insufficient quantities of O_2 will be available to satisfy increased O_2 demand, and coronary insufficiency may occur $(\rightarrow \mathbf{D})$

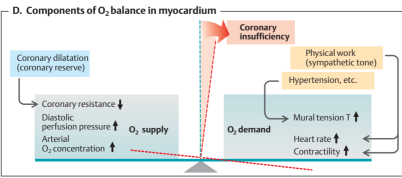
Myocardial O2 demand increases with cardiac output (increased pressure-volume-work/time). i.e., in response to increases in heart rate and/or contractility, e.g., during physical exercise ($\rightarrow \mathbf{D}$, right). It also increases as a function of mural tension (Tventr) times the duration of systole (tension-time index). Since $T_{ventr} = P_{ventr} \cdot r_{ventr} / 2w$ (Laplace's law \rightarrow Eq. 8.4b, p. 188), O₂ demand is greater when the ventricular pressure (Pventr) is high and the stroke volume small than when P_{ventr} is low and the stroke volume high, even when the same amount of work (P \times V) is performed. In the first case, the efficiency of the heart is reduced. When the ventricular pressure Pventr is elevated, e.g., in hypertension, the myocardium therefore requires more O₂ to perform the same amount of work (\rightarrow **D**, right).

Since the myocardial metabolism is aerobic, an increased O_2 demand quickly has to lead to vasodilatation. The following factors are involved in the **coronary vasodilatation**:

- Metabolic factors: (a) oxygen deficiency since O₂ acts as a vasoconstrictor; (b) Adenosine; oxygen deficiencies result in insufficient quantities of AMP being re-converted to ATP, leading to accumulation of adenosine, a degradation product of AMP. This leads to A₂ receptor-mediated vasodilatation; (c) Accumulation of lactate and H¹ ions (from the anaerobic myocardial metabolism); (d) prostaglandin I₂.
- ◆ Endothelial factors: ATP (e.g., from platelets), bradykinin, histamine and acetylcholine are vasodilators. They liberate nitric oxide (NO) from the endothelium, which diffuses into vascular muscle cells to stimulate vasodilatation (→ p. 279 E).
- Neurohumoral factors: Norepinephrine released from sympathetic nerve endings and adrenal epinephrine have a vasodilatory effect on the distal coronary vessels via β_2 adrenoceptors.
- Myocardial energy sources. The myocardium can use the available glucose, free fatty acids, lactate and other molecules for ATP production. The oxidation of each of these three energy substrates consumes a certain fraction of myocardial O_2 (O_2 extraction coefficient); accordingly, each contributes approx. one-third of the produced ATP at rest. The myocardium consumes increasing quantities of *lactate* from the skeletal muscles during physical exercise (\rightarrow **A**, \rightarrow **p**. 72 and 282).







The blood flow must be regulated to ensure an adequate blood supply, even under changing environmental conditions and stress (cf. p.74). This implies (a) optimal regulation of cardiac activity and blood pressure (homeostasis), (b) adequate perfusion of all organ systems, and (c) shunting of blood to active organ systems (e.g., muscles) at the expense of the resting organs (e.g., gastrointestinal tract) to keep from overtaxing the heart (→A).

Regulation of blood flow to the organs is mainly achieved by changing the diameter of blood vessels. The muscle tone (tonus) of the vascular smooth musculature changes in response to (1) local stimuli (\rightarrow B2a|b), (2) hormonal signals (\rightarrow B3 a|b) and (3) neuronal signals (\rightarrow B1 a|b). Most blood vessels have an intermediary muscle tone at rest (resting tone). Many vessels dilate in response to denervation, resulting in a basal tone. This occurs due to spontaneous depolarization of smooth muscle in the vessels (see also p.70).

Local Regulation of Blood Flow (Autoregulation)Autoregulation has two **functions**:

- ◆ Autoregulatory mechanisms help to maintain a constant blood flow to certain organs when the blood pressure changes (e.g., renal vessels constrict in response to rises in blood pressure; → p. 150).
- ♦ Autoregulation also functions to adjust the blood flow according to changes in metabolic activity of an organ (metabolic autoregulation); the amount of blood flow to the organ (e.g., cardiac and skeletal muscle; \rightarrow A and p. 201) can thereby increase many times higher than the resting level.

Types of autoregulatory mechanism:

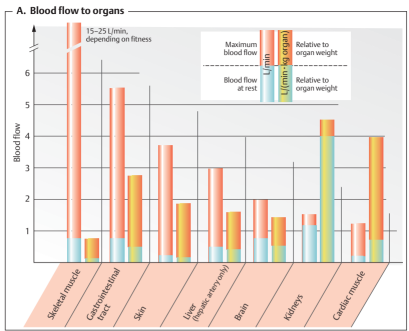
- Myogenic effects arising from the vascular musculature of the lesser arteries and arterioles (Bayliss effect) ensure that these vessels contract in response to blood pressure-related dilatation (→ B2a) in certain organs (e.g., kidneys, gastrointestinal tract and brain), but not in others (e.g., skin and lungs).
- ◆ Oxygen deficiencies generally cause the blood vessels to *dilate*. Hence, the degree of blood flow and O₂ uptake increase with increasing O₂ consumption. In the lungs, on the

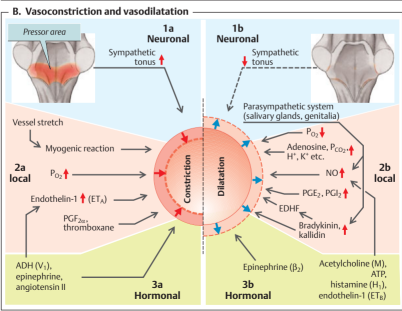
- other hand, a low Po_2 in the surrounding alveoli causes the vessels to *contract* (*hypoxic vasoconstriction*; \rightarrow p. 122).
- ◆ Local metabolic (chemical) effects: An increase in local concentrations of *metabolic products* such as CO₂, H⁺, ADP, AMP, adenosine, and K⁺ in the interstitium has a *vasodilatory effect*, especially in precapillary arterioles. The resulting rise in blood flow not only improves the supply of substrates and O₂, but also accelerates the efflux of these metabolic products from the tissue. The blood flow to the *brain* and *myocardium* (→ p. 210) is almost entirely subject to local metabolic control. Both local metabolic effects and O₂ deficiencies lead to an up to 5-fold increase in blood flow to an affected region in response to the decreased blood flow (*reactive hyperemia*).
- Vasoactive substances: A number of vasoactive substances such as prostaglandins play a role in autoregulation (see below).

Hormonal Control of Circulation

Vasoactive substances. Vasoactive hormones either have a direct effect on the vascular musculature (e.g., epinephrine) or lead to the local release of vasoactive substances (e.g., nitric oxide, endothelin) that exert local paracrine effects (\rightarrow **B**).

- Nitric (mon)oxide (NO) acts as a vasodilatory agent. NO is released from the endothelium when acetylcholine (M receptors), ATP, endothelin (ET_B receptors), or histamine (H₁ receptors) binds with an endothelial cell (\rightarrow p. 278). NO then diffuses to and relaxes vascular myocytes in the vicinity.
- ◆ Endothelin-1 can lead to vasodilatation by inducing the release of NO from the endothelium by way of ET_B receptors (see above), or can cause vasoconstriction via ET_A receptors in the vascular musculature. When substances such as angiotensin II or ADH (=vasopressin; V₁ receptor) bind to an endothelial cell, they release endothelin-1, which diffuses to and constricts the adjacent vascular muscles with the aid of ET_A receptors.
- ♦ Epinephrine (E): High concentrations of E from the adrenal medulla (\rightarrow p. 86) have a *vasoconstrictive* effect (α_1 -adrenoceptors), whereas low concentrations exert *vasodilatory* effects by way of β_2 adrenoceptors in the *myo*-





cardium, skeletal muscle and liver $(\rightarrow C)$. The effect of E mainly depends on which type of adrenoceptor is predominant in the organ, α_1 adrenoceptors are predominant in the blood vessels of the kidney and skin.

- **◆ Eicosanoids** (→ p. 269): Prostaglandin (PG) $F_{2\alpha}$ and thromboxane A_2 (released from platelets, \rightarrow p. 102) and B₂ have vasoconstrictive effects, while PGI₂ (= prostacyclin, e.g. released from endothelium) and PGE2 have vasodilatory effects. Another vasodilator released from the endothelium (e.g., by bradykinin; see below) opens K+ channels in vascular myocytes and hyperpolarizes them, leading to a drop in the cytosolic Ca2+ concentration. This endothelium-derived hyperpolarizing factor (EDHF), has been identified as a 11,12-epoxyeicosatrienoic acid (11.12-EET).
- ◆ Bradykinin and kallidin are vasodilatory agents cleaved from kininogens in blood plasma by the enzyme kallikrein. Histamine also acts as a vasodilator. All three substances influence also vessel permeability (e.g., during infection) and blood clotting.

Neuronal Regulation of Circulation

Neuronal regulation of blood flow $(\rightarrow B1a/b)$ mainly involves the lesser arteries and greater arterioles (\rightarrow p. 188), while that of venous return to the heart (\rightarrow p. 188) can be controlled by dilating or constricting the veins (changes in their blood storage capacity). Both mechanisms are usually controlled by the sympathetic nervous system (\rightarrow B1a and p. 78ff.), whereby norepinephrine (NE) serves as the postganglionic transmitter (except in the sweat glands). NE binds with the α_1 adrenoceptors on blood vessels, causing them to constrict (\rightarrow **B**). Vasodilatation is usually achieved by decreasing the tonus of the sympathetic system (\rightarrow **B1b**). This does not apply to blood vessels in salivary glands (increased secretion) or the genitals (erection), which dilate in response to parasympathetic stimuli. In this case, vasoactive substances (bradykinin and NO, respectively) act as the mediators. Some neurons release calcitonin gene-related peptide (CGRP), a potent vasodilator.

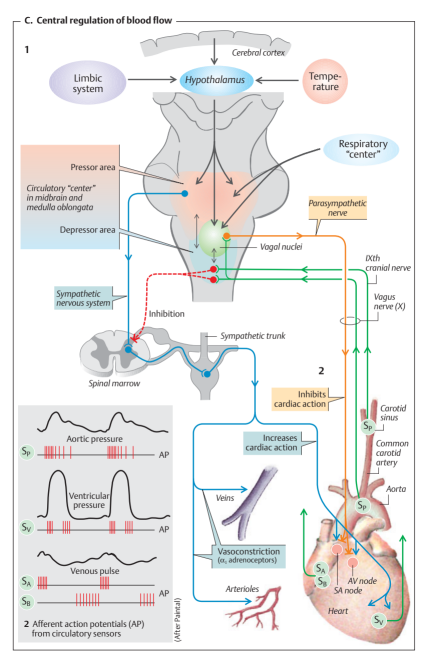
Neuronal regulation of blood flow to organs occurs mainly: (a) via central co-innervation (e.g., an impulse is simultaneously sent from

the cerebral cortex to circulatory centers when a muscle group is activated, or (b) via neuronal feedback from the organs whose activity level and metabolism have changed. If the neuronal and local metabolic mechanisms are conflicting (e.g., when sympathetic nervous stimulation occurs during skeletal muscle activity), the metabolic factors will predominate. Vasodilatation therefore occurs in the active muscle while the sympathetic nervous system reduces the blood flow to the inactive muscles. Blood flow to the skin is mainly regulated by neuronal mechanisms for the purpose of controlling heat disposal (temperature control; → p. 224). Hypovolemia and hypotension lead to centralization of blood flow, i.e., vasoconstriction in the kidney (oliguria) and skin (pallor) occurs to increase the supply of blood to vital organs such as the heart and central nervous system (\rightarrow p. 218).

During exposure to extremely low temperatures, the cold-induced vasoconstriction of cutaneous vessels is periodically interrupted to supply the skin with blood to prevent tissue damage (**Lewis response**). Axoaxonal reflexes in the periphery play a role in this response, as afferent cutaneous nerve fibers transmit signals to efferent vasomotor axons. Skin reddening in response to scratching (dermatographism) is also the result of axoaxonal reflexes.

Central regulation of blood flow $(\rightarrow C)$ is the responsibility of the CNS areas in the medulla oblongata and pons. They receive information from circulatory sensors (S) or receptors (a) in the high-pressure system (barosensors or pressure sensors, SP, in the aorta and carotid artery); (b) in the low-pressure system (stretch sensors in the vena cava and atria, S_A and S_B); and (c) in the left ventricle (Sv). The sensors measure arterial blood pressure (SP), pulse rate (S_P and S_V) and filling pressure in the low pressure system (indirect measure of blood volume). The A sensors (SA) mainly react to atrial contraction, whereas the B sensors (SB) react to passive filling stretch (\rightarrow C2). If the measured values differ from the set-point value, the circulatory control centers of the CNS transmit regulatory impulses through efferent nerve fibers to the heart and blood vessels $(\rightarrow \mathbf{D} \text{ and p. 5 C2}).$

Situated laterally in the circulatory "center" is a **pressor area** (\rightarrow **C**, reddish zone), the neu-



rons of which (blue arrows) continuously transmit sympathetic nerve impulses to the heart to increase its activity (heart rate, conduction and contractility). Their effects on vessels are predominantly vasoconstrictive (resting tone). The pressor area is in close contact with more medial neurons (**depressor area**, light blue area in \mathbf{C}). The pressor and depressor area are connected to the dorsal nuclei of the vagus nerve ($\rightarrow \mathbf{C}$, green), the stimulation of which reduces the heart rate and cardiac impulse conduction rate ($\rightarrow \mathbf{C}$, orange arrows).

Homeostatic circulatory reflexes include signals along afferent nerve tracts ($\rightarrow D3a/b$) that extend centrally from the pressosensors in the aorta and carotid sinus (\rightarrow **C**, green tracts). The main purpose of homeostatic control is to maintain the arterial blood pressure at a stable level. Acute increases in blood pressure heighten the rate of afferent impulses and activate the depressor area. By way of the vagus nerve, parasympathetic neurons ($\rightarrow \mathbf{C}$, orange tract) elicit the depressor reflex response, i.e., they decrease the cardiac output (CO). In addition, inhibition of sympathetic vessel innervation causes the vessels to dilate, thereby reducing the peripheral resistance (TPR; \rightarrow D4a/b). Both of these mechanisms help to lower acute increases in blood pressure. Conversely, an acute drop in blood pressure leads to activation of pressor areas, which stimulates a rise in CO and TPR as well as venous vasoconstriction $(\rightarrow C$, blue tracts), thereby raising the blood pressure back to normal.

Due to the fast adaptation of pressosensors (differential characteristics, \rightarrow p. 312ff.), these regulatory measures apply to acute changes in blood pressure. Rising, for example, from a supine to a standing position results in rapid redistribution of the blood volume. Without control (orthostatic homeostatic reflex: → p. 204), the resulting change in venous return would lead to a sharp drop in arterial blood pressure. The circulatory centers also respond to falling Po₂ or rising Pco₂ in the blood (cross-links from respiratory center) to raise the blood pressure as needed.

In individuals with chronic **hypertension**, the input from the pressosensors is normal because they are fully adapted. Therefore, circulatory control centers cannot respond to and decrease the high pressures. On the contrary, they may even help to "fix" the

blood pressure at the high levels. Chronic hypertension leads to stiffening of the carotid sinus. This may also contribute to decreasing the sensitivity of carotid pressosensors in hypertension.

A temporary increase in venous return (e.g., after an intravenous infusion) also leads to an increase in heart action (\rightarrow **D**, right). This mechanism is known as the **Bainbridge reflex**. The physiological significance of this reflex is, however, not entirely clear, but it may complement the Frank–Starling mechanism (\rightarrow p. 202ff.).

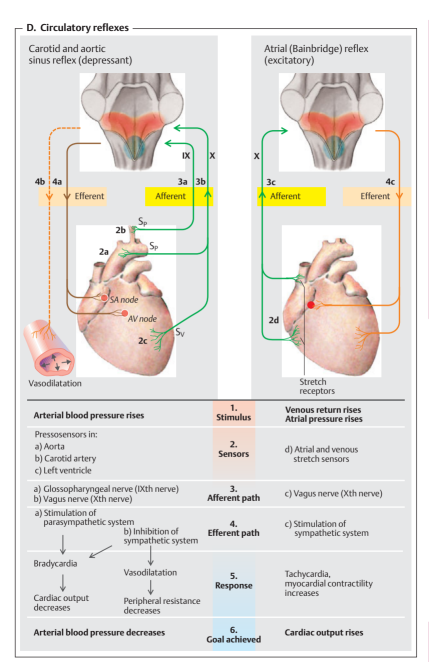
Hypertension

Hypertension is defined as a chronic increase in the systemic arterial blood pressure. The general criterion for diagnosis of hypertension is consistent elevation of resting blood pressure to more than 90 mmHg diastolic (→ p. 206). Untreated or inadequately managed hypertension results in stress and compensatory hypertrophy of the left ventricle which can ultimately progress to *left heart failure*. Individuals with hypertension are also at risk for arteriosclerosis and its sequelae (myocardial infarction, stroke, renal damage, etc.). Therefore, hypertension considerably shortens the life expectancy of a large fraction of the population.

The **main causes of hypertension** are (a) increased extracellular fluid (ECF) volume with increased venous return and therefore increased cardiac output (volume hypertension) and (b) increased total peripheral resistance (resistance hypertension). As hypertension always leads to vascular changes resulting in increased peripheral resistance, type *a* hypertension eventually proceeds to type *b* which, regardless of how it started, ends in a vicious circle.

The ECF volume increases when more NaCl (and water) is absorbed than excreted. The usually high intake of dietary salt may therefore play a role in the development of **essential hypertension** (primary hypertension), the most common type of hypertension, at least in patients sensitive to salt. Volume hypertension can even occur when a relatively low salt intake can no longer be balanced. This can occur in renal insufficiency or when an adrenocortical tumor produces uncontrolled amounts of aldosterone, resulting in Na* retention.

Other important cause of hypertension is **pheochromocytoma**, a tumor that secretes epinephrine and norepinephrine and therefore raises the CO and TPR. **Renal hypertension** can occur due to renal artery stenosis and renal disease. This results in the increased secretion of *renin*, which in turn raises the blood pressure via the renin–angiotensin–aldosterone (RAA) system (\rightarrow p. 184).



Circulatory Shock

Shock is characterized by acute or subacute progressive *generalized failure of the circulatory system* with disruption of the microcirculation and *failure to maintain adequate blood flow to vital organs*. In most cases, the **cardiac output (CO) is insufficient** due to a variety of reasons, which are explained below.

- ◆ Hypovolemic shock is characterized by reduced central venous pressure and reduced venous return, resulting in an inadequate stroke volume (Frank–Starling mechanism). The blood volume can be reduced due to bleeding (hemorrhagic shock) or any other conditions associated with the external loss of fluids from the gastrointestinal tract (e.g., severe vomiting, chronic diarrhea), the kidneys (e.g., in diabetes mellitus, diabetes insipidus, high-dose diuretic treatment) or the skin (burns, profuse sweating without fluid intake). An internal loss of blood can also occur, e.g., due to bleeding into soft tissues, into the mediastinum or into the pleural and abdominal space.
- Cardiogenic shock: Acute heart failure can be caused by acute myocardial infarction, acute decompensation of heart failure or impairment of cardiac filling, e.g. in pericardial tamponade. The central venous pressure is higher than in hypovolemic shock.
 Shock can occur due to hormonal causes, such as adrenocortical insufficiency, diabetic coma or insulin overdose (hypoglycemic shock).
- Vasogenic shock: Reduced cardiac output can also be due to peripheral vasodilatation (absence of pallor) and a resultant drop of venous return. This occurs in Gram-positive septicemia (septic shock), anaphylactic shock, an immediate hypersensitivity reaction (food or drug allergy, insect bite/sting) in which vasoactive substances (e.g., histamines) are released.

Symptoms. Hypovolemic and cardiovascular shock are characterized by *decreased blood pressure* (weak pulse) *increased heart rate*, *pallor* with cold sweats (not observed with shock caused by vasodilatation), reduced urinary output (*oliguria*) and extreme *thirst*.

Shock index. The ratio of pulse rate (beats/min) to systolic blood pressure (mmHg), or *shock index*, provides a rough estimate of the extent of volume loss. An index of up to 0.5 indicates normal or < 10% blood loss; up to 1.0 = < 20-30% blood loss and impending shock; up to 1.5 = > 30-50% blood loss and manifest shock.

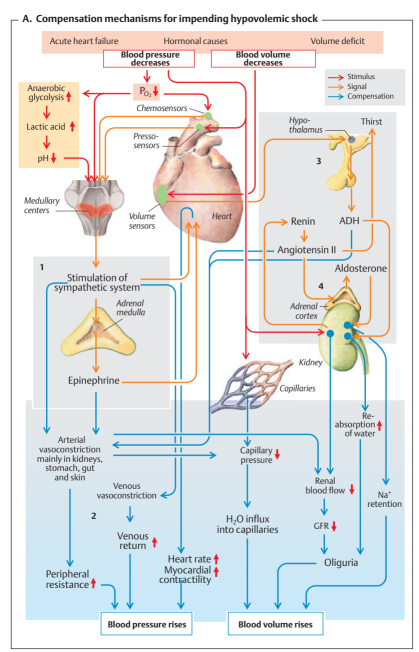
Most of the symptoms described reflect the counterregulatory measures taken by the

body during the *non-progressive phase* of shock in order to ward off progressive shock (\rightarrow **A**). Rapid-acting mechanisms for *raising the blood pressure* and slower-acting mechanisms to *compensate for volume losses* both play a role.

Blood pressure compensation (\rightarrow A left): A drop in blood pressure *increases sympathetic tonus* (\rightarrow A1 and p. 214). Arterial vasoconstriction (absent in shock due to vasodilatation) shunts the reduced cardiac output from the skin (pallor), abdominal organs and kidneys (oliguria) to vital organs such as the coronary arteries and brain. This is known as **centralization of blood flow** (\rightarrow A2). Sympathetic constriction of venous capacitance vessels (which raises ventricular filling), tachycardia and positive inotropism increase the diminished cardiac output to a limited extent.

Compensation for volume deficits $(\rightarrow A,$ right): When shock is imminent, the resultant drop in blood pressure and peripheral vasoconstriction lead to a reduction of capillary filtration pressure, allowing interstitial fluid to enter the bloodstream. Atrial stretch sensors detect the decrease in ECF volume (reduced atrial filling) and transmit signals to stop the atria from secreting atriopeptin (= ANP) and to start the secretion of antidiuretic hormone (ADH) from the posterior lobe of the pituitary (Gauer-Henry reflex; \rightarrow p. 170). ADH induces vasoconstriction (V1 receptors) and fluid retention (V₂ receptors). The drop in renal blood pressure triggers an increase in renin secretion and activation of the renin-angiotensin-aldosterone (RAA) system (\rightarrow p. 184). If these measures are successful in warding off the impending shock, the lost red blood cells are later replaced (via increased renal erythropoietin secretion, \rightarrow p. 88) and the plasma protein concentration is normalized by increased hepatic synthesis.

Manifest (or progressive) shock will develop if these homeostatic compensation mechanisms are unable to prevent impending shock and the patient does not receive medical treatment (infusion, etc.). Severe hypotension (< 90 mmHg systolic or < 60 mmHg mean blood pressure) can persist for extended periods, even in spite of volume replacement. The resulting development of hypoxia leads to organ damage and multiple organ failure, ultimately culminating in irreversible shock and death.



Fetal and Neonatal Circulation

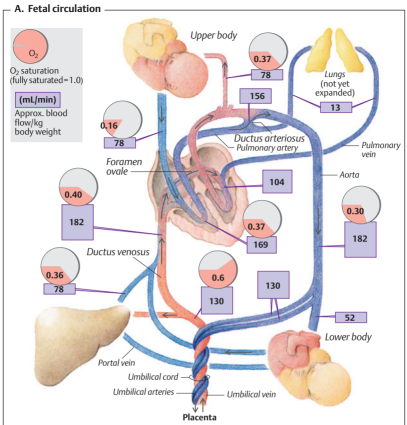
Placenta. The maternal placenta acts as the "gut" (absorption of nutrients), "kidneys" (removal of waste products) and "lungs" of the fetus (uptake of O_2 and elimination of CO_2). Although the *fetal* O_2 -hemoglobin dissociation curve is shifted to the left compared to that of adults (\rightarrow p. 129 C), only 60% (0.6) of placental hemoglobin is saturated with O_2 (\rightarrow **A**).

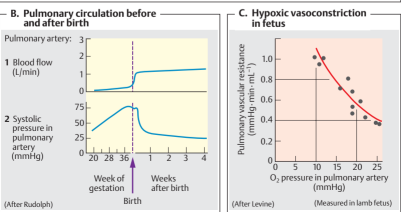
Fetal blood is distributed according to need. Inactive or hardly active organs receive little blood. The **fetal cardiac output** (from both ventricles) is about 0.2 L/min per kg body weight. The **fetal heart rate** rises from an initial 65 min⁻¹ (week 5) to 130–160 min⁻¹ in later weeks. Approx. 50% of the blood ejected from the heart flows through the placenta, the other half supplies the body (35%) and lungs (15%) of the fetus. This is supplied by the left and right heart, which function essentially *in parallel*. The serial connection of the systemic circulation to the pulmonary circuit (as in adults) is not fully necessary in the fetus.

Fetal circulation. The blood flows through the fetal body as follows $(\rightarrow A)$: After being arterialized in the placenta, the blood passes into the fetus via the umbilical vein and part of it travels through the ductus venosus (Arantii), thereby bypassing the liver. When entering the inferior vena cava, the blood mixes with venous blood from the lower half of the body. Guided by special folds in the vena cava, the mixed blood passes directly from right atrium to the left atrium through an opening in the atrial septum (foramen ovale). From the left atrium, it then proceeds to the left ventricle. While in the right atrium, the blood mingles with venous blood from the superior vena cava (only slight mixing), which is received by the right ventricle. Only about one-third of this blood reaches the lungs (due to high flow resistance since the lungs are not yet expanded, and due to hypoxic vasoconstriction, \rightarrow C and p. 122). The other two-thirds of the blood travels through the ductus arteriosus (Botalli) to the aorta (right-to-left shunt). Due to the low peripheral resistance (placenta), the blood pressure in the aorta is relatively low-only about 65 mmHg towards the end of pregnancy. The arteries of the head and upper body are supplied with partly arterialized blood from the left ventricle (\rightarrow A). This is important since brain tissue is susceptible to hypoxia. The remaining blood leaves the aorta and mixes with venous blood from the ductus arteriosus. As a result, the blood supplied to the lower half of the body has a relatively low O₂ concentration (O₂ saturation = 0.3; \rightarrow A). The majority of this blood returns via the *umbilical arteries* to the placenta, where it is oxygenated again.

Circulation during birth. The exchange of O₂. nutrients, and waste materials through the placenta stops abruptly during birth, This leads to a rise in blood Pco2, triggering chemosensors (\rightarrow p. 132) that induce a strong breathing reflex. The resultant inspiratory movement causes negative pressure (suction) in the thoracic cavity, which removes the blood from the placenta and umbilical vein (placental transfusion) and expands the lungs. The unfolding of the lungs and the rise in alveolar Po₂ reduces the resistance in the pulmonary circulation, and the blood flow increases while the pressure decreases (\rightarrow **B1, 2**). Meanwhile, the resistance in the systemic circulation increases due to occlusion or clamping of the umbilical cord. This changes the direction of blood flow in the ductus arteriosus, resulting in a left-to-right shunt. The pulmonary circulation therefore receives aortic blood for a few days after birth. The right atrial filling volume decreases due to the lack of placental blood, while that of the left atrium increases due to the increased pulmonary blood flow. Due to the resultant pressure gradient from the left to right atrium and to a decrease in vasodilatory prostaglandins, the foramen ovale usually closes within about 2 weeks after birth. The ductus arteriosus and ductus venosus also close, and the systemic and pulmonary circulation now form serial circuits.

Shunts occur when the foramen ovale or ductus arteriosus remains open, placing a strain on the heart. In **patent foramen ovale** (atrial septum defect), the blood flows from left atrium \rightarrow right atrium (left-to-right shunt) \rightarrow right ventricle (*volume overload*) \rightarrow lungs \rightarrow left atrium. In **patent ductus arteriosus**, the blood flows from aorta \rightarrow pulmonary artery (=left-to-right shunt) \rightarrow lungs (*pressure overload*) \rightarrow aorta.





Thermal Balance and Thermoregulation

Thermal Balance

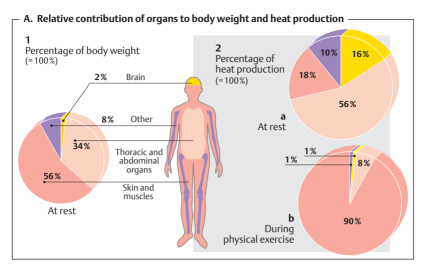
The body temperature of humans remains relatively constant despite changes in the environmental temperature. This homeothermy applies only to the **core temperature** (≈ 37 °C) of the body. The extremities and skin ("shell") exhibit **poikilothermy**, i.e., their temperature varies to some extent with environmental temperature. In order to maintain a constant core temperature, the body must balance the amount of heat it produces and absorbs with the amount it loses; this is thermoregulation $(\rightarrow p.224)$.

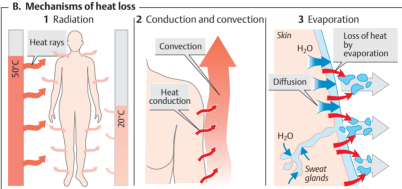
Heat production. The amount of heat produced is determined by energy metabolism $(\rightarrow p. 228)$. At rest, approximately 56% of total heat production occurs in the internal organs and about 18% in the muscles and skin (\rightarrow A2, top). During physical exercise, heat production increases several-fold and the percentage of heat produced by muscular work can rise to as much as 90% (\rightarrow A2, bottom). To keep warm, the body may have to generate additional voluntary (limb movement) and involuntary (shivering) muscle contractions. Newborns also have tissue known as brown fat, which enables them to produce additional heat without shivering (\rightarrow p. 225). Cold stimulates a reflex pathway resulting in norepinephrine release (β_3 -adrenergic receptors) in fatty tissues, which in turn stimulates (1) lipolysis and (2) the expression of lipoprotein lipase (LPL) and thermogenin, LPL increases the supply of free fatty acids (\rightarrow p. 254). Thermogenin localized in the inner mitochondrial membrane is an uncoupling protein that functions as an H+ uniporter (UCP1, \rightarrow p. 230). It short-circuits the H⁺ gradient across the inner mitochondrial membrane (\rightarrow p. 17/B2), thereby uncoupling the (heat-producing) respiratory chain of ATP production.

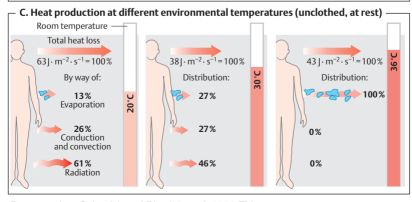
Heat produced in the body is absorbed by the bloodstream and conveyed to the body surface. In order for this internal flow of heat to occur, the temperature of the body surface must be lower than that of the body interior. The blood supply to the skin is the chief determinant of heat transport to the skin (\rightarrow p. 224).

Heat loss occurs by the physical processes of radiation, conduction, convection, and evaporation ($\rightarrow \mathbf{B}$).

- Radiation (→ B1, C). The amount of heat lost by radiation from the skin is chiefly determined by the temperature of the radiator (fourth power of its absolute temperature). Heat net-radiates from the body surface to objects or individuals when they are cooler than the skin, and net-radiates to the body from objects (sun) that are warmer than the skin. Heat radiates from the body into the environment when no radiating object is present (night sky). Heat radiation does not require the aid of any vehicle and is hardly affected by the air temperature (air itself is a poor radiator). Therefore, the body loses heat to a cold wall (despite warm air in between) and absorbs radiation from the sun or an infrared radiator without air (space) or cold air, respectively, in between.
- **2. Conduction** and **convection** (\rightarrow **B2, C**). These processes involve the transfer of heat from the skin to cooler air or a cooler object (e.g. sitting on rock) in contact with the body (conduction). The amount of heat lost by conduction to air increases greatly when the warmed air moves away from the body by natural convection (heated air rises) or forced convection (wind).
- **3. Evaporation** (\rightarrow **B3, C**). The first two mechanisms alone are unable to maintain adequate temperature homeostasis at high environmental temperatures or during strenuous physical activity. Evaporation is the means by which the body copes with the additional heat. The water lost by evaporation reaches the skin surface by diffusion (insensible perspiration) and by neuron-activated sweat glands $(\rightarrow B3$, pp. 73ff. and 225 D). About 2428 kJ (580 kcal) of heat are lost for each liter of water evaporating and thereby cooling the skin. At temperatures above 36°C or so, heat loss occurs by evaporation only ($\rightarrow \mathbf{C}$, right). At even higher environmental temperatures, heat is absorbed by radiation and conduction/convection. The body must lose larger amounts of heat by evaporation to make up for this. The surrounding air must be relatively dry in order for heat loss by evaporation to occur. Humid air retards evaporation. When the air is extremely humid (e.g., in a tropical rain forest), the average person cannot tolerate temperatures above 33 °C, even under resting conditions.







Thermoregulation

Thermoregulation maintains the **core temperature** $(\rightarrow A)$ at a constant **set point** $(\approx 37^{\circ}C)$ despite fluctuations in heat absorption, production, and loss $(\rightarrow p.222)$. The core temperature exhibits *circadian variation*. It fluctuates by about $0.6^{\circ}C$ and is lowest around 3 a.m., and highest around 6 p.m. $(\rightarrow p.381\ C)$. The set point changes are controlled by an intrinsic *biological clock* $(\rightarrow p.334)$. Extended set-point fluctuations happen during the menstrual cycle $(\rightarrow p.299/A3)$ and fever.

The *control center* for body temperature and *central thermosensors* are located in the **hypothalamus** (\rightarrow p. 330). Additional thermosensors are located in the spinal cord and skin (\rightarrow p. 314). The control center compares the actual core temperature with the set-point value and initiates measures to counteract any deviations (\rightarrow **D** and p. 4f.).

When the **core temperature rises** above the set point (e.g., during exercise), the body increases the internal heat flow (\rightarrow p. 222) by dilating the blood vessels of the skin. Moreover, arteriovenous anastomoses open in the periphery, especially in the fingers. The blood volume transported per unit time then not only conveys more heat, but also reduces the countercurrent exchange of heat between the arteries and their accompanying veins $(\rightarrow B)$. In addition, venous return in the extremities is re-routed from the deep, accompanying veins to the superficial veins. Sweat secretion also increases. The evaporation of sweat cools the skin, thereby creating the core/skin temperature gradient needed for the internal heat flow. Central warm sensors emit the signals that activate the sweat glands. (In this case, the thermosensors of the skin do not detect warmth because their environment is cooler than the core temperature). The efferent nerve fibers to the sweat glands are cholinergic fibers of the sympathetic nervous system ($\rightarrow \mathbf{D}$).

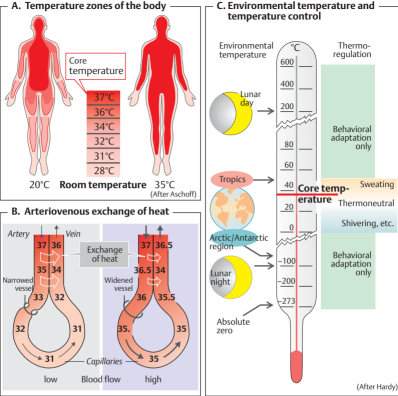
Acclimatization to high environmental temperatures (e.g., in the tropics) is a slow process that often takes years. Characteristically, the sweat secretion rate rises, the salt content of the sweat decreases, and thirst and thus H₂O intake increase.

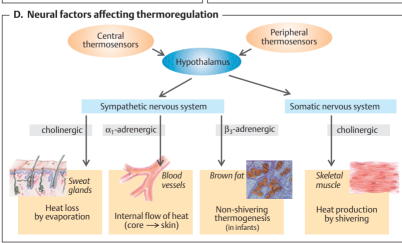
When the **core temperature falls** below set point, the body checks heat loss by constricting the blood vessels in the shell (\rightarrow **A**, left) and increases heat production by generating voluntary and involuntary (*shivering*) muscle activity (\rightarrow **D**). Although infants can quickly become hypothermic because of their high surface/volume ratio, their *brown fat* allows them to produce additional heat (*non-shivering thermogenesis*; \rightarrow **p**. 222). Upon exposure to low ambient temperatures, these three mechanisms are activated by the cold receptors of the skin (\rightarrow **p**. 314) *before* the core temperature falls.

The range of ambient temperatures between the sweating and shivering thresholds is known as the **thermoneutral zone**. It lies between ca. 27 °C and 32 °C in the nearly unclothed test subject. The only thermoregulatory measure necessary within this range is variation of blood flow to the skin. The narrow range of this zone shows the thermoregulatory importance of **behavior**. It involves choosing the appropriate clothing, seeking shade, heating or cooling our dwellings, etc. Behavioral adaptation is the chief factor in survival at extreme ambient temperatures (\rightarrow **C**).

The thermoneutral zone is subjectively perceived as the **comfort zone**. 95% of all subjects wearing normal office attire and performing normal office activities perceive an indoor climate with the following conditions to be comfortable: ambient and radiant (wall) temperature ≈ 23 °C, wind velocity < 0.1 m/s, and relative humidity ≈ 50 %. A resting, unclothed subject feels comfortable at about 28 °C and ca. 31 °C to 36 °C in water depending on the thickness of subcutaneous fat (heat isolator).

Fever. Exogenous (e.g., bacteria) and endogenous pyrogens (various interleukins and other cytokines from macrophages) can cause the set-point temperature to rise above normal. This is triggered by prostaglandin PGE₂ in the hypothalamus. In the initial phase of fever, the core temperature (although at its normal level) is too low compared to the elevated set-point. This results in shivering to raise the core temperature. As the fever decreases, i.e. the set-point returns toward the normal temperature, the core temperature is now too warm compared to the normalized set-point, resulting in vasodilatation and sweating to lower the core temperature again.





Nutrition and Digestion

Nutrition

An adequate diet must meet the body's energy requirements and provide a minimum of carbohydrates, proteins (incl. all essential amino acids) and fats (incl. essential fatty acids). Minerals (incl. trace elements), vitamins, and sufficient quantities of water are also essential. To ensure a normal passage time, especially through the colon, the diet must also provide a sufficient amount of roughage (indigestible plant fibers-cellulose, lignin, etc.).

The total energy expenditure (TEE) or total metabolic rate consists of (1) the basal metabolic rate (BMR), (2) the activity energy costs, and the (3) diet-induced thermogenesis (DIT; → p. 228, 231 A). TEE equals BMR when measured (a) in the morning (b) 20 h after the last meal, (3) resting, reclining, (4) at normal body temp., and (5) at a comfortable ambient temp. (\rightarrow p. 224). The BMR varies according to sex, age, body size and weight. The BMR for a young adult is ca. 7300 kJ/day (≈ 1740 kcal/ day; see p. 374 for units) in men, and ca. 20% lower in women, During physical activity, TEE increases by the following factors: 1.2-fold for sitting quietly, 3.2-fold for normal walking, and 8-fold for forestry work. Top athletes can perform as much as 1600W (= I/s) for two hours (e.g., in a marathon) but their daily TEE is much lower. TEE also increases at various degrees of injury (1.6-fold for sepsis, 2.1-fold for burns), 1 °C of fever increases TEE 1.13-fold.

Protein, fats and carbohydrates are the three basic energy substances $(\rightarrow B)$.

An adequate intake of **protein** is needed to maintain a proper nitrogen balance, i.e., balance of dietary intake and excretory output of nitrogen. The minimum requirement for protein is 0.5 g/kg BW per day (functional minimum). About half of dietary protein should be animal protein (meat, fish, milk and eggs) to ensure an adequate supply of essential amino acids such as histidine, isoleucine, leucine, lysine, methionine, phenylalanine, threonine, tryptophan and valine (children also require arginine). The content of most vegetable proteins is only about 50% of animal protein.

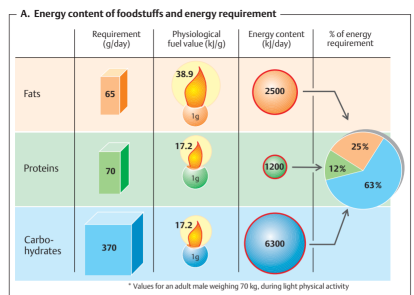
Carbohydrates (starch, sugar, glycogen) and fats (animal and vegetable fats and oils) pro-

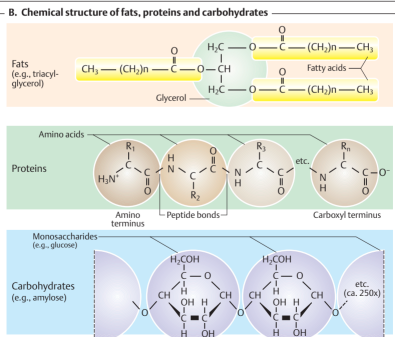
vide the largest portion of the energy requirement. They are basically interchangeable sources of energy. The energy contribution of carbohydrates can fall to about 10% (normally 60%) before metabolic disturbances occur.

Fat is not essential provided the intake of fat-soluble vitamins (vitamins E, D, K and A) and essential fatty acids (linoleic acid) is sufficient. About 25-30% of dietary energy is supplied by fat (one-third of which is supplied as essential fatty acids; $\rightarrow A$), although the proportion rises according to energy requirements (e.g., about 40% during heavy physical work). Western diets contain generally too much energy (more fats than carbohydrates!) considering the generally low level of physical activity of the Western lifestyle. Alcohol also contains superfluous energy (ca. 30 kJ/g = 7.2 kcal/g). The excessive intake of dietary energy leads to weight gain and obesity $(\rightarrow p.230)$.

An adequate intake of minerals (inorganic compounds), especially calcium (800 mg/day; \rightarrow p. 290ff.), iron (10–20 mg/day; \rightarrow p. 90) and iodine (0.15 mg/day; \rightarrow p. 288), is essential for proper body function. Many trace elements (As, F, Cu, Si, V, Sn, Ni, Se, Mn, Mo, Cr, Co) are also essential. The normal diet provides sufficient quantities of them, but excessive intake has toxic effects.

Vitamins (A, B₁, B₂, B₆, B₁₂, C, D₂, D₃, E, H (biotin), K1, K2, folic acid, niacinamide, pantothenic acid) are compounds that play a vital role in metabolism (usually function as coenzymes). However, the body cannot produce (or sufficient quantities of) them. A deficiency of vitamins (hypovitaminosis) can lead to specific conditions such as night blindness (vit. A), scurvy (vit. C), rickets (vit. D = calciferol; \rightarrow p. 292), anemia (vit. B_{12} = cobalamin; folic acid; \rightarrow p. 90), and coagulation disorders (vit. K; \rightarrow p. 104). An excessive intake of certain vitamins like vitamin A and D, on the other hand, can be toxic (hypervitaminosis).





Energy Metabolism and Calorimetry

The chemical energy of foodstuffs is first converted into energy-rich substances such as creatine phosphate and adenosine triphosphate (ATP). Energy, work and amount of heat are expressed in joules (J) or calories (cal) $(\rightarrow p. 374)$. The energy produced by hydrolysis of ATP (\rightarrow p. 41) is then used to fuel muscle activity, to synthesize many substances, and to create concentration gradients (e.g., Na+ or Ca^{2+} gradients: $\rightarrow p. 26ff.$). During all these energy conversion processes, part of the energy is always converted to heat (\rightarrow p. 38ff.).

In oxidative (aerobic) metabolism $(\rightarrow p$. 39 C), carbohydrates and fat combine with O2 to yield CO2, water, high-energy compounds (ATP etc.) and heat. When a foodstuff is completely oxidized, its biologically useable energy content is therefore equivalent to its physical caloric value (CV).

The **bomb calorimeter** (\rightarrow **A**), a device consisting of an insulated combustion chamber in a tank of water. is used to measure the CV of foodstuffs. A known quantity of a foodstuff is placed in the combustion chamber of the device and incinerated in pure O2. The surrounding water heats up as it absorbs the heat of combustion. The degree of warming is equal to the caloric value of the foodstuff.

Fats and carbohydrates are completely oxidized to CO2 and H2O in the body. Thus, their physiological fuel value (PFV) is identical to their CV. The mean PFV is 38.9 kJ/g (= 9.3 kcal/ g) for fats and $17.2 \, \text{kJ/g}$ (= 4.1 kcal/g) for digestible carbohydrates (→ p. 227 A). In contrast, proteins are not completely broken down to CO₂ and water in the human body but yield urea, which provides additional energy when it is oxidized in the bomb calorimeter. The CV of proteins (ca. 23 kJ/g) is therefore greater than their PFV, which is a mean of about 17.2 kJ/g or 4.1 kcal/g (\rightarrow p. 227 A).

At rest, most of the energy supplied by the diet is converted to heat, since hardly any external mechanical work is being performed. The heat produced is equivalent to the internal energy turnover (e.g., the work performed by the heart and respiratory muscles or expended for active transport or synthesis of substances).

In **direct calorimetry** $(\rightarrow B)$, the amount of heat produced is measured directly. The test subject, usually an experimental animal, is placed in a small chamber immersed in a known volume of ice. The amount of heat produced is equivalent to the amount of heat absorbed by the surrounding water or ice. This is respectively calculated as the rise in water temperature or the amount of ice that melts.

In indirect calorimetry, the amount of heat produced is determined indirectly by measuring the amount of O_2 consumed (\dot{V}_{O_2}) : \rightarrow p. 120). This method is used in humans. To determine the total metabolic rate (or TEE; \rightarrow p. 226) from \dot{V}_{O_2} , the caloric equivalent (CE) of a foodstuff oxidized in the subject's metabolism during the measurement must be known. The CE is calculated from the PFV and the amount of O2 needed to oxidize the food. The PFV of glucose is 15.7 kJ/g and 6 mol of O₂ (6×22.4L) are required to oxidize 1 mol (= 180 g) of glucose (\rightarrow C). The oxidation of 180 g of glucose therefore generates 2827 kJ of heat and consumes 134.4 L of O2 resulting in a CE of 21 kJ/L. This value represents the CE for glucose under standard conditions (0 °C; \rightarrow **C**). The mean CE of the basic nutrients at 37°C is 18.8 kJ/L O₂ (carbohydrates), 17.6 kJ/L O₂ (fats) and 16.8 kJ/L O₂ (proteins).

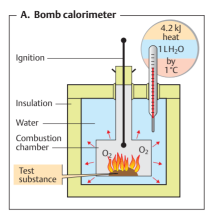
The oxidized nutrients must be known in order to calculate the metabolic rate from the CE. The respiratory quotient (RQ) is a rough measure of the nutrients oxidized. RQ = \dot{V}_{CO_2} / \dot{V}_{O_2} (\rightarrow p. 120). For pure carbohydrates oxidized, RQ = 1.0. This can be illustrated for glucose as follows:

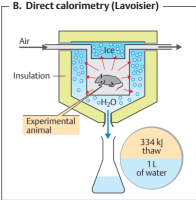
 $C_6H_{12}O_6 + 6 O_2$ $6 CO_2 + 6 H_2O$ [10.1] The oxidation of the fat *tripalmitin* yields: 2 C₅₁H₉₈O₆ + 145 O₂ 102 CO2 + 98 H2O

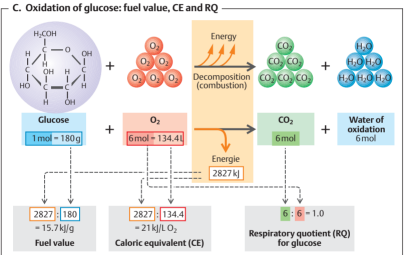
[10.2]

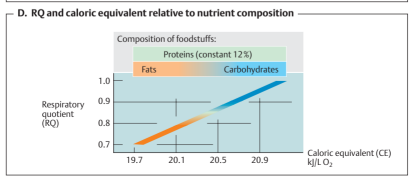
The RQ of tripalmitin is therefore 102/145 = 0.7. Since the protein fraction of the diet stays relatively constant, each RQ between 1 and 0.7 can be assigned a CE (\rightarrow **D**). Using the known CE, the TEE can be calculated as $CE \cdot \dot{V}_{O_2}$.

Food increases the TEE (diet-induced thermogenesis, DIT) because energy must be consumed to absorb and store the nutrients. The DIT of protein is higher than that of other substances, e.g., glucose.









Energy Homeostasis and Body Weight

Fat depots are by far the body's largest energy reserve. Accurate long-term homeostasis of energy absorption and consumption is necessary to keep the size of the fat depots constant, i.e., to maintain *lipostasis*. Considering that a person's body weight (BW) mainly varying with the weight of the fat depots, it is obvious that energy homeostasis is synonymous with the regulation of body weight (→ A).

The **body mass index (BMI)** is commonly used to determine whether an individual is underweight, overweight or in the normal weight range. BMI is calculated from body weight (kg) and height (m) as follows:

BMI =
$$(weight/height)^2$$
 [10.3]

The normal BMI range is 19–24 in women and 20–25 in men. The "normal" BMI range is defined as the values at which the mean life expectancy is highest. An abnormally high body mass index (BMI > 24 or 25 = overweight; BMI > 30 = obesity) reduces the life expectancy since this is often associated with diabetes mellitus (type II), hypertension and cardiac disease.

The following **regulatory mechanisms** serve to keep the fat depots and thus the body weight constant $(\rightarrow B)$:

- The hypothalamus, the center responsible for feedback control of BW, maintains communication with the limbic system, cerebral cortex and brain stem. It sends and receives:
- Afferent messages concerning the size of fat depots. **Leptin**, a 16-kDa proteohormone produced by fat cells, is the main indicator for this. The plasma leptin concentration rises as fat cell mass increases.
- ♦ Efferent commands (a) to reduce nutrient absorption and increase energy consumption when plasma leptin are high ("fat reserve high!"), and (b) to increase nutrient absorption and decrease energy consumption when plasma leptin levels are low ("fat reserve low!") (\rightarrow B).

Leptin receptors. Leptin binds with *type b leptin receptors* (**Ob-Rb**) of the hypothalamus (mainly in dorsomedial, ventromedial, lateral, paraventricular and arcuate nuclei). Certain neurons with Ob-Rb lie in front of the bloodbrain barrier. (T lymphocytes and B cells of the pancreas are also equipped with Ob-Rb receptors.)

Effects of leptin. In contrast to primary starvation, the weight loss induced by leptin is restricted to the body's fat depots and completes the feedback loop of the regulatory process. The effects of leptin are chiefly mediated by two neurotransmitters located in the hypothalamus: α -MSH and NPY (\rightarrow B).

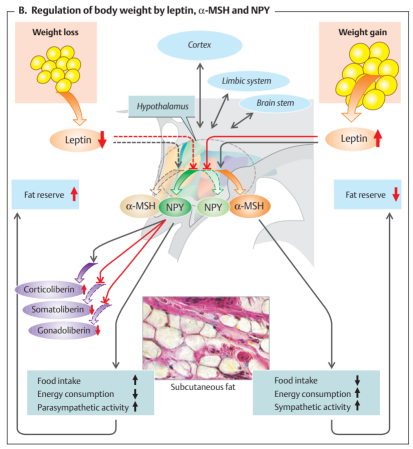
• α-MSH. Leptin stimulates the release of α-MSH (α-melanocyte-stimulating hormone), one of the melanocortins (MC) synthesized from POMC (\rightarrow p. 280). α-MSH inhibits the absorption of nutrients and increases sympathetic nervous activity and energy consumption via MC4 receptors in various areas of the hypothalamus.

The **mechanism** by which α-MSH increases energy consumption is not entirely clear, but an involuntary increase in ordinary skeletal muscle activity and tone appears to occur. In addition, **uncoupling proteins** (type UCP2 and UCP3) that were recently discovered in skeletal muscle and white fat make the membranes of the mitochondria more permeable to H^+ , thereby uncoupling the respiratory chain. As a result, chemical energy is converted into more heat and less ATP. The action of these UCPs, the expression of which may be stimulated by α -MSH, is therefore similar to that of thermogenin (UCP1; \rightarrow p. 222).

 NPY. Leptin inhibits the hypothalamic release of NPY (neuropeptide Y), a neuropeptide that stimulates hunger and appetite, increases the parasympathetic activity and reduces energy consumption.

Leptin deficiency. Since NPY increases the secretion of gonadotropin-releasing hormone (GnRH), extreme weight loss results in *amenorrhea* (\rightarrow B). Certain *genetic defects* can lead to impaired leptin production (*ab* [obesity] *gene*) or impaired leptin receptor function (*db* [diabetes] *gene*). The symptoms include arrested development in puberty and childhood *obesity*.

Other neurotransmitters and neuropeptides also play a role in the long-term regulation of fat depots. Some like $orexin\ A/B$ and norepinephrine (α_2 -adrenoceptors) are orexigenic, i.e. they stimulate the appetite, while others like CCK, CRH, CART (cocaine and amphetamine-regulated transcript), insulin, and serotonin are anorexigenic. Peptides like CCK, GLP-1 (glucagon-like peptide amides), somatostatin, glucagon, and GRP (gastrin-releasing peptide) signal satiety, i.e., that one has had enough to eat. Together with gustatory stimuli and stretch receptors of the stomach wall, these satiety peptides help to limit the amount of food consumed with each meal.



Gastrointestinal (GI) Tract: Overview, Immune Defense, Blood Flow

Food covering the body's energy and nutrient requirements (\rightarrow p. 228ff.) must be swallowed, processed and broken down (digestion) before it can be absorbed from the intestines. The three-layered GI musculature ensures that the GI contents are properly mixed and transported. The passage time through the different GI segments varies and is largely dependent on the composition of the food (see A for mean passage times).

Solid food is chewed and mixed with **saliva**, which lubricates it and contains immunocompetent substances (see below) and enzymes. The **esophagus** rapidly transports the food bolus to the stomach. The lower esophageal sphincter opens only briefly to allow the food to pass. The **proximal stomach** mainly serves as a food reservoir. Its tone determines the rate at which food passes to the **distal stomach**, where it is further processed (chyme formation) and its proteins are partly broken down. The distal stomach (including the pylorus) is also responsible for portioning chyme delivery to the small intestine. The stomach also secretes $intrinsic factor (\rightarrow p. 90)$.

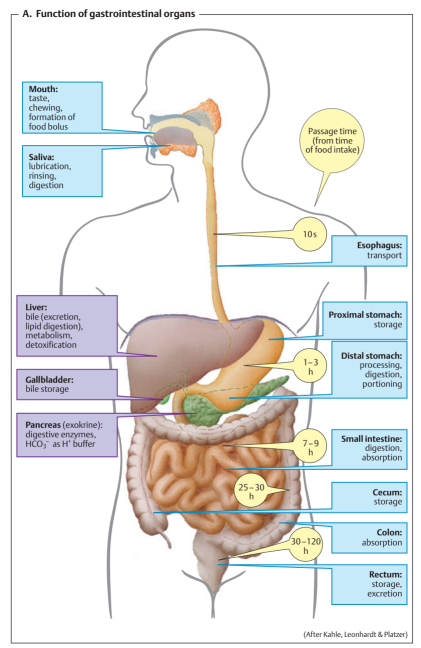
In the small intestine, enzymes from the pancreas and small intestinal mucosa break down the nutrients into absorbable components. HCO₃ - in pancreatic juices neutralizes the acidic chyme. Bile salts in bile are essential for fat digestion. The products of digestion (monosaccharides, amino acids, dipeptides, monoglycerides and free fatty acids) as well as water and vitamins are absorbed in the small intestine.

Waste products (e.g. bilirubin) to be excreted reach the feces via bile secreted by the liver. The liver has various other metabolic functions. It serves, for example, as an obligatory relay station for metabolism and distribution of substances reabsorbed from the intestine (via the portal vein, see below), synthesizes plasma proteins (incl. albumin, globulins, clotting factors, apolipoproteins etc.) and detoxifies foreign substances (biotransformation) and metabolic products (e.g., ammonia) before they are excreted.

The **large intestine** is the last stop for water and ion absorption. It is colonized by *bacteria* and contains storage areas for feces (**cecum**, **rectum**).

Immune defense. The large internal surface area of the GI tract (roughly 100 m²) requires a very effective immune defense system. Saliva contains mucins, immunoglobulin A (IgA) and lysozyme that prevent the penetration of pathogens. Gastric juice has a bactericidal effect. Peyer's patches supply the GI tract with immunocompetent lymph tissue. M cells (special membranous cells) in the mucosal epithelium allow antigens to enter Peyer's patches. Together with macrophages, the Peyer's patches can elicit immune responses by secreting $IqA (\rightarrow p. 98)$. IgA is transported to the intestinal lumen by transcytosis ($\rightarrow p.30$). In the epithelium, IgA binds to a secretory component, thereby protecting it from digestive enzymes. Mucosal epithelium also contains intraepithelial lymphocytes (IEL) that function like T killer cells (\rightarrow p. 98). Transmitter substances permit reciprocal communication between IEL and neighboring enterocytes. Macrophages of the hepatic sinusoids (Kupffer's cells) are additional bastions of immune defense. The physiological colonies of intestinal flora in the large intestine prevent the spread of pathogens, IgA from breast milk protects the GI mucosa of neonates.

Blood flow to the stomach, gut, liver, pancreas and spleen (roughly 30% of cardiac output) is supplied by the three main branches of the abdominal aorta. The intestinal circulation is regulated by local reflexes, the autonomic nervous system, and hormones. Moreover, it is autoregulatory, i.e., largely independent of systemic blood pressure fluctuations. Blood flow to the intestines rises sharply after meals (acetylcholine, vasoactive intestinal peptide VIP, etc. function as vasodilatory transmitters) and falls during physical activity (transmitters: norepinephrine, etc.). The venous blood carries substances reabsorbed from the intestinal tract and enters the liver via the portal vein. Some components of reabsorbed fat are absorbed by the intestinal lymph, which transports them to the greater circulation while bypassing the liver.



Neural and Hormonal Integration

Endocrine and paracrine hormones and neurotransmitters control GI motility, secretion, perfusion and growth. Reflexes proceed within the mesenteric and submucosal plexus (enteric nervous system, ENS), and external innervation modulates ENS activity.

Local reflexes are triggered by stretch sensors in the walls of the esophagus, stomach and gut or by chemosensors in the mucosal epithelium and trigger the contraction or relaxation of neighboring smooth muscle fibers. *Peristaltic reflexes* extend further towards the oral (ca. 2 mm) and anal regions (20–30 mm). They are mediated in part by interneurons and help to propel the contents of the lumen through the GI tract (*peristalsis*).

External innervation of the GI tract (cf. p.78ff.) comes from the *parasympathetic nervous system* (from lower esophagus to ascending colon) and *sympathetic nervous system*. Innervation is also provided by *visceral afferent fibers* (in sympathetic or parasympathetic nerves) through which the afferent impulses for *supraregional reflexes* flow.

ENS function is largely independent of external innervation, but **external innervation** has some advantages (a) rapid transfer of signals between relatively distant parts of the GI tract via the abdominal ganglia (short visceral afferents) or CNS (long visceral afferents); (b) GI tract function can be ranked subordinate to overall body function (c) GI tract activity can be processed by the brain so the body can become aware of them (e.q., stomach ache).

Neurotransmitters. Norepinephrine (NE) is released by the adrenergic postganglionic neurons, and acetylcholine (ACh) is released by pre- and postganglionic (enteric) fibers (→ p. 78ff.). VIP (vasoactive intestinal peptide) mediates the relaxation of circular and vascular muscles of the GI tract. Met- and leuenkephalin intensify contraction of the pyloric, ileocecal and lower esophageal sphincters by binding to opioid receptors. GRP (gastrin-releasing peptide) mediates the release of gastrin. CGRP (calcitonin gene-related peptide) stimulates the release of somatostatin (SIH).

All **endocrine hormones** effective in the GI tract are *peptides* produced in endocrine cells

of the mucosa. (a) Gastrin and cholecystokinin (CCK) and (b) secretin and GIP are structurally similar; so are glucagon (\rightarrow p. 282ff.) and VIP. High concentrations of hormones from the same family therefore have very similar effects.

Gastrin occurs in short (G17 with 17 amino acids, AA) and long forms (G34 with 34 AA). G17 comprises 90% of all antral gastrin. Gastrin is secreted in the antrum and duodenum. Its release (\rightarrow A1) via gastrin-releasing peptide (GRP) is subject to neuronal control; gastrin is also released in response to stomach wall stretching and protein fragments in the stomach. Its secretion is inhibited when the pH of the gastric/duodenal lumen falls below 3.5 (\rightarrow A1). The main effects of gastrin are acid secretion and gastric mucosal growth (\rightarrow A2).

Cholecystokinin, CCK (33 AA) is produced throughout small intestinal mucosa. Longchain fatty acids, AA and oligopeptides in the lumen stimulate the release of CCK (\rightarrow A1). It causes the gallbladder to contract and inhibits emptying of the stomach. In the pancreas, it stimulates growth, production of enzymes and secretion of HCO_3^- (via secretin, see below) (\rightarrow A2).

Secretin (27 AA) is mainly produced in the duodenum. Its release is stimulated by acidic chyme (\rightarrow A1). Secretin inhibits acid secretion and gastric mucosal growth and stimulates HCO_3^- secretion (potentiated by CCK), pancreatic growth and hepatic bile flow (\rightarrow A2).

GIP (glucose-dependent insulinotropic peptide, 42 AA; formerly called gastric inhibitory polypeptide = enterogastrone) is produced in the duodenum and jejunum and released via protein, fat and carbohydrate fragments (e.g., glucose) (\rightarrow A1). GIP inhibits acid secretion (\rightarrow A2) and stimulates insulin release (this is why oral glucose releases more insulin than intravenous glucose).

Motilin (22 AA) is released by neurons in the small intestine and regulates interdigestive motility (\rightarrow A1, 2).

Paracrine transmitters. Histamine, somatostatin and prostaglandin are the main paracrine transmitters in the GI tract.

Saliva

The functions of saliva are reflected by its constituents. Mucins serve to lubricate the food. making it easier to swallow, and to keep the mouth moist to facilitate masticatory and speech-related movement. Saliva dissolves compounds in food, which is a prerequisite for taste buds stimulation (\rightarrow p. 338) and for dental and oral hygiene. Saliva has a low NaCl concentration and is hypotonic, making it suitable for rinsing of the taste receptors (NaCl) while eating. Infants need saliva to seal the lips when suckling. Saliva also contains α -amylase, which starts the digestion of starches in the mouth, while immunoglobulin A and lysozyme are part of the immune defense system $(\rightarrow p. 94ff.)$. The high HCO₃⁻ concentration in saliva results in a pH of around 7, which is optimal for α-amylase-catalyzed digestion. Swallowed saliva is also important for buffering the acidic gastric juices refluxed into the esophagus (\rightarrow p. 238). The secretion of profuse amounts of saliva before vomiting also prevents gastric acid from damaging the enamel on the teeth. Saliva secretion is very dependent on the body water content. A low content results in decreased saliva secretion-the mouth and throat become dry, thereby evoking the sensation of thirst. This is an important mechanism for maintaining the fluid balance $(\rightarrow pp. 168 \text{ and } 184).$

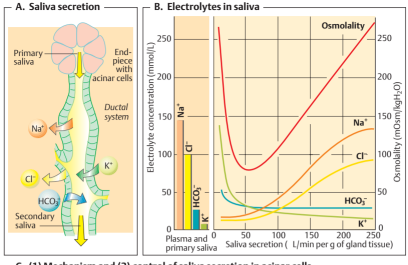
Secretion rate. The rate of saliva secretion varies from 0.1 to $4\,\text{mL/min}$ $(10\text{--}250\,\mu\text{L/min})$ per gram gland tissue), depending on the degree of stimulation $(\rightarrow$ B). This adds up to about 0.5 to 1.5 L per day. At 0.5 mL/min, 95% of this rate is secreted by the parotid gland (serous saliva) and submandibular gland (mucinrich saliva). The rest comes from the sublingual glands and glands in the buccal mucosa.

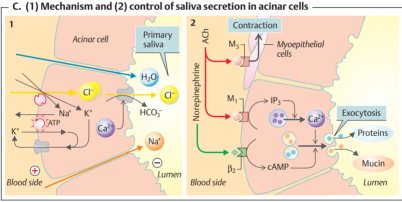
Saliva secretion occurs in *two steps*: The *acini* (*end pieces*) produce **primary saliva** (\rightarrow **A**, **C**) which has an electrolyte composition similar to that of plasma (\rightarrow **B**). Primary saliva secretion in the acinar cells is the result of *transcellular Cl- transport*: Cl⁻ is actively taken up into the cells (secondary active transport) from the blood by means of a Na+-K⁺-2Cl⁻ cotransport carrier and is released into the lumen (together with HCO₃⁻) via anion chan-

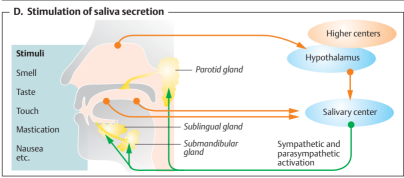
nels, resulting in a lumen-negative transepithelial potential (LNTP) that drives Na+ paracellularly into the lumen. Water also follows passively (osmotic effect). Primary saliva is modified in excretory ducts, yielding secondary saliva. As the saliva passes through the excretory ducts, Na+ and Cl- are reabsorbed and K⁺ and (carbonic anhydrase-dependent) HCO₃⁻ is secreted into the lumen. The saliva becomes hypotonic (far below 100 mOsm/kg $H_2O: \rightarrow B$) because Na⁺ and Cl⁻ reabsorption is greater than K⁺ and HCO₃⁻ secretion and the ducts are relatively impermeable to water $(\rightarrow B)$. If the secretion rate rises to values much higher than 100 μL/(min·g), these processes lag behind and the composition of secondary saliva becomes similar to that of primary saliva $(\rightarrow B)$.

Salivant stimuli. Reflex stimulation of saliva secretion occurs in the larger salivary glands $(\rightarrow \mathbf{D})$. Salivant stimuli include the smell and taste of food, tactile stimulation of the buccal mucosa, mastication and nausea. Conditioned reflexes also play a role. For instance, the routine clattering of dishes when preparing a meal can later elicit a salivant response. Sleep and dehydration inhibit saliva secretion. Saliva secretion is stimulated via the sympathetic and parasympathetic nervous systems $(\rightarrow \mathbf{C2})$:

- Norepinephrine triggers the secretion of highly viscous saliva with a high concentration of mucin via β_2 adrenoreceptors and cAMP. VIP also increases the cAMP concentration of acinar cells.
- Acetylcholine: (a) With the aid of M₁ cholinoceptors and IP₃ (→ pp. 82 and 274), acetylcholine mediates an increase in the cytosolic Ca²+ concentration of acinar cells. This, in turn, increases the conductivity of luminal anion channels, resulting in the production of watery saliva and increased exocytosis of salivary enzymes. (b) With the aid of M₃ cholinergic receptors, ACh mediates the contraction of myoepithelial cells around the acini, leading to emptying of the acini. (c) ACh enhances the production of kallikreins, which cleave bradykinin from plasma kininogen. Bradykinin and VIP (—p. 234) dilate the vessels of the salivary glands. This is necessary because maximum saliva secretion far exceeds resting blood flow.







Deglutition

The upper third of the esophageal wall consists of striated muscle, the rest contains smooth muscle. During the process of swallowing, or deglutition, the tongue pushes a bolus of food into the throat (\rightarrow A1). The nasopharynx is reflexively blocked, (\rightarrow A2), respiration is inhibited, the vocal chords close and the epiglottis seals off the trachea (\rightarrow A3) while the *upper esophageal sphincter* opens (\rightarrow A4). A peristaltic wave forces the bolus into the stomach (\rightarrow A5, B1,2). If the bolus gets stuck, stretching of the affected area triggers a *secondary peristaltic wave*.

The **lower esophageal sphincter** opens at the start of deglutition due to a *vagovagal reflex* (*receptive relaxation*) mediated by VIP- and NO-releasing neurons (\rightarrow **B3**). Otherwise, the lower sphincter remains closed to prevent the reflux of aggressive gastric juices containing pepsin and HCl.

Esophageal motility is usually checked by measuring pressure in the lumen, e.g., during a peristaltic wave (\rightarrow **B1-2**). The resting pressure within the lower sphincter is normally 20–25 mmHg. During receptive relaxation, esophageal pressure drops to match the low pressure in the proximal stomach (\rightarrow **B3**), indicating opening of the sphincter. In achalasia, receptive relaxation fails to occur and food collects in the esophagus.

Pressure in the lower esophageal sphincter is decreased by VIP, CCK, NO, GIP, secretin and progesterone (→ p. 234) and increased by acetylcholine, gastrin and motilin. Increased abdominal pressure (external pressure) also increases sphincter pressure because part of the lower esophageal sphincter is located in the abdominal cavity.

Gastroesophageal reflux. The sporadic reflux of gastric juices into the esophagus occurs fairly often. Reflux can occur while swallowing (lower esophageal sphincter opens for a couple of seconds), due to unanticipated pressure on a full stomach or to transient opening of the sphincter (lasts up to 30 seconds and is part of the eructation reflex). Gastric reflux greatly reduces the pH in the distal esophagus.

Protective mechanisms to prevent damage to the esophageal mucosa after gastroesophageal reflux include 1. **Volume clearance**, i.e., the rapid return of refluxed fluid to the stomach via the esophageal peristaltic reflex. A refluxed volume of 15 mL, for example, remains in the

esophagus for only 5 to 10 s (only a small amount remains). 2. **pH clearance**. The pH of the residual gastric juice left after volume clearance is still low, but is gradually increased during each act of swallowing. In other words, the saliva that is swallowed buffers the residual gastric juice.

Vomiting

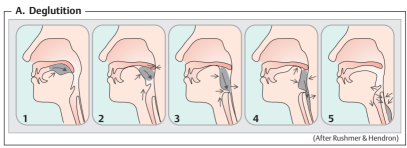
Vomiting mainly serves as a protective reflex but is also an important clinical symptom of conditions such as intracranial bleeding and tumors. The act of vomiting is heralded by nausea, increased salivation and retching (→C). The vomiting center is located in the medulla oblongata within the reticular formation. It is mainly controlled by chemosensors of the area postrema, which is located on the floor of the fourth ventricle; this is called the chemosensory trigger zone (CTZ). The bloodbrain barrier is less tight in the area postrema.

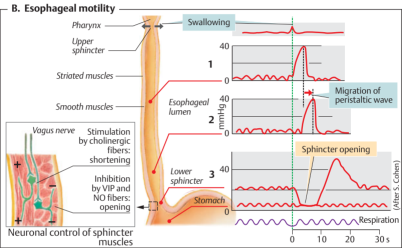
The CTZ is activated by nicotine, other toxins, and dopamine agonists like apomorphine (used as an emetic). Cells of the CTZ have receptors for neurotransmitters responsible for their neuronal control. The vomiting center can also be activated independent of the CTZ, for example, due to abnormal stimulation of the organ of balance (kinesia, motion sickness), overextension of the stomach or intestines, delayed gastric emptying and inflammation of the abdominal organs. Nausea and vomiting often occur during the first trimester of pregnancy (morning sickness) and can exacerbate to hyperemesis gravidarum leading to vomiting-related disorders (see below).

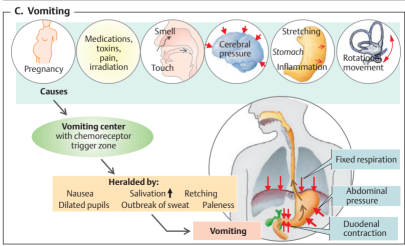
During the act of vomiting, the diaphragm remains in the inspiratory position and the abdominal muscles quickly contract exerting a high pressure on the stomach. Simultaneous contraction of the duodenum blocks the way to the gut; the lower esophageal sphincter then relaxes, resulting in ejection of the stomach contents via the esophagus.

The sequelae of **chronic vomiting** are attributable to reduced food intake (*malnutrition*) and the related loss of gastric juices, swallowed saliva, fluids and intestinal secretions. In addition to *hypovolemia*, nonrespiratory *alkalosis* due to the loss of gastric acid (10–100 mmol H*/L gastric juice) also develops. This is accompanied by *hypokalemia* due to the loss of K* in the vomitus (nutrients, saliva, gastric juices) and urine (hypovolemia-related *hyperaldosteronism*; \rightarrow p. 180 ff.).

238







Stomach Structure and Motility

Structure. The cardia connects the esophagus to the upper stomach (fundus), which merges with the body (cornus) followed by the antrum of the stomach. The lower outlet of the stomach (pylorus) merges with the duodenum $(\rightarrow A)$. Stomach size is dependent on the degree of gastric filling, but this distension is mainly limited to the proximal stomach ($\rightarrow A$. B). The stomach wall has an outer laver of longitudinal muscle fibers (only at curvatures: regulates stomach length), a layer of powerful circular muscle fibers, and an inner layer of oblique muscle fibers. The mucosa of the tubular glands of the fundus and corpus contain chief cells (CC) and parietal cells (PC) (\rightarrow A) that produce the constituents of gastric juice ($\rightarrow p$. 242). The gastric mucosa also contains endocrine cells (that produce gastrin in the antrum, etc.) and mucous neck cells (MNC).

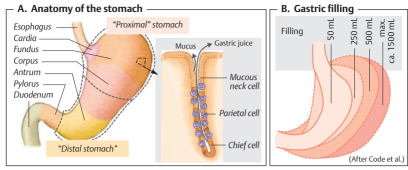
Functional anatomy. The stomach can be divided into a proximal and a distal segment $(\rightarrow A)$. A vagovagal reflex triggered by swallowing a bolus of food causes the lower esophageal sphincter to open $(\rightarrow p.238)$ and the proximal stomach to dilate for a short period (receptive relaxation). This continues when the food has entered the stomach (vagovagal accommodation reflex). As a result. the internal pressure hardly rises in spite of the increased filling. Tonic contraction of the proximal stomach, which mainly serves as a reservoir, slowly propel the gastric contents to the distal stomach. Near its upper border (middle third of the corpus) is a pacemaker zone (see below) from which peristaltic waves of contraction arise due mainly to local stimulation of the stomach wall (in response to reflex stimulation and gastrin; \rightarrow **D1**). The peristaltic waves are strongest in the antrum and spread to the pylorus. The chyme is thereby driven towards the pylorus (\rightarrow C5, 6, 1), then compressed (\rightarrow C2, 3) and propelled back again after the pylorus closes (\rightarrow C3, 4). Thereby, the food is processed, i.e., ground, mixed with gastric juices and digested, and fat is emulsified.

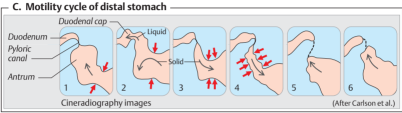
The distal stomach contains **pacemaker cells** (*interstitial Cajal cells*), the membrane potential of which oscillates roughly every 20 s, producing characteristic *slow waves* (→ p. 244). The velocity (0.5–4 cm/s) and amplitude

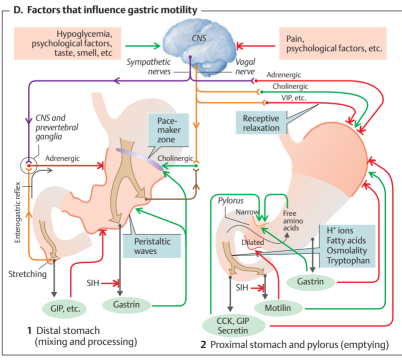
(0.5–4 mV) of the waves increases as they spread to the pylorus. Whether and how often contraction follows such an excitatory wave depends on the sum of all neuronal and hormonal influences. Gastrin increases the response frequency and the pacemaker rate. Other hormones like GIP inhibit this motility directly, whereas somatostatin (SIH) does so indirectly by inhibiting the release of GRP (\rightarrow D1 and p.234).

Gastric emptying. Solid food remains in the stomach until it has been broken down into small particles (diameter of < 1 mm) and suspended in chyme. The chyme then passes to the duodenum. The time required for 50% of the ingested volume to leave the stomach varies, e.g., 10-20 min for water and 1-4 hours for solids (carbohydrates < proteins < fats). Emptying is mainly dependent on the tone of the proximal stomach and pylorus. Motilin stimulates emptying of the stomach (tone of proximal stomach rises, pylorus dilates), whereas decreases in the pH or osmolality of chyme or increases in the amount of long-chain free fatty acids or (aromatic) amino acids inhibit gastric emptying. Chemosensitive enterocytes and brush cells of the small intestinal mucosa. enterogastric reflexes and certain hormones (CCK, GIP, secretin and gastrin; \rightarrow p. 234) mediate these regulatory activities ($\rightarrow D2$). The pylorus is usually slightly open during the process (free flow of "finished" chyme). It contracts only 1) at the end of "antral systole" (see above) in order to retain solid food and 2) when the duodenum contracts in order to prevent the reflux of harmful bile salts. If such reflex does occur, refluxed free amino acids not normally present in the stomach elicit reflex closure of the pylorus (\rightarrow **D2**).

Indigestible substances (bone, fiber, foreign bodies) do not leave the stomach during the digestive phase. Special contraction waves called *migrating motor complexes* (MMC) pass through the stomach and small intestine roughly every 1.5 hours during the ensuing interdigestive phase, as determined by an intrinsic "biological clock." These peristaltic waves transport indigestible substances from the stomach and bacteria from the small intestine to the large intestine. This "clearing phase" is controlled by *motilin*.







Gastric Juice

The *tubular glands* of the gastric fundus and corpus secrete 3–4 L of gastric juice each day. *Pepsinogens* and *lipases* are released by chief cells and *HCl* and *intrinsic factor* (\rightarrow p. 260) by parietal cells. *Mucins* and HCO_3^- are released by mucous neck cells and other mucous cells on the surface of the gastric mucosa.

Pepsins function as endopeptidases in protein digestion. They are split from pepsinogens exocytosed from chief cells in the glandular and gastric lumen at a pH of < 6. Acetylcholine (ACh), released locally in response to H $^+$ (and thus indirectly also to gastrin) is the chief activator of this reaction.

Gastric acid. The pH of the gastric juice drops to ca. 0.8 during peak HCl secretion. Swallowed food buffers it to a pH of 1.8–4, which is optimal for most pepsins and gastric lipases. The low pH contributes to the *denaturation* of dietary proteins and has a *bactericidal effect*.

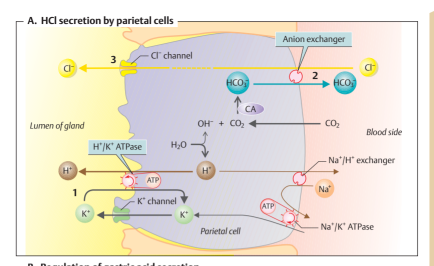
HCl secretion (\rightarrow **A**). The H^+/K^+ -ATPase in the luminal membrane of parietal cells drives H+ ions into the glandular lumen in exchange for K^+ (primary active transport, \rightarrow **A1** and p. 26), thereby raising the H⁺ conc. in the lumen by a factor of ca. 10⁷. K⁺ taken up in the process circulates back to the lumen via luminal K+ channels. For every H⁺ ion secreted, one HCO₃⁻ ion leaves the blood side of the cell and is exchanged for a Cl- ion via an anion antiporter $(\rightarrow A2)$. (The HCO₃⁻ ions are obtained from CO₂ + OH-, a reaction catalyzed by carbonic anhydrase, CA). This results in the intracellular accumulation of Cl-ions, which diffuse out of the cell to the lumen via Cl^- channels ($\rightarrow A3$). Thus, one Cl- ion reaches the lumen for each H+ ion secreted.

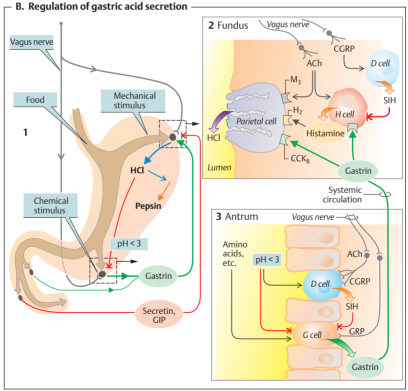
The activation of parietal cells (see below) leads to the opening of canaliculi, which extend deep into the cell from the lumen of the gland (\rightarrow B). The canaliculi are equipped with a brush border that greatly increases the luminal surface area which is densely packed with membrane-bound H*/K* ATPase molecules. This permits to increase the secretion of H* ions from 2 mmol/hour at rest to over 20 mmol/hour during digestion.

Gastric acid secretion is stimulated in phases by neural, local gastric and intestinal factors (\rightarrow **B**). Food intake leads to reflex secretion of gastric juices, but deficient levels of glucose in the brain can also trigger the reflex. The optic, gustatory and olfactory nerves are the afferents for this partly conditioned reflex $(\rightarrow p. 236)$, and efferent impulses flow via the vagus nerve. ACh directly activates parietal cells in the fundus (M_3 cholinoceptors \rightarrow **B2**). GRP (gastrin-releasing peptide) released by neurons stimulates gastrin secretion from G cells in the antrum (\rightarrow B3). Gastrin released in to the systemic circulation in turn activates the parietal cells via CCKB receptors (= gastrin receptors). The glands in the fundus contain H (histamine) cells or ECL cells (enterochromaffin-like cells), which are activated by gastrin (CCK_B receptors) as well as by ACh and β_3 adrenergic substances (\rightarrow **B2**). The cells release histamine, which has a paracrine effect on neighboring parietal cells (H2 receptor). Local gastric and intestinal factors also influence gastric acid secretion because chyme in the antrum and duodenum stimulates the secretion of gastrin (\rightarrow **B1** and p. 235, A).

Factors that inhibit gastric juice secretion: (a) A pH of < 3.0 in the antral lumen inhibits G cells (negative feedback, \rightarrow B1, 3) and activates antral D cells, which secrete SIH (\rightarrow p. 234), which in turn has a paracrine effect. SIH inhibits H cells in the fundus as well as G cells in the antrum (\rightarrow B2, 3). CGRP released by neurons (\rightarrow p. 234) activates D cells in the antrum and fundus, (\rightarrow B2, 3). (c) Secretin and GIP released from the small intestine have a retrograde effect on gastric juice secretion (\rightarrow B1). This adjusts the composition of chyme from the stomach to the needs of the small intestine.

Protection of the gastric mucosa from destructive gastric juices is chiefly provided by (a) a *layer of mucus* and (b) HCO_3^- secretion by the underlying mucous cells of the gastric mucosa. HCO_3^- diffuses through the layer of mucus and buffers the acid that diffuses into it from the lumen. *Prostaglandins* PGE₂ and PGI₂ promote the secretion of HCO_3^- . Anti-inflammatory drugs that inhibit cyclooxygenase 1 and thus prostaglandin production (\rightarrow p. 269) impair this mucosal protection and can result in ulcer development.





Small Intestinal Function

The main function of the small intestine (SI) is to finish digesting the food and to absorb the accumulated breakdown products as well as water, electrolytes and vitamins.

Structure. The SI of live human subjects is about 2 m in length. It arises from the pylorus as the duodenum and continues as the *jejunum*, and ends as the *ileum*, which merges into the large intestine. From outside inward, the SI consists of an outer serous coat (tunica serosa, \rightarrow **A1**), a layer of longitudinal muscle fibers $(\rightarrow A2)$, the myenteric plexus (Auerbach's plexus. \rightarrow A3), a layer of circular muscle fibers (\rightarrow A4), the submucous plexus (Meissner's plexus, \rightarrow **A5**) and a mucous layer (tunica mucosa, \rightarrow A6), which is covered by epithelial cells (→ A13-15). The SI is supplied with blood vessels ($\rightarrow A8$), lymph vessels ($\rightarrow A$ **9**), and nerves $(\rightarrow A10)$ via the mesentery $(\rightarrow A7)$. The surface area of the epithelial-luminal interface is roughly 300-1600 times larger (> 100 m²) than that of a smooth cylindrical pipe because of the Kerckring's folds (\rightarrow A11), the intestinal villi (\rightarrow A12), and the enterocytic microvilli, or the brush border (\rightarrow **A13**).

Ultrastructure and function. Goblet cells (→ A15) are interspersed between the resorbing enterocytes (\rightarrow A14). The mucus secreted by goblet cells acts as a protective coat and lubricant. Intestinal glands (crypts of Lieberkühn, \rightarrow A16) located at the bases of the villi contain (a) undifferentiated and mitotic cells that differentiate into villous cells (see below), (b) mucous cells, (c) endocrine and paracrine cells that receive information about the composition of chyme from chemosensor cells, and (d) immune cells (\rightarrow p. 232). The chyme composition triggers the secretion of endocrine hormones and of paracrine mediators (\rightarrow p. 234). The tubuloacinar duodenal glands (Brunner's glands), located deep in the intestinal wall (tela submucosa) secrete a HCO3--rich fluid containing urogastrone (human epidermal growth factor), an important stimulator of epithelial cell proliferation.

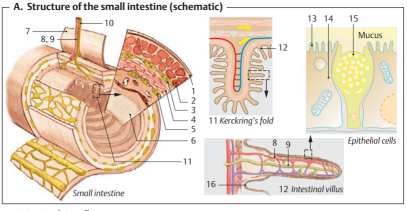
Cell replacement. The tips of the villi are continually shed and replaced by new cells from the crypts of Lieberkühn. Thereby, the entire SI epithelium is renewed every 3–6 days. The dead cells disintegrate in the lumen, thereby releasing enzymes, stored iron, etc.

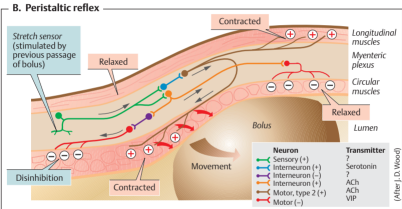
Intestinal motility is autonomously regulated by the enteric nervous system, but is influenced by hormones and external innervation (→ p. 234). Local pendular movements (by longitudinal muscles) and segmentation (contraction/relaxation of circular muscle fibers) of the SI serve to mix the intestinal contents and bring them into contact with the mucosa. This is enhanced by movement of the intestinal villi (lamina muscularis mucosae). Reflex peristaltic waves (30–130 cm/min) propel the intestinal contents towards the rectum at a rate of ca. 1 cm/min. These waves are especially strong during the interdigestive phase (→ p. 240).

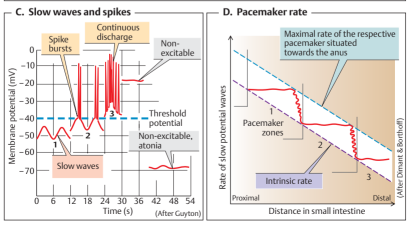
Peristaltic reflex. Stretching of the intestinal wall during the passage of a bolus $(\rightarrow B)$ triggers a reflex that constricts the lumen behind the bolus and dilates that ahead of it. Controlled by interneurons, cholinergic type 2 motoneurons with prolonged excitation simultaneously activate circular muscle fibers behind the bolus and longitudinal musculature in front of it. At the same time the circular muscle fibers in front of the bolus are inhibited (accommodation) while those behind it are disinhibited ($\rightarrow B$ and p. 234).

Pacemakers. The intestine also contains pacemaker cells (*interstitial Cajal cells*). The membrane potential of these cells oscillates between 10 and 20 mV every 3–15 min, producing *slow waves* (\rightarrow C1). Their amplitude can rise (less negative potential) or fall in response to neural, endocrine or paracrine stimuli. A series of action potentials (spike bursts) are fired once the membrane potential rises above a certain threshold (ca. –40 mV) (\rightarrow C2). *Muscle spasms* occur if the trough of the wave also rises above the threshold potential (\rightarrow C3).

Impulse conduction. The spike bursts are conducted to myocytes via *gap junctions* (\rightarrow p. 70). The myocytes then contract rhythmically at the same frequency (or slower). Conduction in the direction of the anus dwindles after a certain distance (\rightarrow D, pacemaker zone), so more distal cells (with a lower intrinsic rate) must assume the pacemaker function. Hence, peristaltic waves of the small intestine only move in the anal direction.







Pancreas

The *exocrine part* of the pancreas secretes 1–2 L of pancreatic juice into the duodenum each day. The pancreatic juice contains *bicarbonate* (HCO₃⁻), which neutralizes (pH 7–8) HCl-rich chyme from the stomach, and mostly inactive precursors of *digestive enzymes* that break down proteins, fats, carbohydrates and other substances in the small intestine.

Pancreatic secretions are similar to saliva in that they are produced in two stages: (1) Cl⁻ is secreted in the acini by active secondary transport, followed by passive transport of Na⁺ and water (\rightarrow p. 237 C1). The electrolyte composition of these primary secretions corresponds to that of plasma (\rightarrow A1 and A2). Primary pancreatic secretions also contain digestive proenzymes and other proteins (exocytosis; \rightarrow p. 30). (2) HCO_3^- is added to the primary secretions (in exchange for Cl-) in the secretory ducts: Na⁺ and water follow by passive transport. As a result, the HCO₃⁻ concentration of pancreatic juice rises to over 100 mmol/L, while the Cl- concentration falls $(\rightarrow A3)$. Unlike saliva $(\rightarrow p. 237 B)$, the osmolality and Na⁺/K⁺ concentrations of the pancreatic juice remain constant relative to plasma (→ A1 and A2). Most of the pancreatic juice is secreted during the digestive phase ($\rightarrow A3$).

 HCO_3^- is secreted from the luminal membrane of the ductules via an anion exchanger that simultaneously reabsorbs Cl^- from the lumen (\rightarrow B1). Cl^- returns to the lumen via a Cl^- channel, which is more frequently opened by secretin to ensure that the amount of HCO_3^- secreted is not limited by the availability of Cl^- (\rightarrow B2). In cystic fibrosis (mucoviscidosis), impairment of this CFTR channel (cystic fibrosis transmembrane conductance regulator) leads to severe disturbances of pancreatic function. The HCO_3^- involved is the product of the $CO_2 + OH^-$ reaction catalyzed by carbonic anhydrase (CA). For each HCO_3^- molecule secreted, one H^+ ion leaves the cell on the blood side via an Na^+/H^+ exchanger (\rightarrow B3).

Pancreatic juice **secretion is controlled** by cholinergic (vagal) and hormonal mechanisms (CCK, secretin). Vagal stimulation seems to be enhanced by CCK_A receptors in cholinergic fibers of the acini (\rightarrow **A2,3, B, C** and p. 234). Fat in the chyme stimulates the release of CCK, which, in turn, increases the *(pro)enzyme con-*

tent of the pancreatic juice (\rightarrow C). Trypsin in the small intestinal lumen deactivates CCK release via a feedback loop (\rightarrow D). Secretin increases HCO₃⁻ and water secretion by the ductules. CCK and acetylcholine (ACh) potentiate this effect by raising the cytosolic Ca²⁺ concentration. Secretin and CCK also affect the pancreatic enzymes.

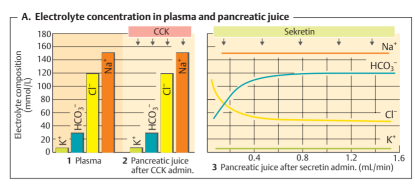
Pancreatic enzymes are essential for digestion. They have a pH optimum of 7–8. Insufficient HCO₃⁻ secretion (e.g., in cystic fibrosis) results in inadequate neutralization of chyme and therefore in impaired digestion.

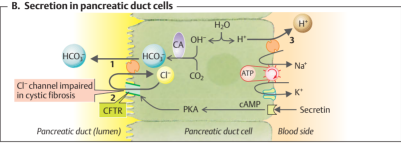
Proteolysis is catalyzed by proteases, which are secreted in their inactive form, i.e., as proenzymes: trypsinogen 1-3, chymotrypsinogen A and B, proelastase 1 and 2 and procarboxypeptidase A1, A2, B1 and B2. They are not activated until they reach the intestine, where an enteropeptidase first converts trypsinogen to trypsin $(\rightarrow \mathbf{D})$, which in turn converts chymotrypsinogen into active chymotrypsin. Trypsin also activates many other pancreatic proenzymes including proelastases and procarboxypeptidases. Pathological activation of the proenzymes within the pancreas causes the organ to digest itself (acute pancreatic necrosis). Trypsins, chymotrypsins and elastases are endoproteases, i.e., they split certain peptide bonds within protein chains. Carboxypeptidases A and B are exopeptidases, i.e., they split amino acids off the carboxyl end of the chain.

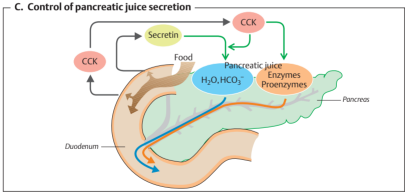
Carbohydrate catabolism. α -Amylase is secreted in active form and splits starch and glycogen into maltose, maltotriose and α -limit dextrin. These products are further digested by enzymes of the intestinal epithelium (\rightarrow p. 259).

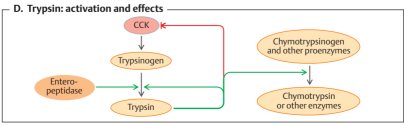
Lipolysis. Pancreatic lipase (see p. 252ff.) is the most important enzyme for lipolysis. It is secreted in its active form and breaks triacylglycerol to 2-monoacylglycerol and free fatty acids. Pancreatic lipase activity depends on the presence of *colipases*, generated from pro-colipases in pancreatic secretions (with the aid of trypsin). Bile salts are also necessary for fat digestion (→ p. 248).

Other important pancreatic enzymes include (pro-) phospholipase A₂, RNases, DNases, and a carboxylesterase.









Bile

Bile components. Bile contains electrolytes, bile salts (bile acids), cholesterol, lecithin (phosphatidylcholine), bilirubin diglucuronide, steroid hormones, medications etc. $(\rightarrow A)$. Bile salts are essential for fat digestion. Most of the other components of bile leave the body via the feces (excretory function of the liver \rightarrow p. 250).

Bile formation. Hepatocytes secrete ca. 0.7 L/day of bile into biliary canaliculi (\rightarrow A), the fine canals formed by the cell membranes of adjacent of hepatocytes. The sinusoidal and canalicular membranes of the hepatocytes contain numerous carriers that absorb bile components from the blood and secrete them into the canaliculi, resp.

Bile salts (BS). The liver synthesizes *cholate* and *chenodeoxycholate* (primary bile salts) from *cholesterol*. The intestinal bacteria convert some of them into secondary bile salts such as *deoxycholate* and *lithocholate*. Bile salts are conjugated with taurine or glycine in the liver and are secreted into the bile in this form $(\rightarrow A)$. This conjugation is essential for micelle formation in the bile and gut.

Hepatic bile salt carriers. Conjugated bile salts in sinusoidal blood are actively taken up by NTCP (Na⁺ taurocholate cotransporting polypeptide; secondary active transport), and transported against a steep concentration gradient into the canaliculi (primary active transport) by the ATP-dependent carrier hBSEP (human bile salt export pump), also referred to as cBAT (canalicular bile acid transporter).

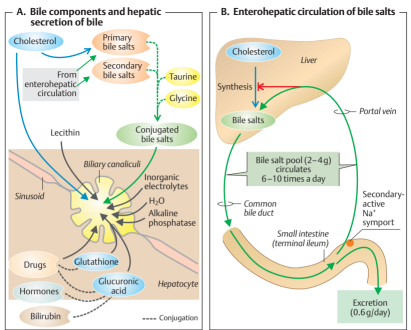
Enterohepatic circulation of BS. Unconjugated bile salts are immediately reabsorbed from the bile ducts (cholehepatic circulation). Conjugated bile salts enter the duodenum and are reabsorbed from the terminal ileum by the Natsymport carrier ISBT (=ileal sodium bile acid cotransporter) and circulated back to the liver (enterohepatic circulation; \rightarrow B) once they have been used for fat digestion (\rightarrow p. 252). The total bile pool (2–4g) recirculates about 6–10 times a day, depending on the fat content of the diet. Ca. 20–30 g of bile salts are required for daily fat absorption.

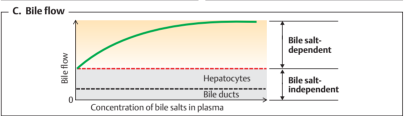
Choleresis. Enterohepatic circulation raises the bile salt concentration in the portal vein to a high level during the digestive phase. This (a) inhibits the hepatic synthesis of bile salts (cholesterol- 7α -hydroxylase; negative feedback; \rightarrow **B**) and (b) stimulates the secretion of bile salts into the biliary canaliculi. The latter effect increases the bile flow due to osmotic water movement, i.e., causes bile salt-dependent choleresis (\rightarrow **C**). Bile salt-independent choleresis is, caused by secretion of other bile components into the canaliculi as well as of HCO_3^- (in exchange for CI^-) and H_2O into the bile ducts (\rightarrow **C**). The latter form is increased by the vagus nerve and secretin.

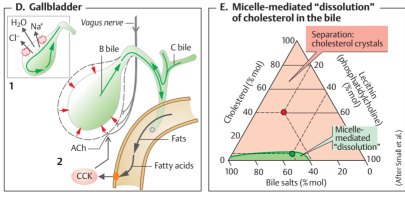
Gallbladder. When the sphincter of Oddi between the common bile duct and duodenum is closed, *hepatic bile* (C *bile*) is diverted to the gallbladder, where it is concentrated (1:10) and stored (\rightarrow **D**). The gallbladder epithelium reabsorbs Na*, Cl- and water (\rightarrow **D**1) from the stored bile, thereby greatly raising the concentration of specific bile components (bile salts, bilirubin-di-glucuronide, cholesterol, phosphatidylcholine, etc.). If bile is used for fat digestion (or if a peristaltic wave occurs in the interdigestive phase, \rightarrow p. 240), the gallbladder contracts and its contents are mixed in portions with the duodenal chyme (\rightarrow **D2**).

Cholesterol in the bile is transported inside *micelles* formed by aggregation of cholesterol with lecithin and bile salts. A change in the ratio of these three substances in favor of cholesterol (\rightarrow **E**) leads to the precipitation of cholesterol crystals responsible for gallstone development in the highly concentrated *gallbladder bile* (B *bile*). The red and green dots in **E** show the effects of two different ratios.

Gallbladder contraction is *triggered* by CCK (\rightarrow p. 234), which binds to CCK_A receptors, and the neuronal plexus of the gallbladder wall, which is innervated by preganglionic parasympathetic fibers of the vagus nerve (\rightarrow **D2**). CGRP (\rightarrow p. 234) and substance P (\rightarrow p. 86) released by sensory fibers appear to stimulate the gallbladder musculature indirectly by increasing acetylcholine release. The sympathetic nervous system *inhibits* gallbladder contractions via α_2 adrenoreceptors located on cholinergic fiber terminals. As *cholagogues*, fatty acids and products of protein digestion (\rightarrow p. 234) as well as egg yolk and MgSO₄ effectively stimulate CCK secretion.







Excretory Liver Function—Bilirubin

The liver detoxifies and excretes many mostly lipophilic substances, which are generated during metabolism (e.g., bilirubin or steroid hormones) or come from the intestinal tract (e.g., the antibiotic chloramphenicol). However, this requires prior biotransformation of the substances. In the first step of the process, reactive OH, NH2 or COOH groups are enzymatically added (e.g., by monooxygenases) to the hydrophobic substances. In the second step, the substances are conjugated with glucuronic acid, acetate, glutathione, glycine, sulfates, etc. The conjugates are now water-soluble and can be either further processed in the kidneys and excreted in the urine, or secreted into bile by liver cells and excreted in the feces. Glutathione conjugates, for example, are further processed in the kidney excreted as mercapturic acids in the urine.

Carriers. The canalicular membrane of hepatocytes contains various carriers, most of which are directly fueled by ATP (see also p. 248). The principal carriers are: MDR1 (multidrug resistance protein 1) for relatively hydrophobic, mainly cationic metabolites, MDR3 for phosphatidylcholine (\rightarrow p. 248), and cMOAT (canalicular multispecific organic anion transporter = multidrug resistance protein MRP2) for conjugates (formed with glutathione, glucuronic acid or sulfate) and many other organic anions.

Bilirubin sources and conjugation. Ca. 85% of all bilirubin originates from the hemoglobin in erythrocytes; the rest is produced by other hemoproteins like cytochrome ($\rightarrow A$ and B). When degraded, the globulin and iron components (\rightarrow p. 90) are cleaved from hemoglobin. Via intermediate steps, biliverdin and finally bilirubin, the yellow bile pigment, are then formed from the porphyrin residue. Each gram of hemoglobin yields ca. 35 mg of bilirubin. Free unconjugated bilirubin ("indirect" bilirubin) is poorly soluble in water, yet lipidsoluble and toxic. It is therefore complexed with albumin when present in the blood (2 mol bilirubin: 1 mol albumin), but not when absorbed by hepatocytes ($\rightarrow A$). Bilirubin is conjugated (catalyzed by glucuronyltransferase) with 2 molecules of UDP-glucuronate (synthesized from glucose, ATP and UTP) in the liver cells yielding bilirubin diglucuronide ("direct"

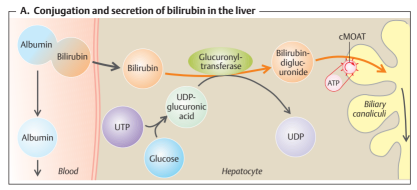
bilirubin). It is a water-soluble substance secreted into the biliary canaliculi by primary active transport mechanisms (cMOAT, see above).

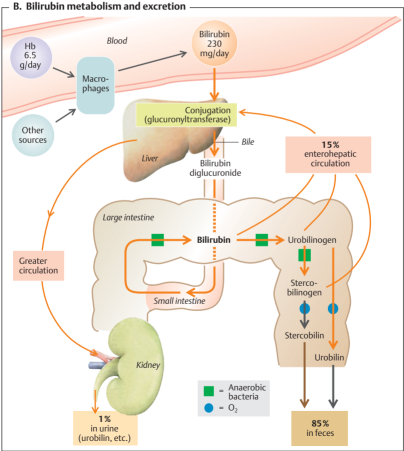
Bilirubin excretion. 200–250 mg of bilirubin is excreted in the bile each day. Ca. 90% of it is excreted in the feces. In the gut, bacteria break bilirubin down into the colorless compound, stercobilinogen (\rightarrow **B**). It is partly oxidized into stercobilin, the brown compound that colors the stools. About 10% of all bilirubin diglucuronide is deconjugated by intestinal bacteria and returned to the liver in this lipophilic form (partly as stercobilinogen) via enterohepatic circulation. A small portion (ca. 1%) reaches the systemic circulation and is excreted by the kidneys as *urobilinogen* = stercobilinogen (see below) (\rightarrow **B**). The renal excretion rate increases when the liver is damaged.

Jaundice. The plasma bilirubin concentration normally does not exceed 17 µmol/L (= 1 mg/dL). Concentrations higher than 30 µmol/L (1.8 mg/dL) lead to yellowish discoloration of the sclera and skin, resulting in jaundice (icterus). Types of jaundice:

- 1. **Prehepatic jaundice**. When excessive amounts of bilirubin are formed, for example, due to increased hemolysis, the liver can no longer cope with the higher load unless the plasma bilirubin concentration rises. Thus, *unconjugated (indirect) bilirubin* is mainly elevated in these patients.
- 2. Intrahepatic jaundice. The main causes are (a) liver cell damage due to toxins (Amanita) or infections (viral hepatitis) resulting in the impairment of bilirubin transport and conjugation; (b) deficiency or absence of the glucuronyltransferase system in the newborn (Crigler–Najjar syndrome); (c) inhibition of glucuronyltransferase, e.g., by steroids; (d) impaired secretion of bilirubin into the biliary canaliculi due to a congenital defect (Dubin–Johnson syndrome) or other reasons (e.g., drugs, steroid hormones).
- 3. **Posthepatic jaundice**: Impairment of the flow of bile occurs due to an obstruction (e.g., stone or tumor) in the bile ducts, usually accompanied by elevated serum concentrations of *conjugated (direct) bilirubin* and alkaline phosphatase—both of which are normal components of bile.

Types 2a, 2d and 3 jaundice are associated with increased urinary concentrations of conjugated bilirubin, leading to brownish discoloration of the urine. In type 3 jaundice, the stools are gray due to the lack of bilirubin in the intestine and the resulting absence of sterrobilin formation.





Lipid Digestion

The average **intake of fats** (butter, oil, margarine, milk, meat, sausages, eggs, nuts etc.) is roughly 60–100 g/day, but there is a wide range of individual variation (10–250 g/day). Most fats in the diet (90%) are neutral fats or *triacylglycerols* (triglycerides). The rest are phospholipids, cholesterol esters, and fatsoluble vitamins (vitamins A, D, E and K). Over 95% of the lipids are normally absorbed in the small intestine.

Lipid digestion (\rightarrow **A**). Lipids are poorly soluble in water, so special mechanisms are required for their digestion in the watery environment of the gastrointestinal tract and for their subsequent absorption and transport in plasma (\rightarrow p. 254). Although small quantities of undegraded triacylglycerol can be absorbed, dietary fats must be hydrolyzed by enzymes before they can be efficiently absorbed. Optimal enzymatic activity requires the prior mechanical *emulsification* of fats (mainly in the distal stomach, \rightarrow p. 240) because emulsified *lipid droplets* (1–2 μ m; \rightarrow **B1**) provide a much larger surface (relative to the mass of fat) for lipases.

Lipases, the fat digesting enzymes, originate from the lingual glands, gastric fundus (chief and mucous neck cells) and pancreas ($\rightarrow A$ and p. 246). About 10-30% of dietary fat intake is hydrolyzed in the stomach, while the remaining 70-90% is broken down in the duodenum and upper jejunum. Lingual and gastric lipases have an acid pH optimum, whereas pancreatic lipase has a pH optimum of 7-8. Lipases become active at the fat/oil and water interface $(\rightarrow B)$. Pancreatic lipase (triacylglycerol hydrolase) develops its lipolytic activity (max. 140 g fat/min) in the presence of colipase and Ca²⁺. Pro-colipase in pancreatic juice yields colipase after being activated by trypsin. In most cases, the pancreatic lipases split triacylglycerol (TG) at the 1st and 3rd ester bond. This process requires the addition of water and yields free fatty acids (FFA) and 2-monoacylglycerol.

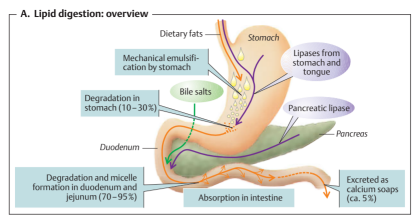
A **viscous-isotropic phase** with aqueous and hydrophobic zones then forms around the enzyme (\rightarrow **B2**). Ca^{2+} excesses or monoacylglycerol deficiencies result in the conversion of the fatty acids into calcium soaps, which are later excreted.

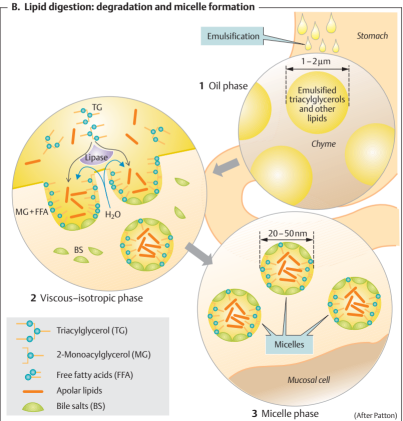
Phospholipase A₂ (from pro-phospholipase A₂ in pancreatic juice—activated by trypsin) cleaves the 2nd ester bond of the phospholipids (mainly phosphatidylcholine = lecithin) contained in micelles. The presence of *bile salts* and Ca^{2+} is required for this reaction.

An unspecific carboxylesterase (= unspecific lipase = cholesterol ester hydrolase) in pancreatic secretions also acts on cholesterol esters on micelles as well as all three ester bonds of TG and esters of vitamins, A, D and E.

This lipase is also present in human breast **milk** (but not cow's milk), so breast-fed infants receive the digestive enzyme required to break down milk fat along with the milk. Since the enzyme is heat-sensitive, pasteurization of human milk significantly reduces the infant's ability to digest milk fat to a great extent.

2-Monoacylglycerols, long-chain free fatty acids and other lipids aggregate with bile salts $(\rightarrow p.248)$ to spontaneously form micelles in the small intestine (\rightarrow **B3**). (Since short-chain fatty acids are relatively polar, they can be absorbed directly and do not require bile salts or micelles). The micelles are only about 20-50 nm in diameter, and their surface-tovolume ratio is roughly 50 times larger than that of the lipid droplets in emulsions. They facilitate close contact between the products of fat digestion and the wall of the small intestine and are therefore essential for lipid absorption. The polar side of the substances involved (mainly conjugated bile salts, 2-monoacylglycerol and phospholipids) faces the watery environment, and the non-polar side faces the interior of the micelle. Totally apolar lipids (e.g., cholesterol esters, fat-soluble vitamins and lipophilic poisons) are located inside the micelles. Thus, the apolar lipids remain in the lipophilic milieu (hydrocarbon continuum) during all these processes until they reach the lipophilic brush border membrane of the epithelium. They are then absorbed by the mucosa cells via dissolution in the membrane or by a passive transport mechanism (e.g., carriers in the case of free fatty acids). Although fat absorption is completed by the time the chyme reaches the end of the jejunum, the bile salts released from micelles are only absorbed in the terminal ileum and then recycled (enterohepatic circulation; \rightarrow p. 249 B).





Lipid Distribution and Storage

Lipids in the blood are transported in lipo**proteins, LP**s $(\rightarrow A)$, which are molecular aggregates (microemulsions) with a core of very hydrophobic lipids such as triacylglycerols (TG) and cholesterol esters (CHO-esters) surrounded by a layer of amphipathic lipids (phospholipids, cholesterol), LPs also contain several types of proteins, called apolipoproteins. LPs are differentiated according to their size, density, lipid composition, site of synthesis, and their apolipoprotein content. Apolipoproteins (Apo) function as structural elements of LPs (e.g. ApoAII and ApoB48), ligands (ApoB100, ApoE, etc.) for LP receptors on the membranes of LP target cells, and as enzyme activators (e.g. ApoAI and ApoCII).

Chylomicrons transport lipids (mainly *triacylglycerol*, *TG*) from the gut to the periphery (via intestinal lymph and systemic circulation; \rightarrow **D**), where their ApoCII activates endothelial lipoprotein lipase (LPL), which *cleaves FFA* from TG. The FFA are mainly absorbed by myocytes and fat cells (\rightarrow **D**). With the aid of ApoE, the *chylomicron remnants* deliver the rest of their TG, cholesterol and cholesterol ester load to the hepatocytes by receptor-mediated endocytosis (\rightarrow **B**, **D**).

Cholesterol (CHO) and the TG imported from the gut and newly synthesized in the liver are exported inside **VLDL** (very low density lipoproteins) from the liver to the periphery, where they by means of their ApoCII also activate **LPL**, resulting in the *release of FFA* (\rightarrow **D**). This results in the loss of ApoCII and exposure of ApoE. VLDL remnants or **IDL** (intermediatedensity lipoproteins) remain. Ca. 50% of the IDL returns to the liver (mainly bound by its ApoE on LDL receptors; see below) and is reprocessed and exported from the liver as VLDL (\rightarrow **B**).

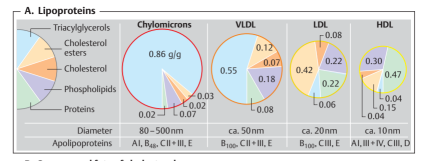
The other 50% of the IDL is converted to **LDL** (low density lipoprotein) after coming in contact with hepatic lipase (resulting in loss of ApoE and exposure of ApoB100). Two-thirds of the LDLs deliver their CHO and CHO-esters to the liver, the other third transfers its CHO to extrahepatic tissue (\rightarrow **B**). Binding of ApoB100 to **LDL receptors** is essential for both processes (see below).

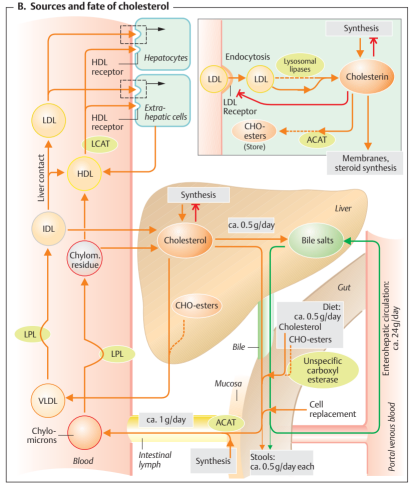
High-density lipoproteins (HDL) exchange certain apoproteins with chylomicrons and VLDL and absorb superfluous CHO from the extrahepatic cells and blood (→ B). With their ApoAl, they activate the plasma enzyme LCAT (lecithin–cholesterol acyltransferase), which is responsible for the partial esterification of CHO. HDL also deliver cholesterol and CHOesters to the liver and steroid hormone-producing glands with HDL receptors (ovaries, testes, adrenal cortex).

Triacylglycerol (TG)

Dietary TGs are broken down into free fatty acids (FFA) and 2-monoacylglycerol (MG) in the gastrointestinal tract ($\rightarrow \mathbf{C}$ and p. 252). Since short-chain FFAs are water-soluble, they can be absorbed and transported to the liver via the portal vein, Long-chain FFAs and 2-monoacylglycerols are not soluble in water. They are re-synthesized to TG in the mucosa cells $(\rightarrow \mathbb{C})$. (The FFAs needed for TG synthesis are carried by FFA-binding proteins from the cell membrane to their site of synthesis, i.e., the smooth endoplasmic reticulum.) Since TGs are not soluble in water, they are subsequently loaded onto chylomicrons, which are exocytosed into the extracellular fluid, then passed on to the intestinal lymph (thereby by-passing the liver), from which they finally reach the greater circulation (\rightarrow **C**, **D**). (Plasma becomes cloudy for about 20-30 minutes after a fatty meal due to its chylomicron content). The liver also synthesizes TGs, thereby taking the required FFAs from the plasma or synthesizing them from glucose. Hepatic TGs are loaded onto **VLDL** (see above) and subsequently secreted into the plasma ($\rightarrow \mathbf{D}$). Since the export capacity of this mechanism is limited, an excess of FFA or glucose ($\rightarrow \mathbf{D}$) can result in the accumulation of TGs in the liver (fatty liver).

Free fatty acids (FFAs) are high-energy substrates used for energy metabolism (\rightarrow p. 228). Fatty acids circulating in the blood are mainly transported in the form of TG (in *lipoproteins*) whereas plasma FFA are *complexed with albumin*. Fatty acids are removed from TGs of chylomicrons and VLDL by lipoprotein lipase (LPL) localized on the luminal surface of the capillary endothelium of many organs (mainly in fat tissue and muscles) (\rightarrow D). ApoCII on the





surface of TGs and VLDL activates LPL. The **insulin** secreted after a meal *induces LPL* (\rightarrow **D**), which promotes the rapid degradation of reabsorbed dietary TGs. LPL is also activated by *heparin* (from endothelial tissue, mast cells, etc.), which helps to eliminate the chylomicrons in cloudy plasma; it therefore is called a *clearance factor*. Albumin-complexed FFAs in plasma are mainly transported to the following **target sites** (\rightarrow **D**):

- Cardiac muscle, skeletal muscle, kidneys and other organs, where they are oxidized to CO_2 and H_2O in the mitochondria (β oxidation) and used as a **source of energy**.
- ♦ Fat cells (\rightarrow **D**), which either **store** the FFAs or use them to synthesize TG. When energy requirements increase or intake decreases, the FFAs are cleaved from triacylglycerol in the fat cells (lipolysis) and transported to the area where they are needed (\rightarrow **D**). Lipolysis is stimulated by epinephrine, glucagon and cortisol and inhibited by insulin (\rightarrow p. 282ff.).
- The *liver*, where the FFAs are oxidized or used to synthesize TG.

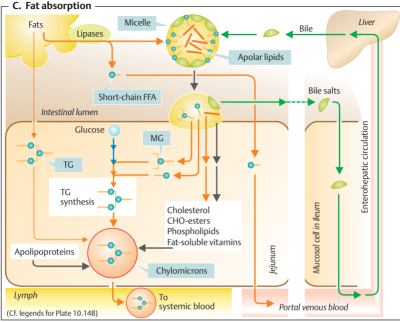
Cholesterol (CHO)

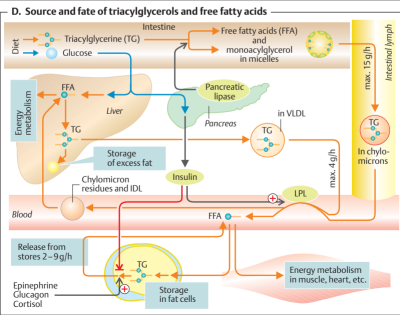
Cholesterol esters (CHO-esters), like TGs, are apolar lipids. In the watery milieu of the body, they can only be transported when incorporated in lipoproteins (or bound to proteins) and can be used for metabolism only after they have been converted to CHO, which is more polar (\rightarrow B). CHO-esters serve as *stores* and in some cases the transported form of CHO. CHO-esters are present in all lipoproteins, but are most abundant (42%) in LDL (\rightarrow A).

Cholesterol is an important constituent of cell membranes (\rightarrow p. 14). Moreover, it is a precursor for *bile salts* (\rightarrow B and p. 248), *vitamin D* (\rightarrow p. 292), and *steroid hormones* (\rightarrow p. 294ff.). Each day ca. 0.6 g of CHO is lost in the feces (reduced to *coprosterol*) and sloughed off skin. The bile salt loss amounts to about 0.5 g/day. These losses (minus the dietary CHO intake) must be compensated for by continuous resynthesis of CHO in the intestinal tract and liver (\rightarrow B). CHO supplied by the diet is absorbed in part as such and in part in esterified form (\rightarrow B, lower right). Before it is reabsorbed, CHO-esters are split by *unspecific pancreatic carboxylesterase* to CHO, which is absorbed in

the upper part of the small intestine $(\rightarrow \mathbf{B},$ bottom). Mucosal cells contain an enzyme that re-esterifies part of the absorbed CHO: ACAT (acvl-CoA-cholesterol acvltransferase) so that both cholesterol and CHO-esters can be integrated in chylomicrons ($\rightarrow A$). CHO and CHOesters in the *chylomicron remnants* (see above) are transported to the liver, where lysosomal acid lipases again break the CHO-esters down into CHO. This CHO and that taken up from other sources (LDL, HDL) leave the liver (\rightarrow **B**): 1. by excretion into the bile (\rightarrow p. 248), 2. by conversion into bile salts which also enter the bile (\rightarrow p. 249 B), and 3. by incorporation into VLDL, the hepatic lipoprotein for export of lipids to other tissues. Under the influence of LPL (see above), the VLDL yield IDL and later **LDL** (\rightarrow **B**, left). The LDL transport CHO and CHO-esters to cells with LDL receptors (hepatic and extrahepatic cells; \rightarrow **B**, top). The receptor density on the cell surface is adjusted according to the prevailing CHO requirement. Like hepatic cells (see above) extrahepatic cells take up the LDL by receptor-mediated endocytosis, and lysosomal acid lipases reduce CHOesters to CHO (\rightarrow **B**, top right). The cells can then insert the CHO in their cell membranes or use it for steroid synthesis. A cholesterol excess leads to (a) inhibition of CHO synthesis in the cells (3-HMG-CoA-reductase) and (b) activation of ACAT, an enzyme that esterifies and stores CHO in the form of its ester (see above).

Hyperlipoproteinemia. An excess of lipids in the blood can be reflected by elevation of triacylglycerol levels and/or CHO levels (> 200-220 mg/dL serum; affects about one in five adults in Western countries). In the most severe form, familial hypercholesterolemia, a genetic defect causes elevated plasma CHO concentrations from birth on, which can result in myocardial infarction in juvenile age. The disease is caused by genetic defects of the high-affinity LDL receptors. The serum CHO level rises since the cells take up smaller quantities of cholesterol-rich LDLs. Extrahepatic tissues synthesize larger quantities of CHO because 3-HMG-CoA-reductase fails to inhibit CHO synthesis due to the decreased absorption of LDLs. As a result, more LDLs bind to the low-affinity scavenger receptors that mediate the storage of CHO in macrophages, cutaneous tissues, and blood vessels. Hypercholesterolemia therefore increases the risk of arteriosclerosis and coronary disease.





Digestion and Absorption of Carbohydrates and Protein

Carbohydrates provide half to two-thirds of the energy requirement (\rightarrow p. 226). At least 50% of dietary carbohydrates consist of starch (amylose and amylopectin), a polysaccharide; other important dietary carbohydrates are cane sugar (saccharose = sucrose) and milk sugar (lactose). Carbohydrate digestion starts in the mouth (\rightarrow A1 and p. 236). Ptyalin, an α amylase found in saliva, breaks starches down into oligosaccharides (maltose, maltotriose, a limit dextrins) in a neutral pH environment. This digestive process continues in the proximal stomach, but is interrupted in the distal stomach as the food is mixed with acidic gastric juices. A pancreatic α -amylase, with a pH optimum of 8 is mixed into the chyme in the duodenum. Thus, polysaccharide digestion can continue to the final oligosaccharide stage mentioned above. The carbohydrates can be only absorbed in the form of monosaccharides. Thus, the enzymes maltase and isomaltase integrated in the luminal brush border membrane of enterocytes break down maltose, maltotriose and α limit dextrins into **alucose** as the final product. As in the renal tubules $(\rightarrow p. 158)$, glucose is first actively taken up by the Na+ symport carrier SGLT1 into mucosal cells (\rightarrow A2, p. 29 B1) before passively diffusing into the portal circulation via GLUT2, the glucose uniport carrier (facilitated diffusion; → p. 22). The hydrolysis of saccharose, lactose, and trehalose is catalyzed by other brush border enzymes: lactase, saccharase (sucrase) and trehalase. In addition to glucose, these reactions release galactose (from lactose), which is absorbed by the same carriers as glucose. and fructose, which crosses the enterocytes by passive uniporters, GLUT5 in the luminal and GLUT2 in the basolateral membrane (\rightarrow A2).

Lactase deficiency. Lactose cannot be broken down and absorbed unless sufficient lactase is available. Lactase deficiencies lead to diarrhea 1) because water is retained in the intestinal lumen due to osmotic mechanisms, and 2) because intestinal bacteria convert the lactose into toxic substances.

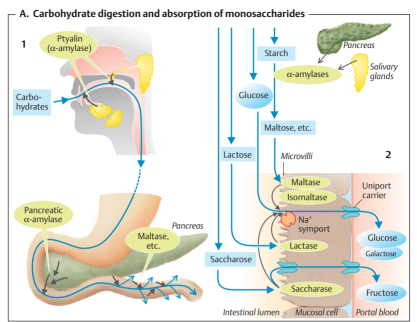
Protein digestion starts in the stomach (\rightarrow B1). *HCl* in the stomach denatures proteins and converts the three secreted *pepsinogens* into

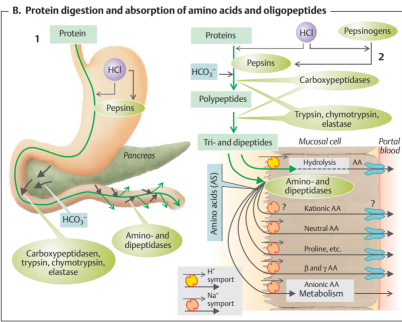
about eight different pepsins. At a pH of 2-5, these endopeptidases split off proteins at sites where tyrosine or phenylalanine molecules are incorporated in the peptide chain. The pepsins become inactive in the small intestine (pH 7-8). Pancreatic juice also contains proenzymes of other peptidases that are activated in the duodenum (\rightarrow p. 246). The endopeptidases trypsin, chymotrypsin and elastase hydrolyze the protein molecules into short-chain pentides, Carboxypentidase A and B (from the pancreas) as well as dipeptidases and aminopeptidase (brush border enzymes) act on proteins at the end of the peptide chain, breaking them down into tripeptides, dipeptides, and (mostly) individual amino acids. These cleavage products are absorbed in the duodenum and ieiunum.

Amino acids (AA) are transported by a number of specific carriers (\rightarrow B2) similar to those found in the kidneys (\rightarrow p. 158). Neutral (without net charge) and anionic ("acid") Lamino acids are transported with Na+ symporters (secondary active transport; \rightarrow p. 28) from the intestinal lumen into mucosal cells, from which they passively diffuse with carriers into the blood. Cationic ("basic") L-amino acids such as L-arginine+, L-lysine+ and L-ornithine+ are partly taken up into the enterocytes by Na⁺ independent mechanisms, as the membrane potential is a driving force for their uptake. Anionic amino acids like L-glutamateand L-aspartate which, for the most part, are broken down in the mucosal cells, also have their own (Na+ and K+ dependent) carrier systems. Neutral amino acids use several different transporters.

AA absorption disorders can be congenital and affect various amino acid groups. These disorders are often associated with defects of renal tubular reabsorption (renal aminoaciduria, e.g. cystinuria).

Dipeptides and tripeptides can be absorbed as intact molecules by a symport carrier (PepT1). The carrier is driven by an H^+ gradient (\rightarrow **B2**), which in turn is generated by H^+ secretion (tertiary active H^+ -peptide symport, \rightarrow p. 29 B5). Amino acids generally are much more rapidly absorbed as dipeptides and tripeptides than as free amino acids. Once they enter the cells, the peptides are hydrolyzed to free amino acids.





Vitamin Absorption

Since higher animals cannot synthesize **cobalamins** (**vitamin B**₁₂), they must obtain this *cobalt-containing coenzyme* from the diet. Animal products (liver, kidneys, fish, eggs, milk) are the main source.

Cobalamin biochemistry. Aqua- and OH-cobalamin are precursors of the two active forms, methyl- and adenosylcobalamin. *Methylcobalamin* is needed to form methionine from homocysteine; cobalamin transfers the methyl group required for this from N⁵-methyltetrahydrofolate (see below) to homocysteine. Some enzymes, e.g. methyl-malonyl-CoA mutase, need *adenosylcobalamin* to break and form carbon-carbon bonds.

Cobalamins are relatively large and lipophobic molecules that require transport proteins $(\rightarrow A)$. During passage through the GI tract, plasma and other compartments, cobalamins bind to (1) intrinsic factor (IF), which is secreted by gastric parietal cells; (2) transcobalamin II (**TC II**) in plasma; and (3) R proteins in plasma (TC I), and granulocytes (TC III), saliva, bile, milk, etc. Gastric acid releases cobalamin from dietary proteins. In most cases, the cobalamin then binds to R protein in saliva or (if the pH is high) to IF (\rightarrow A1). The R protein is digested by trypsin in the duodenum, resulting in the release of cobalamin, which is then bound by (trypsinresistant) intrinsic factor. The mucosa of the terminal ileum has highly specific receptors for the cobalamin-IF complex. IT binds to these receptors and is absorbed by receptor-mediated endocytosis, provided a pH of > 5.6 and Ca²⁺ ions are available (\rightarrow **A2**). The receptor density and, thus, the absorption rate increases during pregnancy. Cobalamin binds to TC I, II and III in plasma (→A3). TC II mainly distributes cobalamin to all cells undergoing division (TC II receptors, endocytosis). TC III (from granulocytes) transports excess cobalamin and unwanted cobalamin derivatives to the liver (TC III receptors), where it is either stored or excreted in the bile. TC I has a half-life of roughly 10 days and serves as a short-term depot for cobalamin in the plasma.

A vegan diet or disturbed cobalamin absorption can lead to severe **deficiency symptoms** like pernicious anemia and spinal cord damage (funicular myelosis).

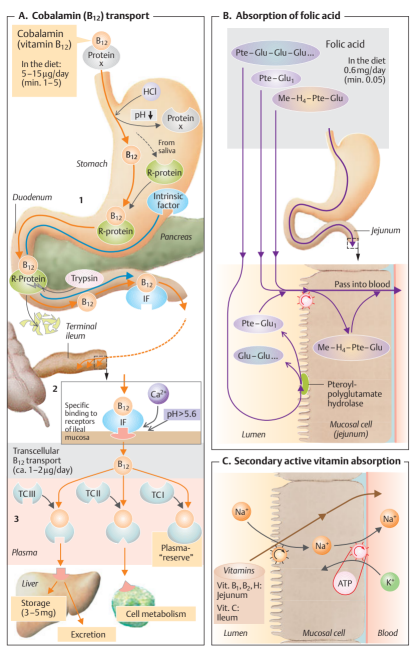
It takes years for these symptoms to manifest as the body initially has a reserve of 1000 times the daily requirement of $1 \mu q (\rightarrow p. 90)$.

Folic acid/folate (= pteroylglutamic acid). N^5 , N¹⁰-methylenetetrahydrofolate, the metabolically active form of folic acid (daily requirement: 0.1-0.2 mg) is needed for DNA synthesis (formation of deoxythymidylate from deoxyuridylate). Folic acid in the diet usually occurs in forms that contain up to seven glutamyl residues (y-linked peptide chain: Pte-Gluz) instead of pteroylglutamic acid (Pte-Glu₁). Since only Pte-Glu₁ can be absorbed from the lumen of the proximal jejunum $(\rightarrow B)$, its polyglutamyl chain must be shortened before absorption. This is done by pterovlpolyglutamate hydrolases located in the luminal membrane of enterocytes. The absorption of Pte-Glu₁ is mediated by a specific active transporter. In mucosal cells, Pte-Glu1 is than broken down to vield N⁵-methyltetrahydrofolate (5-Me-H₄-folate) and other metabolites. If already present in the ingested food, these metabolites are absorbed from the intestinal lumen by the aforementioned mechanism. (The same applies to the cytostatic drug, methotrexate.) Methylcobalamin is needed to convert 5-Me-H₄-folate to tetrahydrofolate (see above). The body stores about 7 mg of folic acid, enough for several months (cf. folic acid deficiency, \rightarrow p. 90).

The **other water-soluble vitamins**—B₁ (thiamin), B₂ (riboflavin), C (ascorbic acid), and H (biotin, niacin)—are absorbed via Na⁺ symport carriers (\rightarrow C). Vitamin C is absorbed from the ileum, whereas vitamins B₁, B₂, and H are absorbed from the jejunum. Members of the vitamin B₆ group (pyridoxal, pyridoxine, pyridoxamine) are probably absorbed by passive mechanisms.

Fat-soluble vitamins—A (retinol), D_3 (cholecalciferol), E (tocopherol), K_1 (phylloquinone), and K_2 (menaquinone)—must be incorporated into *micelles* for absorption (cf. lipid digestion, p. 252). The exact absorption mechanism has not yet been explained, though it is known to be partly saturation—and energy-dependent. Fat-soluble vitamins are incorporated into chylomicrons and VLDL for transport in plasma (\rightarrow p. 254ff.).

261



Water and Mineral Absorption

The average intake of water (in beverages and foodstuffs) is roughly 1.5 L per day. An additional 7 L of fluid are secreted into the gastrointestinal (GI) tract (saliva, gastric juices, bile, pancreatic juice and intestinal secretions), whereas only about 0.1 L/day is eliminated in the feces. The digestive tract must therefore absorb a net volume of at least 8.4 L of water per day. GI absorption of water occurs mainly in the jejunum and ileum, with smaller quantities being absorbed by the colon ($\rightarrow A$). Water is driven through the intestinal epithelium by osmosis. When solutes (Na+, Cl-, etc.) are absorbed in the intestine, water follows $(\rightarrow B)$. (The stool contains only small quantities of Na⁺, Cl⁻ and water.) Conversely, the secretion of substances into the lumen or the ingestion of non-absorbable substances leads to water fluxes into the lumen. Poorly absorbable substances therefore act as laxatives (e.g. sulfate, sorbitol, polyethylene glycol).

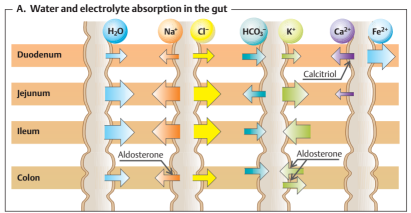
Water absorption is mainly driven by the absorption of Na⁺, Cl⁻ and organic compounds (→ B). The luminal concentration of Na⁺ and Cl⁻ steadily decreases from the duodenum to the colon. That of Na+, for example, is approximately 145 mmol/L in the duodenum, 125 mmol/L in the ileum and only 40 mmol/L in the colon (\rightarrow **C**). Na⁺ is absorbed by various mechanisms, and the Na+-K+-ATPase on the basolateral cell membrane is the primary driving mechanism (\rightarrow p. 26) for all of them (\rightarrow **B, D**). Symport of Na⁺ and organic substances (→ see pp. 26ff. and 258): Na⁺ passively influxes into cells of the duodenum and jejunum via symporter carriers, which actively cotransport glucose, amino acids, phosphates and other compounds (secondary active transport; \rightarrow **D1**). Since this is an electrogenic transport mechanism (\rightarrow p. 28), a lumen-negative transepithelial potential (LNTP; \rightarrow p. 162) that drives Cl^- out of the lumen forms ($\rightarrow D2$).

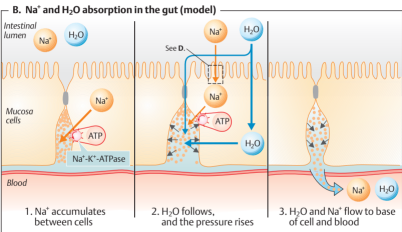
• Parallel transport of Na⁺ and Cl[−]: Na⁺ ions in the lumen of the ileum are exchanged for H⁺ ions (→ D3) while Cl[−] is exchanged for HCO₃[−] at the same time (→ D4). The H⁺ ions combine with HCO₃[−] to yield H₂O + CO₂, which diffuse out of the lumen. Most Na⁺, Cl[−] and, by subsequent osmosis, H₂O is absorbed by this electroneutral transport mechanism. • Na⁺ diffusion: Na⁺ in the colon is mainly absorbed through luminal Na⁺ channels (→ D5). This type of Na⁺ transport is electrogenic and aldosterone-dependent (→ p. 182). The related lumen-negative transepithelial potential (LMTP, see above) either leads to K⁺ secretion or drives Cl⁻ out of the lumen (→ D2).

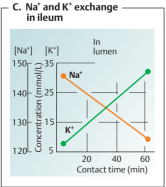
The CI secretion mechanism of epithelial cells (mainly Lieberkühn's crypts, → p. 245, A16) is similar to that of the acini of salivary glands ($\rightarrow p$, 236). The efflux of Cl⁻ into the lumen and the associated efflux of Na⁺ and water are stimulated by cAMP and requlated by neurons and hormones such as VIP (vasoactive intestinal peptide) and prostaglandins. The physiological function of this form of H₂O secretion could be to dilute viscous chyme or to ensure the recirculation of water (from the crypts \rightarrow lumen \rightarrow villi → crypts) to promote the absorption of poorly soluble substances. Cholera toxin inhibits the GTPase of the G_s proteins (\rightarrow p. 274), thereby maintaining a maximal cAMP concentration and therefore a marked increase in CI- secretion. In response to it, large quantities of water and Na⁺ are secreted into the lumen, which can lead to severe diarrhea (up to 1 L/hour!).

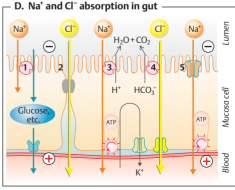
In addition to HCO_3^- from pancreatic juice, HCO_3^- is also secreted into the intestinal lumen by the small and large intestine (\rightarrow A). K⁺ is secreted (aldosterone-dependent) by crypt cells of the colon (luminal K⁺ concentration \approx 90 mmol/l!) and reabsorbed via the H⁺-K⁺ pump of the surface epithelium (similar to the mechanism in the stomach). The aldosterone-dependent K⁺ secretion/absorption ratio determines the net amount of K⁺ excreted (\rightarrow A and p. 180). *Diarrhea* results in losses of K⁺ and HCO_3^- (hypokalemia and metabolic acidosis; \rightarrow p. 142).

Ca²⁺. The stool contains one-third of the dietary Ca^{2+} intake. Ca^{2+} is absorbed in the upper part of the small intestine (\rightarrow **A**) with the aid of intracellular *calcium-binding protein* (CaBP). Calcitriol increases CaBP synthesis, thereby increasing Ca^{2+} absorption (\rightarrow p. 292). Deficiencies of vitamin D or substances that form water-insoluble compounds with Ca^{2+} (phytin, oxalate, fatty acids) decrease Ca^{2+} absorption. Mg^{2+} is absorbed by similar mechanisms, but **iron** (Fe) is absorbed by a different mechanism (\rightarrow p. 90).









Large Intestine, Defecation, Feces

Anatomy. The terminal end of the gastrointestinal tract includes the *large intestine* (*cecum* and *colon*, ca. 1.3 m in length) and *rectum*. The large intestinal mucosa has characteristic pits (crypts), most of which are lined with mucus-forming cells (*goblet cells*). Some of the surface cells are equipped with a brush border membrane and reabsorb ions and water.

The large intestine has two **main functions**: (1) It serves as a *reservoir* for the intestinal contents (cecum, ascending colon, rectum). (2) It absorbs water and electrolytes (\rightarrow p. 262), so the ca. 500–1500 mL of chyme that reaches the large intestine can be reduced to about 100–200 mL. The large intestine is not an essential organ; therefore, large segments of the intestine can be removed—e.g., for treatment of cancer

Water instilled into the **rectum** via an *enema* is reabsorbed. Anally delivered drugs (*suppositories*) also diffuse through the intestinal wall into the bloodstream. Substances administered by this route bypass the liver and also escape the effects of gastric acid and digestive enzymes.

Motility. Different local mixing movements of the large intestine can be distinguished, e.g., powerful segmentation contractions associated with pouch formation (haustration) and anterograde or retrograde peristaltic waves (pacemaker located in transverse colon). Thus, stool from the colon can also be transported to the cecum. Mass movements occur 2−3 times daily (→ A). They are generally stimulated by a meal and are caused by the gastrocolic reflex and gastrointestinal hormones.

The typical sequence of **mass movement** can be observed on X-ray films after administration of a barium meal, as shown in the diagrams (\rightarrow **A1** – **8**). **A1**, barium meal administered at 7:00 a.m. **A2**, 12 noon: the barium mass is already visible in the last loop of the ileum and in the cecum. Lunch accelerates the emptying of the ileum. **A3**, about 5 minutes later, the tip of the barium mass is choked off. **A4**, shortly afterwards, the barium mass fills the transverse colon. **A5**, haustration divides the barium mass in the transverse colon, thereby mixing its contents. **A6–8**, a few minutes later (still during the meal), the transverse colon suddenly contracts around the leading end of the intestinal contents and rapidly propels them to the sigmoid colon.

Intestinal bacteria. The intestinal tract is initially sterile at birth, but later becomes colonized with orally introduced anaerobic

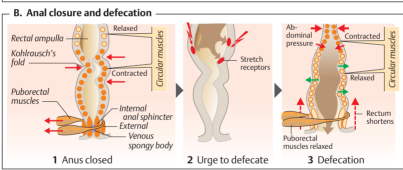
bacteria during the first few weeks of life. The large intestine of a healthy adult contains 1011 to 10¹² bacteria per mL of intestinal contents; the corresponding figure for the ileum is roughly 106/mL. The low pH inside the stomach is an important barrier against pathogens. Consequently, there are virtually no bacteria in the upper part of the small intestine $(0-10^4)$ mL). Intestinal bacteria increase the activity of intestinal immune defenses ("physiological inflammation"), and their metabolic activity is useful for the host. The bacteria synthesize vitamin K and convert indigestible substances (e.g. cellulose) or partially digested saccharides (e.g. lactose) into absorbable shortchain fatty acids and gases (methane, H2, CO2).

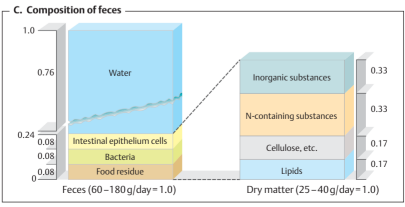
The **anus** is normally closed. Anal closure is regulated by Kohlrausch's valve (transverse rectal fold), the puborectal muscles, the (involuntary) internal and (voluntary) external anal sphincter muscles, and a venous spongy body. Both sphincters contract tonically, the internal sphincter (smooth muscle) intrinsically or stimulated by sympathetic neurons (L_1 , L_2) via α -adrenoceptors, the external sphincter muscle (striated muscle) by the pudendal nerve.

Defecation. Filling of the upper portion of the rectum (rectal ampulla) with intestinal contents stimulates the rectal stretch receptors $(\rightarrow B2)$, causing reflex relaxation of the internal sphincter (accommodation via VIP neurons), constriction of the external sphincter, and an *urge to defecate*. If the (generally voluntary) decision to defecate is made, the rectum shortens, the puborectal and external anal sphincter muscles relax, and (by a spinal parasympathetic reflex via S_2 – S_4) annular contractions of the circular muscles of the descending colon, sigmoid colon and rectum-assisted by increased abdominal pressure—propel the feces out of the body $(\rightarrow B)$. The normal frequency of bowel evacuation can range from 3 times a day to 3 times a week, depending on the dietary content of indigestible fiber (e.g. cellulose, lignin). Frequent passage of watery stools (diarrhea) or infrequent stool passage (constipation) can lead to various disorders.

Stool (feces; \rightarrow **C).** The average adult excretes 60–80 g of feces/day. Diarrhea can raise this over 200 g/d. Roughly $^{1}/_{4}$ of the feces is composed of *dry matter*, about $^{1}/_{3}$ is attributable to be train from the large intesting.

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Hormones and Reproduction

Integrative Systems of the Body

Unlike unicellular organisms, multicellular organisms have numerous specialized groups of cells and organs, the many different functions of which must be expediently integrated and coordinated (see also p. 2). In mammals, the nervous system and endocrine system are chiefly responsible for control and integration. while the immune system serves as an information system for corporal immune defense $(\rightarrow p.94ff.)$. These systems communicate by way of electrical and/or chemical signals $(\rightarrow A)$.

Nerve impulses and hormonal signals serve to control and regulate $(\rightarrow p.4)$ the metabolism and internal milieu (blood pressure, pH, water and electrolyte balance, temperature, etc.), physical growth and maturation, reproductive functions, sexual response, and responses to the social environment. The signals received by sensors (= sensory receptors) in the inner organs, musculoskeletal system, skin and the sensory organs, as well as psychological factors, skeletal muscles and other factors also play a part in regulation and control. The signals are used by many feedback mechanisms in the body (\rightarrow p. 4).

Nerve fibers are specifically adapted for rapid transmission of finely graded signals. The nervous system consists of the central nervous system (CNS; \rightarrow p. 310ff.) and peripheral nervous system. The latter consists of:

- The somatic nervous system, which conducts impulses from non-visceral sensors to a center (afferent neurons) and controls the skeletal musculature (efferent neurons).
- The peripheral autonomic nervous system (→ p. 78ff.), which consists of efferent neurons and mainly functions to control the circulatory system, inner organs and sexual functions. It is supplemented by:
- Visceral afferent neurons, i.e., nerve fibers that conduct signals from inner organs to a center. They are usually located in the same nerves as autonomous fibers (e.g., in vagus nerve); and the
- Enteric nervous system, which integrates the local functions of the esophagus, stomach and gut (\rightarrow p. 234).

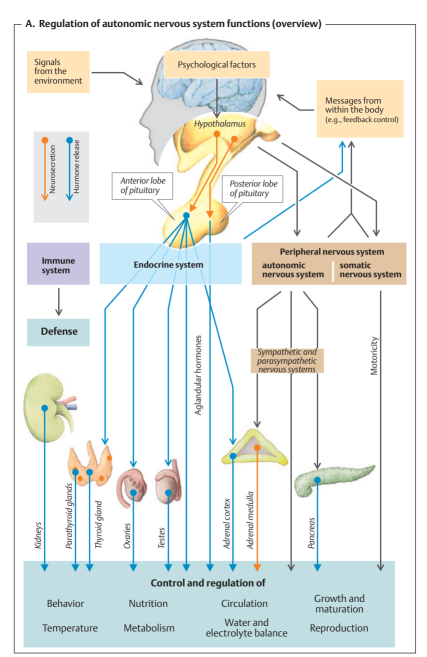
Hormones. Like neurotransmitters (see below) and the immune system's cytokines and chemokines (→ p. 94ff.), hormones serve as messenger substances that are mainly utilized for slower, long-term transmission of signals. Endocrine hormones are carried by the blood to target structures great distances away. Paracrine hormones (and other paracrine transmitters) only act on cells in the immediate vicinity of the cells from which they are released. Hormones that act on the cells that produced the messenger substance are referred to as autocrine hormones.

Hormones are synthesized in specialized glands, tissues and cells (e.g., neuroendocrine cells). Their target organ is either a subordinate endocrine gland (glandotropic hormone) or non-endocrine tissue (aglandotropic hormone). The target cells have high-affinity binding sites (receptors) for their specific hormone, so very low concentrations of the hormone suffice for signal transduction (10⁻⁶ to 10-12 mol/L). The receptors on the target cells pick out the substances specifically intended for them from a wide variety of different messenger substances in their environment.

Hormones work closely with the nervous system to regulate digestion, metabolism, growth, maturation, physical and mental development, maturation, reproduction, adaptation, and the internal milieu of the body (homeostasis) (\rightarrow A). Most of these actions are predominately autonomous functions subject to central control by the hypothalamus, which is controlled by higher centers of the brain $(\rightarrow p.330)$.

Neurotransmitters released at chemical synapses of nerve endings transmit signals to postsynaptic nerve fibers, muscles or glands $(\rightarrow p.50ff.)$. Some neuropeptides released by presynaptic neurons also exert their effects in neighboring synapses, resulting in a kind of "paracrine" action.

Neurons can also secrete hormones, e.g., epinephrine, oxytocin and antidiuretic hormone. Some transmitter substances of the immune system, e.g. thymosin and various cytokines, also have endocrine effects.



Hormones

Hormones are messenger substances that convey information signals relevant to cell function (→ p. 266). **Endocrine hormones**, i.e., those transported in the bloodstream, are produced in *endocrine glands* such as the hypothalamus, thyroid, parathyroid glands, adrenal medulla, pancreatic islets, ovaries and testes. They are also synthesized in diffusely scattered *endocrine cells* of the CNS, in C cells of the thyroid, and in the thymus, atria, kidneys, liver, gastrointestinal tract, etc. **Paracrine hormones**, i.e., those that affect nearby cells only (tissue hormones or *mediators*; see below) are secreted by cells widely distributed throughout the body.

Types of hormone.

- 1. **Peptide hormones** (\rightarrow **A**, dark blue areas) and **glycoprotein hormones** (\rightarrow **A**, light blue areas) are hydrophilic hormones stored in *secretory granules* and released by exocytosis as required. Multiple hormones can be produced from a single gene (\rightarrow e.g., POMC gene, p.280) by variable splicing and posttranslational modification (\rightarrow p. 8 ff.).
- 2. Steroid hormones (\rightarrow A, yellow areas) and calcitriol are chemically related lipophilic hormones metabolized from *cholesterol* (\rightarrow pp. 292ff). They are not stored, but are synthesized as needed.
- 3. Tyrosine derivatives (\rightarrow A, orange areas) include (a) the *hydrophilic catecholamines* dopamine, epinephrine and norepinephrine (\rightarrow p. 84) and (b) *lipophilic thyroid hormones* (T₃, T₄; \rightarrow p. 286).

The **lipophilic hormones** in (2) and (3b) are **transported** in the blood while bound to plasma proteins. Corticosteroids are carried bound to globulin and albumin, testosterone and estrogen to sex hormone-binding globulin and T_3 and T_4 to albumin and two other plasma proteins (\rightarrow p. 286).

Hormone receptors. The receptors (docking sites) for glycoprotein hormones, peptide hormones and catecholamines are transmembrane proteins (\rightarrow p. 14) that bind to their specific hormone on the outer cell surface. Many of these hormones induce the release of intracellular **second messengers** that transmit the hormone signal inside the cell. cAMP,

cGMP, IP3, DAG, Ca2+ and NO function as second messengers (and sometimes as third messengers; \rightarrow p. 274ff.). Some peptide hormones like insulin, prolactin, atriopeptin and numerous growth factors bind to cell surface receptors with cytosolic domains with enzymatic activity (\rightarrow p. 278). Steroid hormones, on the other hand, enter the cells themselves $(\rightarrow p. 278)$. Once they bind to cytosolic receptor proteins, steroid hormones (as well as calcitriol, T₃ and T₄) are transported to the cell nucleus, where they influence transcription (genomic action). A target cell can have different receptors for different hormones (e.g., insulin and glucagon) or different receptors for a single hormone (e.g., α_1 and β_2 adrenoceptors for epinephrine).

Hierarchy of hormones $(\rightarrow A)$. The secretion of hormones is often triggered by neural impulses from the CNS. The hypothalamus is the main neurohormonal control center (\rightarrow p. 280 and 330). Hypothalamic neurons extend to the posterior pituitary (neurohypophysis). The hormones are secreted either by the hypothalamus itself or by the posterior pituitary. Hypothalamic hormones also control hormone release from the anterior pituitary (adenohypophysis). Anterior pituitary glandotropic hormones control peripheral endocrine glands (→ A top, green areas), which release the endhormone $(\rightarrow A)$. The original signal can be amplified or modulated at these relay sites $(\rightarrow p. 272)$.

Pituitary hormones. Hypothalamic hormones control *anterior pituitary* hormone secretion by either stimulating or inhibiting hormone production. They are therefore called *releasing hormones (RH)* or *release-inhibiting hormones (IH)*, resp. (\rightarrow **A** and table). Most anterior pituitary hormones are glandotropic (\rightarrow p.280). The *posterior pituitary* hormones are released by neuronal signals and are mainly aglandotropic (\rightarrow p.280).

Other **endocrine hormones** are secreted largely independent of the hypothalamic-pituitary axis, e.g., pancreatic hormones, parathyroid hormone (PTH), calcitonin and calcitriol, angiotensin II, aldosterone (\rightarrow p. 182ff.), erythropoietin (\rightarrow p. 88) and gastrointestinal hormones (\rightarrow p. 234). Atriopeptin is secreted from the heart atrium in response to stretch

stimuli (\rightarrow p. 170), whereas the release of *melatonin* is subject to afferent neuron control (\rightarrow p. 334).

Some of these hormones (e.g., angiotensin II) and *tissue hormones* or **mediators** exert *paracrine effects* within endocrine and exocrine glands, the stomach wall, other organs, and on inflammatory processes. Bradykinin (\rightarrow pp. 214 and 236), histamine (\rightarrow pp. 100 and 242), serotonin (5-hydroxytryptamine, \rightarrow p. 102) and eicosanoids are members of this group.

Eicosanoids. Prostaglandins (PG), thromboxane (TX), leukotrienes and epoxyeicosatrienoates are eicosanoids (Greek εικοσι = twenty [C atoms]) derived in humans from the fatty acid **arachidonic acid (AA)**. (Prostaglandins derived from AA have the index number 2). AA occurs as an ester in the phospholipid layer of the cell membranes and is obtained from dietary sources (meat), synthesized from linoleic acid, an essential fatty acid, and released by *phospholipase A₂* (→ p. 252).

Pathways of **eicosanoid synthesis** from arachidonic acid (AA):

- 1. Cyclooxygenase pathway: Cyclooxygenase (COX)-1 and COX-2 convert AA into PGG₂, which gives rise to PGH₂, the primary substance of the biologically active compounds PGE₂, PGD₂, PGF_{2α}, PGI₂ (prostacyclin) and TXA₂. COX-1 and 2 are inhibited by nonsteroidal anti-inflammatory drugs (e.g., Aspirin®).
- 2. Lipoxygenase pathway: Leukotriene A₄ is synthesized from AA (via the intermediate 5-HPETE = 5-hydroperoxyeicosatetraenoate) by way of 5-lipoxygenase (especially in neutrophilic granulocytes). Leukotriene A₄ is the parent substance of the leukotrienes C₄, D₄ and E₄. The significance of 12lipoxygenase (especially in thrombocytes) is not yet clear, but 15-lipoxygenase is known to produce vasoactive lipoxins (LXA₄, LXB₄).
- 3. Cytochrome P450-epoxygenase produces epoxyeicosatrienoates (EpETrE = EE).

Typical effects of eicosanoids:

PGE₂ dilates the bronchial and vascular musculature (and keeps the lumen of the fetal ductus arteriosus and foramen ovale open; \rightarrow p. 220), stimulates intestinal and uterine contractions, protects the gastric mucosa (\rightarrow p. 242), inhibits lipolysis, increases the glomerular filtration rate (GFR), plays a role in fever development (\rightarrow p. 224), sensitizes nociceptive nerve endings (pain) and increases the

permeability of blood vessels (inflammation). PGD_2 induces bronchoconstriction. PGI_2 (prostacyclin) synthesized in the endothelium, is vasodilatory and inhibits platelet aggregation. TXA_2 , on the other hand, occurs in platelets, promotes platelet aggregation and acts as a vasoconstrictor (\rightarrow p. 102). 11,12-EpETrE has a vasodilatory effect (= EDHF, \rightarrow p. 214).

Hormones (h.) of the hypothalamus and pituitary

Abbreviation/synonyme

Thyrotropin RH, TRH, TRF

Hypothalamus

The suffix "-liberin" denotes releasing
h. (RH) or factor (RF); "-statin" is used for releaseinhibiting h. (IH) or factors (IF)

Corticoliberin Corticotropin RH, CRH, CRF

Gonadoliberin Gonadotropin RH, Gn-RH; ICSH

Gonadoliberin Gonadotropin RH, Gn-RH; ICSH
Prolactostatin
Somatoliberin Somatotropin RH, SRH, SRF,
GHRH, GRH
Somatostatin** Somatotropin (growth h.) IH, SIH

Anterior lobe of the pituitary

Name*

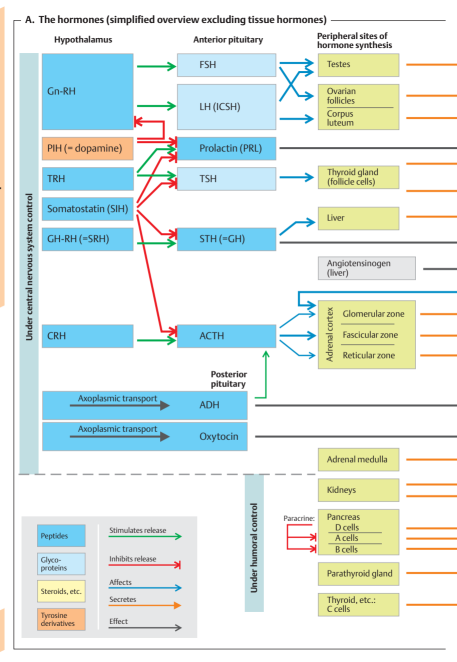
Thyroliberin

Adrenocorticotropic h. (ACTH) Corticotropin Follitropin Follicle-stimulating h. (FSH) Luteinizing h. (LH), interstitial Lutropin cell-stimulating h. (ICSH) Melanotropin α-Melanocyte-stimulating h. $(\alpha$ -MSH), α -melanocortin Somatotropic h. (STH), growth h. Somatotropin Thyrotropin Thyroid stimulating h. (TSH) Prolactin PRL, lactogenic (mammotropic)

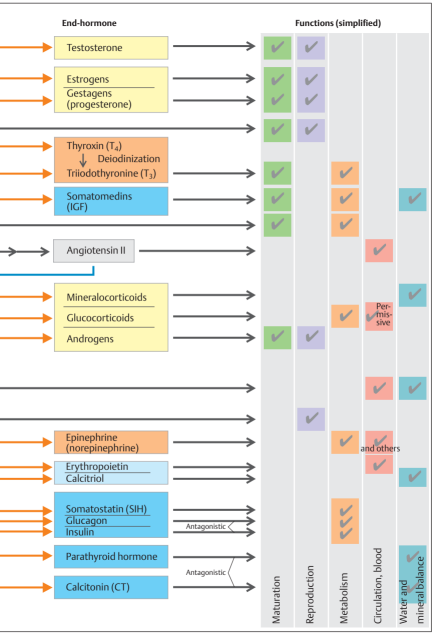
Posterior lobe of the pituitary

Oxytocin –
Adiuretin Anti-diuretic h. ADH,
(arginine-) vasopressin (AVP)

- * Names generally recommended by IUPAC-IUB Committee on Biochemical Nomenclature.
- ** Also synthesized in gastrointestinal organs, etc.



271



272

Humoral Signals: Control and Effects

Hormones and other humoral signals function to provide **feedback control**, a mechanism in which the response to a signal feeds back on the signal generator (e.g., endocrine gland). The speed at which control measures are implemented depends on the rate at which the signal substance is broken down—the quicker the degradation process, the faster and more flexible the control.

In negative feedback control, the response to a feedback signal opposes the original signal. In the example shown in A1, a rise in plasma cortisol in response to the release of corticoliberin (corticotropin-releasing hormone, CRH) from the hypothalamus leads to down-regulation of the signal cascade "CRH ⇒ ACTH ⇒ adrenal cortex," resulting in a decrease in cortisol secretion. In shorter feedback loops. ACTH can also negatively feed back on the hypothalamus (\rightarrow A2), and cortisol, the end-hormone, can negatively feed back on the anterior pituitary (\rightarrow A3). In some cases, the metabolic parameter regulated by a hormone (e.g., plasma glucose concentration) rather then the hormone itself represents the feedback signal. In the example $(\rightarrow B)$, glucagon increases blood glucose levels (while insulin decreases them), which in turn inhibits the secretion of glucagon (and stimulates that of insulin). Neuronal signals can also serve as feedback (neuroendocrine feedback) used, for example, to regulate plasma osmolality $(\rightarrow p. 170)$.

In **positive feedback control**, the response to the feedback amplifies the original signal and heightens the overall response (e.g., in *autocrine regulation*; see below).

The higher hormone not only controls the synthesis and excretion of the end-hormone, but also controls the **growth of peripheral endocrine gland**. If, for example, the end-hormone concentration in the blood is too low despite maximum synthesis and secretion of the existing endocrine cells, the gland will enlarge to increase end-hormone production. This type of **compensatory hypertrophy** is observed for instance in goiter development (→ p. 288) and can also occur after surgical excision of part of the gland.

Therapeutic administration of a hormone (e.g., cortisone, a cortisol substitute) have the same effect on higher hormone secretion (ACTH and CRH in the example) as that of the end-hormone (cortisol in the example) normally secreted by the peripheral gland (adrenal cortex in this case). *Long-term* administration of an end-hormone would therefore lead to inhibition and atrophy of the endocrine gland or cells that normally produce that hormone. This is known as **compensatory atrophy**.

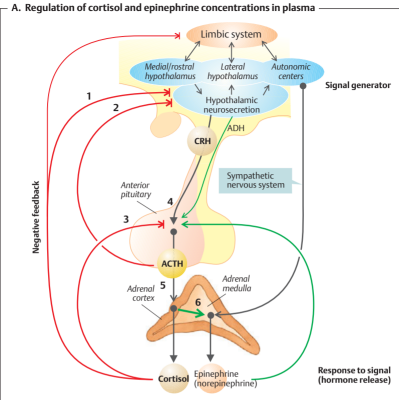
A **rebound effect** can occur if secretion of the higher hormone (e.g., ACTH) is temporarily elevated after discontinuation of end-hormone administration.

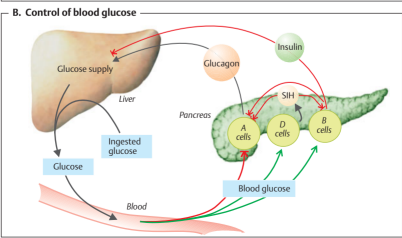
The **principal functions** of *endocrine hormones*, *paracrine hormones* and other humoral transmitter substances are to control and regulate:

- enzyme activity by altering the conformation (allosterism) or inhibiting/stimulating the synthesis of the enzyme (induction);
- transport processes, e.g., by changing the rate of insertion and synthesis of ion channels/ carriers or by changing their opening probability or affinity;
- growth (see above), i.e., increasing the rate of mitosis (proliferation), "programmed cell death" (apoptosis) or through cell differentiation or dedifferentiation:
- ◆ secretion of other hormones. Regulation can occur via endocrine pathways (e.g., ACTH-mediated cortisol secretion; \rightarrow **A5**), a short portal vein-like circuit within the organ (e.g., effect of CRH on ACTH secretion, \rightarrow **A4**), or the effect of cortisol from the adrenal cortex on the synthesis of epinephrine in the adrenal medulla, (\rightarrow **A6**), or via paracrine pathways (e.g., the effect of somatostatin, SIH, on the secretion of insulin and glucagon; \rightarrow **B**).

Cells that have receptors for their own humoral signals transmit **autocrine signals** that function to

- ◆ exert negative feedback control on a target cell, e.g., to discontinue secretion of a transmitter (e.g., norepinephrine; → p. 84);
- coordinate cells of the same type (e.g., in growth);
- ◆ exert positive feedback control on the secreting cell or to cells of the same type. These mechanisms serve to amplify weak signals as is observed in the eicosanoid secretion or in T cell clonal expansion (→ p. 96ff.).





Hormones, neurotransmitters (\rightarrow p. 55 and p. 82), cytokines and chemokines (\rightarrow p. 94ff.) act as *messenger substances* (**first messengers**) that are transported to their respective target cells by extracellular pathways. The target cell has a high-affinity binding site (**receptor**) for its *specific* messenger substance.

Glycoprotein and peptide messengers as well as catecholamines bind to cell surface receptors on the target cell. Binding of the messenger to its receptor usually triggers certain protein-protein interactions (and sometimes protein-phospholipid interactions). This leads to the release of secondary messenger substances (second messengers) that forward the signal within the cell. Cyclic adenosine monophosphate (cAMP), cyclic guanosine monophosphate (cGMP). inositol 1.4.5-trisphosphate (IP₃), 1,2-diacylglycerol (DAG) and Ca2+ are such second messengers. Since the molecular structure of the receptor ensures that the effect of the first messenger will be specific, multiple first messengers can use the same second messenger. Moreover, the intracellular concentration of the second messenger can be raised by one messenger and lowered by another. In many cases, different types of receptors exist for a single first messenger.

cAMP as a Second Messenger

For a cAMP-mediated response to occur, the cell membrane must contain stimulatory (Gs) or inhibitory (Gi) G proteins (guanyl nucleotide-binding proteins) ($\rightarrow A1$). These G proteins consist of three subunits—alpha (α_s or α_i), beta (β) and gamma (γ)—and are therefore heterotrimers. Guanosine diphosphate (GDP) is bound to the α -subunit of an inactive G protein. Once the first messenger (M) binds to the receptor (Rec.), the M-Rec. complex conjugates with the G_s -GDP (or G_i -GDP) molecule (\rightarrow A2). GDP is then replaced by cytosolic GTP, and the $\beta\gamma$ -subunit and the M-Rec. complex dissociate from the α-subunit if Mg²⁺ is present (\rightarrow A3). α_s -GTP or α_i -GTP remain as the final products. Adenylate cyclase on the inside of the cell membrane is activated by α_s - GTP (cytosolic cAMP concentration rises) and inhibited by α_i -GTP (cAMP concentration falls; \rightarrow **A3**).

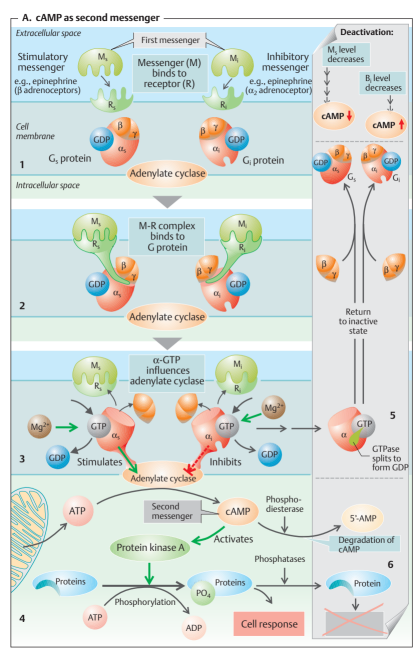
G₅-activating messengers. ACTH, adenosine (A_{2A} and A_{2B} rec.), antidiuretic hormone = vasopressin (V_2 rec.), epinephrine and norepinephrine (β_1 , β_2 , β_3 adrenoceptors), calcitonin, CGRP, CRH, dopamine (D_1 and D_5 rec.), FSH, glucagon, histamine (H_2 rec.), oxytocin (V_2 rec., see above), many prostaglandins (DP, IP, EP₂ and EP₄ rec.), serotonin = 5-hydroxytryptamine (5-HT₄ and 5-HT₇ rec), secretin and VIP activate G_5 proteins, thereby raising cAMP levels. TRH and TSH induce partial activation.

G_I-activating messengers. Some of the above messenger substances also activate G_i proteins (thereby lowering cAMP levels) using a different binding receptor. Acetylcholine (M_2 and M_4 rec.), adenosine (A_1 and A_2 rec.), epinephrine and norepinephrine (α_2 adrenoceptors), angiotensin II, chemokines, dopamine (D_2 , D_3 and D_4 rec.), GABA (GABA $_8$ rec.), glutamate (mGLU $_{2-4}$ and mGLU $_{6-8}$ rec.), melatonin, neuropeptide Y, opioids, serotonin = 5-hydroxytryptamine (5-HT $_1$ rec.), somatostatin and various other substances activate G_1 proteins.

Effects of cAMP. cAMP activates type A protein kinases (PKA = protein kinase A) which then activate other proteins (usually enzymes and membrane proteins, but sometimes the receptor itself) by phosphorylation (→A4). The specific response of the cell depends on the type of protein phosphorylated, which is determined by the type of protein kinases present in the target cell. Phosphorylation converts the proteins from an inactive to an active form or vice versa.

Hepatic glycogenolysis, for instance, is dually increased by cAMP and PKA. Glycogen synthase catalyzing glycogen synthesis is inactivated by phosphorylation whereas glycogen phosphorylase stimulating glycogenolysis is activated by cAMP-mediated phosphorylation.

Signal transduction comprises the entire signaling pathway from the time the first messenger binds to the cell to the occurrence of cellular effect, during which time the signal can be (a) *modified* by other signals and (b) *amplified* by many powers of ten. A single adenylate cyclase molecule can produce numerous cAMP and PKA molecules, which in turn can phosphorylate an enormous number of enzyme molecules. The interposition of more kinases can lead to the formation of long



kinase cascades that additionally amplify the original signal while receiving further regulatory signals.

Deactivation of the signaling cascade (\rightarrow **A, right** panel) is induced by the α-subunit in that its GTP molecule splits off GDP and P_i after reacting with its *GTPase* (\rightarrow **A5**), and the subunit subsequently binds to a βγ subunit to again form the trimeric G protein. *Phosphodiesterase* also converts cAMP into inactive 5'-AMP (\rightarrow **A4**, **A6**), and *phosphatases* dephosphorylate the protein previously phosphorylated by protein kinase A (\rightarrow **A4**). Another way to inactivate a receptor in the presence of high messenger concentrations is to make the receptor insensitive by phosphorylating it (*desensitization*).

Cholera toxin inhibits the GTPase, thereby blocking its deactivating effect on adenylate cyclase (\rightarrow **A5**). This results in extremely high levels of intracellular cAMP. When occurring in intestinal cells, this can lead to severe diarrhea (\rightarrow p. 262). Pertussis (whooping cough) toxin and forskolin also lead to an increase in the cytosolic cAMP concentration. Pertussis toxin does this by inhibiting G_i protein and thereby blocking its inhibitory effect on adenylate cyclase, while forskolin directly activates adenylate cyclase. Theophylline and caffeine inhibit the conversion of cAMP to 5'-AMP, which extends the life span of cAMP and prolongs the effect of the messenger.

Certain **ion channels** are regulated by G_s , G_i and other G proteins $(\mathbf{G_0})$ with or without the aid of adenylate cyclase. Some Ca^{2+} channels are activated by G_s proteins and inactivated by G_o proteins, whereas some K^+ channels are activated by G_o proteins and (the $\beta\gamma$ subunits of) G_i proteins $(\rightarrow p.83 \, B)$. $\mathbf{G_{olf}}$ in olfactory receptors, **transducin** in retinal rods $(\rightarrow p.348 \mathrm{ff.})$, and α **-gustducin** in gustatory sensors are also members of the G protein family $(\rightarrow p.338)$.

IP3 and DAG as Second Messengers

As in the case of G_s proteins, once the first messenger using this transduction pathway binds to its receptor outside the cell, the α_q subunit dissociates from the heterotrimeric $\mathbf{G_q}$ protein and activates phospholipase $\mathsf{C-\beta}$ (PLC- β) on the inside of the cell membrane (\rightarrow B1). PLC- β converts phosphatidylinositol 4,5-bisphosphate (PlP_2), to inositol 1,4,5-trisphosphate (PlP_3) and diacylglycerol (DAG). Pl_3 and DAG function as parallel second messengers with different ac-

tions that are exerted either independently or jointly (\rightarrow **B1**).

 IP_3 is a hydrophilic molecule carried via the cytosol to Ca^{2+} stores within the cell (mainly in the endoplasmic reticulum; \rightarrow p. 36). IP_3 binds there to Ca^{2+} channels to open them (\rightarrow B2), leading to an efflux of Ca^{2+} from the intracellular stores into the cytosol. In the cytosol, Ca^{2+} acts as a third messenger that regulates various cell functions, e.g., by interacting with the cAMP signaling chain. Many Ca^{2+} -related activities are mediated by calmodulin, a calciumbinding protein (\rightarrow pp. 36 and 70).

DAG is a lipophilic molecule that remains in the cell membrane and has two main functions:

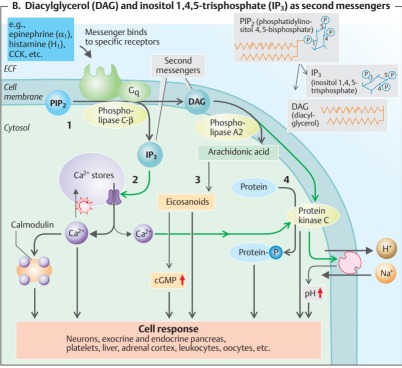
- ◆ DAG is broken down by phospholipase A2 (PLA-2) to yield *arachidonic acid*, a precursor of eicosanoids (→ **B3** and p. 269).
- ◆ DAG activates *protein kinase C* (**PKC**). PKC is Ca^{2+} -dependent (hence the "C") because the Ca^{2+} released by IP₃ (see above) is needed to transfer PKC from the cytosol to the intracellular side of the cell membrane (→ **B4**). Thus activated PKC phosphorylates the serine or threonine residues of many proteins.

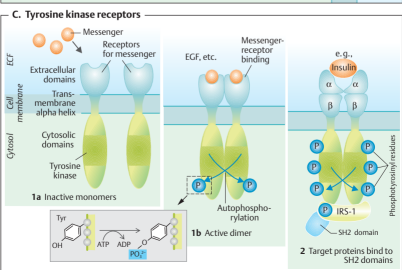
PKC triggers a series of other phosphorylation reactions (high signal amplification) that ultimately lead to the phosphorylation of **MAP kinase** (mitogen-activated protein kinase). It enters the cell nucleus and activates *Elk-1*, a gene-regulating protein. *NF-KB*, another gene-regulating protein, is also released in response to PKC phosphorylation. In addition, PKC activates *Na*/H* antiporters*, thereby raising the cellular **pH**—a stimulus that triggers many other cellular reactions.

IP3 and DAG activating messengers include acetylcholine (M_1 and M_3 cholinoceptors), antidiuretic hormone = vasopressin (V_1 rec.), epinephrine and norepinephrine (α_1 adrenoceptor), bradykinin, CCK, endothelin, gastrin, glutamate (mGLU₁ and mGLU₅ rec.), GRP, histamine (H_1 rec.), leukotrienes, neurotensin, oxytocin and various prostaglandins (FP, TP, and Ep₁ rec.), serotonin = 5-hydroxytryptamine (5-HT₂ rec.), tachykinin, thromboxane A_2 . TRH and TSH induce partial activation.

Deactivation of the signaling cascade can also be achieved through *self-inactivation* of the G proteins involved (GTP cleavage) and phosphatase (see above) as well as by degradation of IP₃.

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Enzyme-Linked Cell Surface Receptors For Messenger Substances

These (*G* protein-independent) receptors, together with their cytosolic domains, act as enzymes that are activated when a messenger binds to the receptor's extracellular domain. There are five classes of these receptors:

- **1. Receptor guanylyl cyclases** convert GTP into the second messenger cGMP, which activates protein kinase G (PKG; see below). The *atriopeptin receptor* belongs to this class.
- **2. Receptor tyrosine kinases** (→ **C**), phosphorylate proteins (of same or different type) at the OH group of their tyrosyl residues. The receptors for *insulin* and various *growth factors* (GF) such as e.g., E[epidermal]GF, PDGF, N[nerve]GF, F[fibroblast]GF, H[hepatocyte]GF, and I[insulin-like]GF-1 belong to this class of receptors.

Signals regarding first messenger binding (e.g., EGF and PDGF) are often transferred inside the cell via binding of two receptors (dimerization; C1a \Rightarrow C1b) and subsequent mutual phosphorylation of their cytosolic domain (autophosphorylation, \rightarrow C1b). The receptor for certain hormones, like insulin and IGF-1, is from the beginning a heterotetramer ($\alpha_2\beta_2$) that undergoes autophosphorylation before phosphorylating another protein (insulin receptor substrate-1, IRS-1) that in turn activates intracellular target proteins containing SH2 domains (\rightarrow C2).

- **3. Receptor serine/threonine kinases**, which like the TGF- β receptor, function similar to kinases in Group 2, the only difference being that they phosphorylate serine or threonine residues of the target protein instead of tyrosine residues (as with PKC; see above).
- **4.** Tyrosine kinase-associated receptors are those where the receptor works in combination with non-receptor tyrosine kinases (chiefly proteins of the Src family) that phosphorylate the target protein. The receptors for STH, prolactin, erythropoietin and numerous cytokines belong to this group.
- **5. Receptor tyrosine phosphatases** remove phosphate groups from tyrosine residues. The CD45 receptor involved in T cell activation belongs to this group.

Hormones with Intracellular Receptors

Steroid hormones (\rightarrow p. 270 ff., yellow areas), calcitriol and thyroid hormones are like other

hormones in that they induce a specific cell response with the difference being that they activate a different type of signaling cascade in the cell. They are *lipid-soluble* substances that freely penetrate the cell membrane.

Steroid hormones bind to their respective *cytoplasmic receptor protein* in the target cell $(\rightarrow \mathbf{D})$. This binding leads to the dissociation of inhibitory proteins (e.g., heat shock protein, HSP) from the receptors. The *hormone-receptor protein complex* (H–R complex) then migrates to the cell nucleus (*translocation*), where it activates (**induces**) or inhibits the transcription of certain genes. The resulting increase or decrease in synthesis of the respective protein (e.g., AIPs; \rightarrow p. 182) is responsible for the actual cell response (\rightarrow **D**).

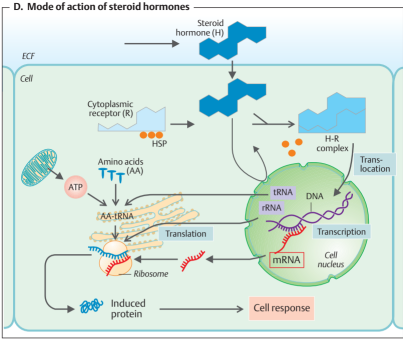
Triiodothyronine (T_3 ; \rightarrow p. 286ff.) and **calcitriol** (\rightarrow p. 292) bind to their respective receptor proteins in the cell nucleus (*nuclear receptors*). These receptors are *hormone-activated transcription factors*. Those of calcitriol can induce the transcription of calcium-binding protein, which plays an important role in interstitial Ca²⁺ absorption (\rightarrow p. 262).

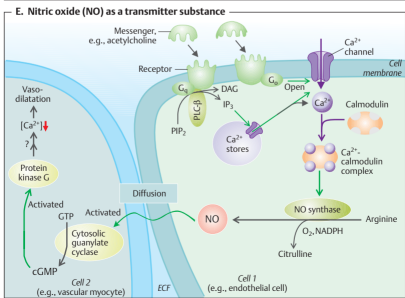
Recent research indicates that steroid hormones and calcitriol also regulate cell function by **non-genomic control mechanisms**.

Nitric Oxide as a Transmitter Substance

In nitrogenergic neurons and endothelial tissues, **nitric (mon)oxide (NO)** is released by $Ca^{2+}/calmodulin-mediated activation of neuronal or endothelial$ **nitric oxide synthase (NOS)** $(<math>\rightarrow$ E). Although NO has a half-life of only a few seconds, it diffuses into neighboring cells (e.g., from endothelium to vascular myocytes) so quickly that it activates **cytoplasmic guanylyl cyclase**, which converts GTP into **cGMP** (\rightarrow E). Acting as a second messenger, cGMP activates *protein kinase G* (**PKG**), which in turn decreases the cytosolic Ca^{2+} concentration $[Ca^{2+}]$ i by still unexplained mechanisms. This ultimately leads to *vasodilatation* (e.g., in coronary arteries).

Penile erections are produced by cGMP-mediated vasodilatation of the deep arteries of the penis (→ p. 308). The erection can be prolonged by drugs that inhibit cGMP-specific phosphodiesterase type 5, thereby delaying the degradation of cGMP (e.g., sildenafil citrate = Viagra®).





Hypothalamic-Pituitary System

In the **hypothalamus**, (1) humoral signals from the periphery (e.g., from circulating cortisol) can be converted to efferent neuronal signals, and (2) afferent neuronal signals can be converted to endocrine messengers (*neurosecretion*).

The first case is possible because the hypothalamus is situated near **circumventricular organs** like the organum vasculosum laminae terminalis (OVLT), subfornical organ, the median eminence of the hypothalamus, and the neurohypophysis. Since there is no blood-brain barrier there, hydrophilic peptide hormones can also enter.

The hypothalamus is closely connected to other parts of the CNS (\rightarrow p. 330). It controls many autonomous regulatory functions and its neuropeptides influence higher brain functions. The hypothalamus is related to the sleeping-waking rhythm (\rightarrow p. 334) and to psychogenic factors. Stress, for example, stimulates the release of cortisol (via CRH, ACTH) and can lead to the cessation of hormone-controlled menstruation (amenorrhea).

Neurosecretion. Hypothalamic neurons synthesize hormones, incorporate them in granules that are transported to the ends of the axons ($axoplasmic transport \rightarrow p.42$), and secrete them into the bloodstream. In this way, oxytocin and ADH are carried from magnocellular hypothalamic nuclei to the neurohypophysis, and RHs and IHs (and ADH) reach the $median \ eminence$ of the hypothalamus (\rightarrow A). The $action \ potential-triggered$ exocytotic release of the hormones into the bloodstream results in Ca²⁺ influx into the nerve endings (\rightarrow p. 50ff.).

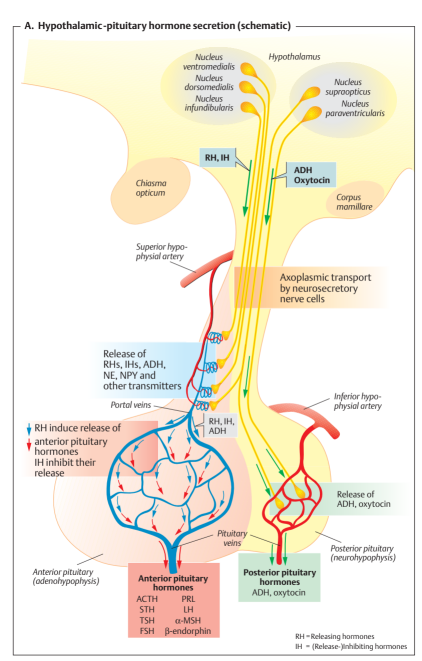
Oxytocin (Ocytocin) and antidiuretic hormone (ADH) are two posterior pituitary hormones that enter the systemic circulation directly. ADH induces water retention in the renal collecting ducts (V_2 -rec.; \rightarrow p. 166) and induces vasoconstriction (endothelial V_1 rec.) by stimulating the secretion of endothelin-1 (\rightarrow p. 212ff.). ADH-bearing neurons also secrete ADH into the portal venous circulation (see below). The ADH and CRH molecules regulate the secretion of ACTH by the adenohypophysis. Oxytocin promotes uterine contractions and milk ejection (\rightarrow p. 304). In nursing

mothers, suckling stimulates nerve endings in the nipples, triggering the secretion of oxytocin (and prolactin, \rightarrow p. 303) via neuro-humoral reflexes.

Releasing hormones (RH) or liberins that stimulate hormone release from the adenohypophysis (Gn-RH, TRH, SRH, CRH; \rightarrow p. 270 ff.) are secreted by hypothalamic neurons into a kind of portal venous system and travel only a short distance to the anterior lobe ($\rightarrow A$). Once in its vascular network, they trigger the release of anterior pituitary hormones into the systemic circulation (\rightarrow A). Some anterior pituitary hormones are regulated by release-inhibiting hormones (IH) or statins, such as SIH and PIH = dopamine (\rightarrow p. 270ff.). Peripheral hormones, ADH (see above) and various neurotransmitters such as neuropeptide Y (NPY). norepinephrine (NE), dopamine, VIP and opioids also help to regulate anterior pituitary functions (\rightarrow p. 272).

The four glandotropic hormones (ACTH. TSH, FSH and LH) and the aglandotropic hormones (prolactin and GH) are secreted from the anterior pituitary $(\rightarrow A)$. The secretion of growth hormone (GH = somatotropic hormone, STH) is subject to control by GH-RH, SIH and IGF-1.GH stimulates protein synthesis (anabolic action) and skeletal growth with the aid of somatomedins (growth factors formed in the liver), which play a role in sulfate uptake by cartilage. Somatomedin C = insulin-like growth factor-1 (IGF-1) inhibits the release of GH by the anterior pituitary via negative feedback control. GH has lipolytic and glycogenolytic actions that are independent of somatomedin activity.

Proopiomelanocortin (POMC) is a peptide precursor not only of ACTH, but (inside or outside the anterior pituitary) also of β-endorphin and α -melanocyte-stimulating hormone (α -MSH = α -melanocortin). β-endorphin has analgesic effects in the CNS and immunomodulatory effects, while α -MSH in the hypothalamus helps to regulate the body weight (\rightarrow p. 230) and stimulates peripheral melanocytes.



Glucose is the *central energy carrier* of the human metabolism. The brain and red blood cells are fully glucose-dependent. The **plasma glucose concentration** (blood sugar level) is determined by the level of glucose *production* and *consumption*.

The following terms are important for proper understanding of carbohydrate metabolism (\rightarrow **A**, **C**):

- **1. Glycolysis** generally refers to the anaerobic conversion of glucose to lactate (\rightarrow p. 72). This occurs in the red blood cells, renal medulla, and skeletal muscles (\rightarrow p. 72). Aerobic oxidation of glucose occurs in the CNS, heart, skeletal muscle and in most other organs.
- **2.** Glycogenesis, i.e., the synthesis of glycogen from glucose (in liver and muscle), facilitates the storage of glucose and helps to maintain a constant plasma glucose concentration. Glycogen stored in a muscle can only be used by that muscle.
- **3. Glycogenolysis** is the breakdown of glycogen to glucose, i.e., the opposite of glycogenesis.
- **4. Gluconeogenesis** is the production of glucose (in liver and renal cortex) from non-sugar molecules such as amino acids (e.g., glutamine), lactate (produced by anaerobic glycolysis in muscles and red cells), and glycerol (from lipolysis).
- **5. Lipolysis** is the breakdown of triacylglycerols into glycerol and free fatty acids.
- **6. Lipogenesis** is the synthesis of triacylglycerols (for storage in fat depots).

Islets of Langerhans in the *pancreas* play a primary role in carbohydrate metabolism. Three cell types (A, B, D) have been identified so far (\rightarrow p. 273 B). 25% of all islet cells are type A (α) cells that produce **glucagon**, 60% are B (β) cells that synthesize **insulin**, and 10% are D (δ) cells that secrete somatostatin (**SIH**). These hormones mutually influence the synthesis and secretion of each other (\rightarrow p. 273 B). Islet cells in the pancreas head synthesize *pancreatic polypeptide*, the physiological function of which is not yet clear. High concentrations of these hormones reach the liver by way of the portal venous circulation.

Function. Pancreatic hormones (1) ensure that ingested food is stored as glycogen and fat (insulin); (2) mobilize energy reserves in response to food deprivation, physical activity or stress (glucagon and the non-pancreatic hormone epinephrine); (3) maintain the plasma

glucose concentration as constant as possible $(\rightarrow A)$; and (4) promote growth.

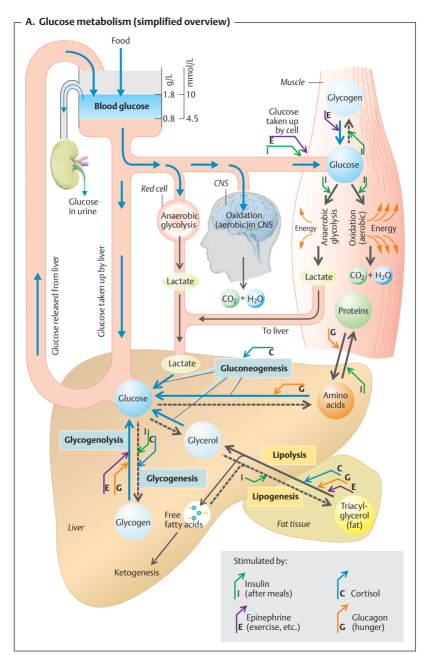
Insulin

Synthesis. Insulin is a 6 kDa peptide (51 amino acids, AA) formed by the C chain cleaved from *proinsulin* (84 AA), the precursor of which is *preproinsulin*, a preprohormone. Insulin contains two peptide chains (A and B) held together by disulfide bridges **Degradation**: Insulin has a half-life of about 5–8 min and is degraded mainly in liver and kidneys.

Secretion. Insulin is secreted in pulsatile bursts, mainly in response to increases in the blood levels of glucose ($\rightarrow B$ right), as follows: plasma glucose $\uparrow \rightarrow$ glucose in B cells $\uparrow \rightarrow$ glucose oxidation $\uparrow \rightarrow$ cytosolic ATP of ATP-gated K+ $\uparrow \rightarrow$ closure channels \rightarrow depolarization \rightarrow opening of voltage-gated Ca^{2+} channels \rightarrow cytosolic $Ca^{2+} \uparrow$. The rising Ca2+ in B cells leads to (a) exocytosis of insulin and (b) re-opening of K+ channels (deactivated by feedback control). Stimulation. Insulin secretion is stimulated mainly during food digestion via acetylcholine (vagus nerve), gastrin, secretin, GIP (\rightarrow p. 234) and GLP-1 (glucagon-like peptide = enteroglucagon), a peptide that dissociates from intestinal proglucagon, Certain amino acids (especially arginine and leucine), free fatty acids, many pituitary hormones and some steroid hormones also increase insulin secretion. Inhibition. Epinephrine and norepinephrine (α_2 adrenoceptors; $\rightarrow A$, B), SIH (\rightarrow p. 273 B) and the neuropeptide galanin inhibit insulin secretion. When hypoglycemia occurs due, e.g., to fasting or prolonged physical exercise, the low blood glucose concentration is sensed by central chemosensors for glucose, leading to reflex activation of the sympathetic nervous system.

The **insulin receptor** is a heterotetramer ($\alpha_2\beta_2$) consisting of two extracellular α subunits and two transmembranous β subunits. The α subunits bind the hormone. Once the β subunits are autophosphorylated, they act as receptor tyrosine kinases that phosphorylate insulin receptor substrate-1 (IRS-1). Intracellular proteins with SH2 domains are phosphorylated by IRS-1 and pass on the signal (\rightarrow p. 277 C3).

Action of insulin (\rightarrow A, B, C). Insulin has *anabolic* and *lipogenic* effects, and promotes the *storage of glucose*, especially in the liver, where it activates enzymes that *promote glycolysis*



and glycogenesis and suppresses those involved in gluconeogenesis. Insulin also increases the number of GLUT-4 uniporters in skeletal myocytes. All these actions serve to lower the plasma glucose concentration (which increases after food ingestion). About two-thirds of the glucose absorbed by the intestines after a meal (postprandial) is temporarily stored and kept ready for mobilization (via glucagon) during the interdigestive phase. This provides a relatively constant supply of glucose for the glucose-dependent CNS and vital organs in absence of food ingestion. Insulin increases the storage of amino acids (AA) in the form of proteins, especially in the skeletal muscles (anabolism). In addition, it promotes growth, inhibits extrahepatic lipolysis ($\rightarrow p, 257, D$) and affects K^+ distribution (\rightarrow p. 180).

Hypoglycemia develops when the insulin concentration is too high. Glucose levels of < 2 mmol/L (35 mg/dL) produce glucose deficiencies in the brain, which can lead to coma and *hypoglycemic shock*.

The excessive intake of carbohydrates can overload glycogen stores. The liver therefore starts to convert glucose into fatty acids, which are transported to and stored in fatty tissues in the form of triacylalycerols (\rightarrow p. 257 D).

Diabetes mellitus (DM). One type of DM is insulin-dependent diabetes mellitus (IDDM), or type 1 DM, which is caused by an insulin deficiency. Another type is non-insulin-dependent DM (NIDDM), or type 2 DM, which is caused by the decreased efficacy of insulin and sometimes occurs even in conjunction with increased insulin concentrations. DM is characterized by an abnormally high plasma glucose concentration (hyperglycemia), which leads to glucosuria ($\rightarrow p. 158$). Large quantities of fatty acids are liberated since lipolysis is no longer inhibited (\rightarrow p. 257 D). The fatty acids can be used to produce energy via acetylcoenzyme A (acetyl-CoA); however, this leads to the formation of acetoacetic acid, acetone (ketosis), and β-oxybutyric acid (metabolic acidosis, \rightarrow p. 142). Because hepatic fat synthesis is insulin-independent and since so many fatty acids are available, the liver begins to store triacylglycerols, resulting in the development of fatty liver.

Glucagon, Somatostatin and Somatotropin

Glucagon released from A cells is a peptide hormone (29 AA) derived from *proglucagon* (glicentin). The granules in which glucagon is stored are secreted by exocytosis. Secretion is **stimulated** by AA from digested proteins (especially alanine and arginine) as well as by hy-

poglycemia (e.g., due to fasting, prolonged physical exercise; \rightarrow **B**) and sympathetic impulses (via β_2 adrenoceptors; \rightarrow **A**). Glucagon secretion is **inhibited** by glucose and SIH (\rightarrow p. 273, B) as well as by high plasma concentrations of free fatty acids.

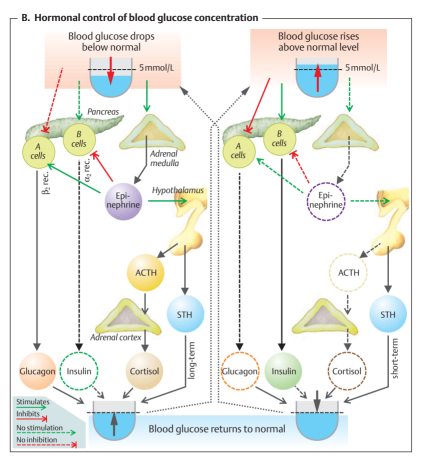
The **actions** of glucagon (\rightarrow **A, B, C**) (via cAMP; \rightarrow p. 274) mainly antagonize those of insulin. Glucagon maintains a normal *blood glucose level between meals* and during phases of increased glucose consumption to ensure a constant energy supply. It does this (a) by increasing glycogenolysis (in liver not muscle) and (b) by stimulating gluconeogenesis from lactate, AA (protein degradation = catabolism) and glycerol (from *lipolysis*).

Increased plasma concentrations of **amino acids** (AA) stimulate insulin secretion which would lead to hypoglycemia without the simultaneous ingestion of glucose. Hypoglycemia normally does not occur, however, since AA also stimulate the release of glucagon, which increases the blood glucose concentration. Glucagon also stimulates gluconeogenesis from AA, so some of the AA are used for energy production. In order to increase protein levels in patients, glucose must therefore be administered simultaneously with therapeutic doses of AA to prevent their metabolic degradation.

Somatostatin (SIH). Like insulin, SIH stored in D cells (**SIH 14** has 14 AA) is released in response to increased plasma concentrations of glucose and arginine (i.e., after a meal). Through *paracrine* pathways (via G_i -linked receptors), SIH inhibits the release of insulin (\rightarrow p. 273, B). Therefore, SIH inhibits not only the release of gastrin, which promotes digestion (\rightarrow p. 243, B3), but also interrupts the insulin-related storage of nutrients. SIH also inhibits glucagon secretion (\rightarrow p. 273 B). This effect does not occur in the presence of a glucose deficiency because of the release of catecholamines that *decrease* SIH secretion.

Somatotropin (STH) = growth hormone (GH). The short-term effects of GH are similar to those of insulin; its action is mediated by somatomedins (\rightarrow p. 280). In the long-term, GH increases the blood glucose concentration and promotes growth.

The effects of **glucocorticoids** on carbohydrate metabolism are illustrated on plate **C** and explained on p. 296.



─ C. Hormonal effects on carbohydrate and fat metabolism ————————————————————————————————————							
	Hormone Function	Insulin Satiated ← Buffer	Glucagon	Epinephrine	Cortisol Supply		
Glucose Uptake by cell Glycolysis Gluconeogene	esis (liver)	Muscle,	-	+ Muscle + +	Muscle,		
Glycogen Synthesis	Lysis	Liver, muscle	Liver	Liver, muscle	Liver		
Fat Synthesis	Lysis	Liver, fat	Fat	Fat	Fat		

The thyroid gland contains spherical follicles $(50-500\,\mu\mathrm{m}\,\mathrm{in}\,\mathrm{diameter})$. Follicle cells synthesize the two iodine-containing thyroid hormones **thyroxine** (T_4 , tetraiodothyronine) and **triiodothyronine** (T_3). T_3 and T_4 are bound to the glycoprotein *thyroglobulin* (\rightarrow B2) and stored in the colloid of the follicles (\rightarrow A1, B1). The synthesis and release of T_3/T_4 is controlled by the *thyroliberin* (= *thyrotropin-releasing hormone*, **TRH**)—*thyrotropin* (**TSH**) *axis* (\rightarrow A, and p. 270ff.). T_3 and T_4 influence physical growth, maturation and metabolism. The *parafollicular cells* (C *cells*) of the thyroid gland synthesize *calcitonin* (\rightarrow p. 292).

Thyroglobulin, a dimeric glycoprotein (660 kDa) is synthesized in the thyroid cells. TSH stimulates the transcription of the thyroglobulin gene. Thyroglobulin is stored in vesicles and released into the colloid by exocytosis (\rightarrow **B1** and p. 30).

Iodine uptake. The iodine needed for hormone synthesis is taken up from the bloodstream as iodide (I⁻). It enters thyroid cells through secondary active transport by the Na⁺-I⁻ symport carrier (NIS) and is concentrated in the cell ca. 25 times as compared to the plasma (\rightarrow B2). Via cAMP, TSH increases the transport capacity of basolateral I⁻ uptake up to 250 times. Other anions competitively inhibit I⁻ uptake; e.g., ClO₄⁻, SCN⁻ and NO₂⁻.

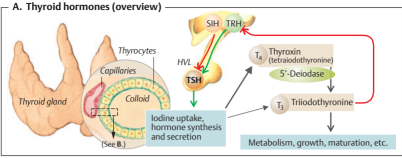
Hormone synthesis. I- ions are continuously transported from the intracellular I- pool to the apical (colloidal) side of the cell by a I-/Cl- antiporter, called pendrin, which is stimulated by **TSH**. With the aid of thyroid peroxidase (TPO) and an H₂O₂ generator, they are oxidized to elementary I₂⁰ along the microvilli on the colloid side of the cell membrane. With the help of TPO, the I⁰ reacts with about 20 of the 144 tvrosyl residues of thyroglobulin $(\rightarrow C)$. The phenol ring of the tyrosyl residue is thereby iodinated at the 3 and/or 5 position, yielding a protein chain containing either diiodotyrosine (DIT) residues and/or monoiodotyrosine (MIT) residues. These steps of synthesis are stimulated by TSH (via IP₃) and inhibited by thiouracil, thiocyanate, glutathione, and other reducing substances. The structure of the thyroglobulin molecule allows the iodinated tyrosyl residues to react with each other in the

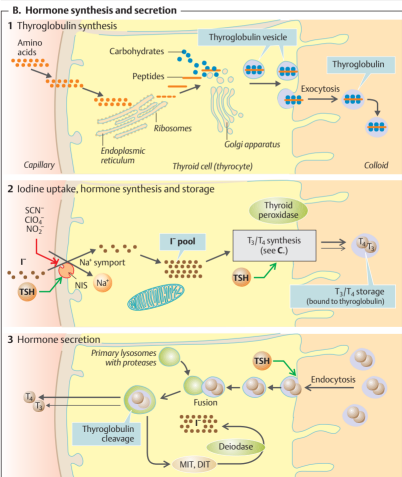
thyrocolloid. The phenol ring of one DIT (or MIT) molecule links with another DIT molecule (ether bridges). The resulting thyroglobulin chain now contains tetraiodothyronine residues ($\rightarrow \mathbf{C}$). These are the storage form of T_4 and T_3 .

TSH also stimulates T_3 and T_4 secretion. The iodinated thyroglobulin in thyrocolloid are reabsorbed by the cells via endocytosis (\rightarrow B3, C). The endosomes fuse with primary lysosomes to form phagolysosomes in which thyroglobulin is hydrolyzed by proteases. This leads to the release of T_3 and T_4 (ca. 0.2 and 1–3 mol per mol of thyroglobulin, respectively). T_3 and T_4 are then secreted into the bloodstream (\rightarrow B3). With the aid of *deiodase*, I^- meanwhile is split from concomitantly released MIT and DIT and becomes reavailable for synthesis.

Control of T₃/T₄ secretion. TSH secretion by the anterior pituitary is stimulated by TRH, a hypothalamic tripeptide (→p.280) and inhibited by **somatostatin (SIH)** (\rightarrow **A** and p. 270). The effect of TRH is modified by T4 in the plasma. As observed with other target cells, the T₄ taken up by the thyrotropic cells of the anterior pituitary is converted to T₃ by 5'-deiodase, T₃ reduces the density of TRH receptors in the pituitary gland and inhibits TRH secretion by the hypothalamus. The secretion of TSH and consequently of T₃ and T₄ therefore decreases (negative feedback circuit). In neonates, cold seems to stimulate the release of TRH via neuronal pathways (thermoregulation, \rightarrow p. 224). TSH is a heterodimer (26 kDa) consisting of an α subunit (identical to that of LH and FSH) and a β subunit. TSH controls all thyroid gland functions, including the uptake of I-, the synthesis and secretion of T_3 and $T_4 (\rightarrow A-C)$, the blood flow and growth of the thyroid gland.

Goiter (struma) is characterized by diffuse or nodular enlargement of the thyroid gland. Diffuse goiter can occur due to an iodine deficiency, resulting in T_3/T_4 deficits that ultimately lead to increased secretion of T5H. Chronic elevation of T5H leads to the proliferation of follicle cells, resulting in goiter development (*hyperplastic goiter*). This prompts an increase in T_3/T_4 synthesis, which sometimes normalizes the plasma concentrations of T_3/T_4 (*euthyroid goiter*). This type of goiter often persists even after the iodide deficiency is rectified.





Hypothyroidism occurs when TSH-driven thyroid enlargement is no longer able to compensate for the T_3/T_4 deficiency (*hypothyroid goiter*). This type of goiter can also occur due to a congenital disturbance of T_3/T_4 synthesis (see below) or thyroid inflammation. **Hyperthyroidism** occurs when a thyroid tumor (*hot node*) or diffuse struma (e.g., in *Grave's disease*) results in the overproduction of T_3/T_4 , independent of TSH. In the latter case, an autoantibody against the TSH receptor binds to the TSH receptor. Its effects mimic those of TSH, i.e., it stimulates T_3/T_4 synthesis and secretion.

 T_3/T_4 transport. T_3 and T_4 occur at a ratio of 1:40 in the plasma, where >99% of them (mainly T_4) are bound to plasma proteins: *thyroxine-binding globulin* (TBG), thyroxine-binding prealbumin (TBPA), and *serum albumin*. TBG transports two-thirds of the T_4 in the blood, while TBPA and serum albumin transport the rest. Less than 0.3% of the total T_3/T_4 in blood occurs in an unbound (free) form, although only the unbound molecules have an effect on the target cells. Certain drugs split T_3 and T_4 from protein bonds, resulting in increased plasma concentrations of the free hormones.

Potency of T₃/**T**₄. T₃ is 3–8 times more potent than T₄ and acts more rapidly (half-life of T₃ is 1 day, that of T₄ 7 days). Only ca. 20% of all circulating T₃ originate from the thyroid; the other 80% are produced by the liver, kidneys, and other target cells that cleave iodide from T₄. The conversion of T₄ to T₃ is catalyzed by microsomal **5**′-**deiodase**, which removes iodine from the 5′ position on the outer ring (\rightarrow **D**). T₃ is therefore the more potent hormone, while T₄ is mainly ascribed a *storage function* in plasma.

The inactive form of T_3 called **reverse T**₃ (**rT**₃) is produced from T_4 when the iodine is split from the inner ring with the aid of a 5- (not 5'-)deiodase. Approximately equal amounts of T_3 and rT_3 are normally produced in the periphery (ca. 25 μ g/day). When a person fasts, the resulting *inhibition of 5'-deiodase* decreases T_3 synthesis (to save energy, see below) while rT_3 synthesis increases. *Pituitary 5'-deiodase* is not inhibited, so TSH secretion (unwanted in this case) is suppressed by the negative feedback.

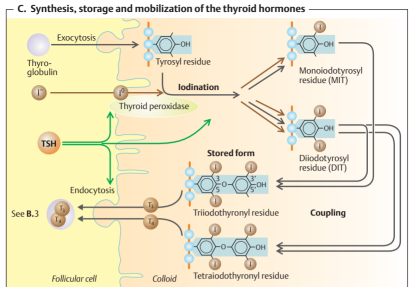
 T_3/T_4 receptors are hormone-sensitive transcription factors located in the cell nuclei. Hormone-receptor complexes bind to regulator

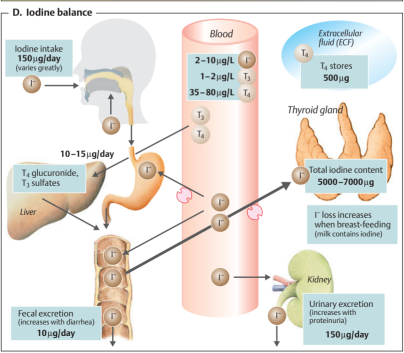
proteins of certain genes in the nuclei and influence their transcription.

The actions of T_3/T_4 are numerous and mainly involve the intermediate metabolism. The thyroid hormones increase the number of mitochondria and its cristae, increase Na+-K+-ATPase activity and modulate the cholesterol metabolism. This results in an increase in energy turnover and a corresponding rise in O₂ consumption and heat production. T3 also specifically stimulates heat production by increasing the expression of the uncoupling protein thermogenin in brown fat (\rightarrow p. 222). T₃ also influences the efficacy of other hormones. Insulin, glucagon, GH and epinephrine lose their energy turnover-increasing effect in hypothyroidism, whereas the sensitivity to epinephrine increases (heart rate increases, etc.) in hyperthyroidism. T3 is thought to increase the density of β-adrenoceptors. T₃ also stimulates growth and maturation, especially of the brain and bones.

Cretinism occurs due to neonatal T_3/T_4 deficiencies and is marked by growth and maturation disorders (dwarfism, delayed sexual development, etc.) and central nervous disorders (intelligence deficits, seizures, etc.). The administration of thyroid hormones in the first six months of life can prevent or reduce some of these abnormalities.

lodine metabolism (\rightarrow **D**). Iodine circulates in the blood as either (1) inorganic Γ (2–10 μg/L), (2) organic non-hormonal iodine (traces) and (3) protein-bound iodine (PBI) within Γ_3 and Γ_4 (35–80 μg iodine/L). The average daily requirement of iodine is ca. 150 μg; larger quantities are required in fever and hyperthyroidism (ca. 250–500 μg/day). Iodine excreted from the body must be replaced by the diet (\rightarrow **D**). Sea salt, seafood, and cereals grown in iodine-rich soil are rich in iodine. Iodized salt is often used to supplement iodine deficiencies in the diet. Since iodine passes into the breast milk, nursing mothers have a higher daily requirement of iodine (ca. 200 μg/day).





Calcium and Phosphate Metabolism

Calcium, particularly ionized calcium (Ca2+). plays a central role in the regulation of numerous cell functions (\rightarrow pp. 36, 62ff., 192, 276). Calcium accounts for 2% of the body weight. Ca. 99% of the calcium occurs in bone while 1% is dissolved in body fluids. The total calcium conc. in serum is normally 2.1-2.6 mmol/L. Ca. 50% of it is **free Ca²⁺** (1.1–1.3 mmol/L) while ca. 10% is bound in complexes and 40% is bound to proteins (mainly albumin: \rightarrow p. 178). Calcium protein binding increases as the pH of the blood rises since the number of Ca²⁺ binding sites on protein molecules also rises with the pH. The Ca2+ conc. accordingly decreases in alkalosis and rises in acidosis (by about 0.21 mmol/L Ca²⁺ per pH unit). Alkalosis (e.g., due to hyperventilation) and hypocalcemia (see below) can therefore lead to tetany.

The **calcium metabolism** is tightly regulated to ensure a balanced intake and excretion of Ca^{2+} (\rightarrow A). The *dietary intake of Ca²⁺* provides around 12–35 mmol of Ca^{2+} each day (1 mmol = 2 mEq = 40 mg). Milk, cheese, eggs and "hard" water are particularly rich in Ca^{2+} . When calcium homeostasis is maintained, most of the ingested Ca^{2+} is excreted in the feces, while the remainder is excreted in the urine (\rightarrow p. 178). When a calcium deficiency exists, up to 90% of the ingested Ca^{2+} is absorbed by the intestinal tract (\rightarrow A and p. 262).

Pregnant and nursing mothers have higher Ca²⁺ requirements because they must also supply the fetus or newborn infant with calcium. The fetus receives ca. 625 mmol/day of Ca²⁺ via the placenta, and nursed infants receive up to 2000 mmol/day via the breast milk. In both cases, the Ca²⁺ is used for bone formation. Thus, many women develop a Ca²⁺ deficiency during or after pregnancy.

Phosphate metabolism is closely related to calcium metabolism but is less tightly controlled. The daily intake of phosphate is about 1.4 g; 0.9 g of intake is absorbed and usually excreted by the kidneys (\rightarrow p. 178). The **phosphate concentration** in serum normally ranges from 0.8–1.4 mmol/L.

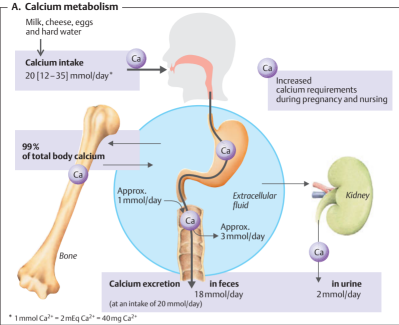
Calcium phosphate salts are sparingly soluble. When the product of Ca²⁺ conc. times phosphate conc. (*solubility product*) exceeds a certain threshold, calcium phosphate starts to

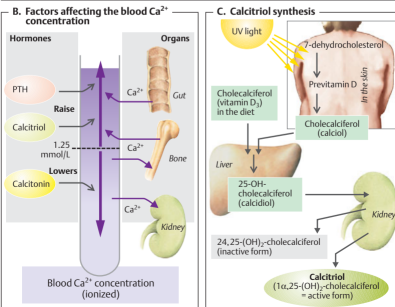
precipitate in solutions, and the deposition of calcium phosphate salts occurs. The salts are chiefly deposited in the bone, but can also precipitate in other organs. The infusion of phosphate leads to a decrease in the serum calcium concentration since calcium phosphate accumulates in bone. Conversely, hypophosphatemia leads to hypercalcemia (Ca²⁺ is released from bone).

Hormonal control. Calcium and phosphate homeostasis is predominantly regulated by *parathyroid hormone* and *calcitriol*, but also by calcitonin to a lesser degree. These hormones mainly affect three organs: the *intestines*, the *kidneys* and the *bone* (\rightarrow **B** and **D**).

Parathyrin or parathyroid hormone (PTH) is a peptide hormone (84 AA) secreted by the parathyroid glands. Ca2+ sensors in cells of the parathyroid glands regulate PTH synthesis and secretion in response to changes in the plasma concentration of ionized Ca^{2+} ($\rightarrow p.36$). More PTH is secreted into the bloodstream whenever the Ca2+ conc. falls below normal (hypocalcemia). Inversely, PTH secretion decreases when the Ca^{2+} level rises ($\rightarrow \mathbf{D}$, left panel). The primary function of PTH is to nor**malize decreased Ca²⁺** conc. in the blood $(\rightarrow D)$. This is accomplished as follows: (1) PTH activates osteoclasts, resulting in bone breakdown and the release of Ca2+ (and phosphate) from the bone; (2) PTH accelerates the final step of calcitriol synthesis in the kidney, resulting in increased reabsorption of Ca2+ from the aut: (3) in the kidney, PTH increases calcitriol synthesis and Ca²⁺ reabsorption, which is particularly important due to the increased Ca2+ supply resulting from actions (1) and (2). PTH also inhibits renal phosphate reabsorption (\rightarrow p. 178), resulting in hypophosphatemia. This, in turn, stimulates the release of Ca2+ from the bone or prevents the precipitation of calcium phosphate in tissue (solubility product; see above).

Hypocalcemia occurs due to a deficiency (*hypoparathyroidism*) or lack of efficiency (*pseudohypoparathyroidism*) of PTH, which can destabilize the resting potential enough to produce *muscle spasms* and *tetany*. These deficiencies can also lead to a secondary calcitriol deficiency. An excess of PTH (*hyperparathyroidism*) and malignant osteolysis overpower the Ca²⁺ control mechanisms, leading to **hypercalcemia**. The long-term elevation of Ca²⁺ results in cal-





cium deposition (e.g., in the kidneys). Ca²⁺ conc. exceeding 3.5 mmol/L lead to coma, renal insufficiency and cardiac arrhythmias.

Calcitonin (CT), or thyrocalcitonin, is a peptide hormone (32). It is mainly synthesized in the parafollicular cells (C cells) of the thyroid gland, which also contain Ca2+ sensors $(\rightarrow p.36)$. Hypercalcemia increases the plasma calcitonin conc. (\rightarrow **D**, **right** panel), whereas calcitonin can no longer be detected when the calcium conc. [Ca2+] falls below 2 mmol/L. Calcitonin normalizes elevated serum Ca2+ conc. mainly by acting on **bone**. Osteoclast activity is inhibited by calcitonin (and stimulated by PTH). Calcitonin therefore increases the uptake of Ca²⁺ by the bone—at least temporarily $(\rightarrow D5)$. Some gastrointestinal hormones accelerate calcitonin secretion, thereby enhancing the postprandial absorption of Ca²⁺ by bone. These effects (and perhaps the restraining effect of calcitonin on digestive activities) function to prevent postprandial hypercalcemia and the (unwanted) inhibition of PTH secretion and increased renal excretion of the just absorbed Ca2+. Calcitonin also acts on the kidneys (\rightarrow **D6**).

Calcitriol $(1,25-(OH)_2$ -cholecalciferol) is a lipophilic, steroid-like hormone synthesized as follows (\rightarrow **C**): **Cholecalciferol** (vitamin D₃) is produced from hepatic 7-dehydrocholesterol in the **skin** via an intermediate product (previtamin D) in response to UV light (sun, tanning lamps). Both substances bind to vitamin D-binding protein (DBP) in the blood, but cholecalciferol is preferentially transported because of its higher affinity. Previtamin D therefore remains in the skin for a while after UV light exposure (short-term storage). Calcidiol (25-OH-cholecalciferol) and calcitriol bind to DBP. An estrogen-dependent rise in DBP synthesis occurs during pregnancy.

Cholecalciferol (vitamin D₃) is administered to compensate for inadequate UV exposure. The recommended daily dosage in children is approximately 400 units = $10\,\mu g$; adults receive half this amount. Plant-derived **ergocalciferol** (vitamin D₂) is equally effective as animal-derived vitamin D₃. The following actions apply for both forms.

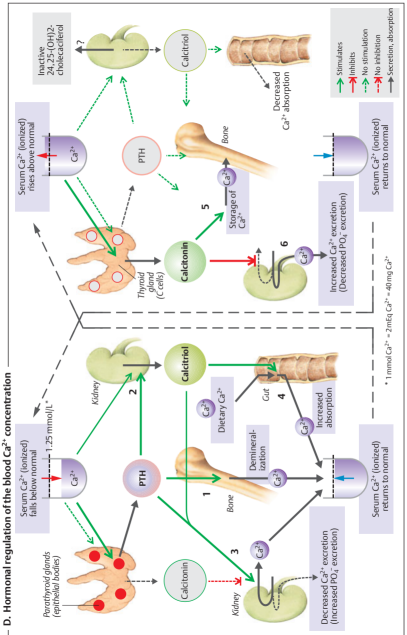
Cholecalciferol is converted to **calcidiol** (25-OH-cholecalciferol) in the liver. Vitamin D is mainly *stored as calcidiol* because the plasma

conc. of calcidiol is $25 \,\mu g/L$, and its half-life is 15 days. **Calcitriol** $(1,25\text{-}(OH)_2\text{-}cholecalciferol)$, the hormonally active form, is mainly synthesized in the **kidneys** (\rightarrow C), but also in the placenta. The plasma conc. of **calcitriol is regulated** by renal $1\text{-}\alpha\text{-hydroxylase}$ (final step of synthesis) and by 24-hydroxylase, an enzyme that deactivates calcitriol.

The **calcitriol concentration rises** in response to *hypocalcemia*-related **PTA** secretion (\rightarrow **D2**), to *phosphate deficiency* and to *prolactin* (lactation). All three inhibit 24-hydroxylase and activate 1- α -hydroxylase. It **decreases** due to several negative feedback loops, i.e. due to the fact that calcitriol (a) directly inhibits 1- α -hydroxylase, (b) inhibits parathyroid hormone secretion, and (c) normalizes the (decreased) plasma conc. of Ca²⁺ and phosphate by increasing the intestinal absorption of Ca²⁺ and phosphate (see below). Calcium and phosphate inhibit 1- α -hydroxylase, while phosphate activates 24-hydroxylase.

Target organs. Calcitriol's primary target is the **gut**, but it also acts on the *bone*, *kidneys*, *placenta*, *mammary glands*, *hair follicles*, *skin* etc. It binds with its nuclear receptor and induces the expression of calcium-binding protein and Ca^{2+} -ATPase (\rightarrow pp. 278 and 36). Calcitriol has also genomic effects. Calcitriol *increases the intestinal absorption of* Ca^{2+} (\rightarrow **D4**) and *promotes mineralization of the bone*, but an **excess of calcitriol** leads to *decalcification* of the bone, an effect heightened by PTH. Calcitriol also increases the transport of Ca^{2+} and phosphate at the kidney (\rightarrow p. 178), placenta and mammary glands.

In transitory **hypocalcemia**, the bones act as a temporary Ca^{2+} buffer $(\rightarrow \mathbf{D})$ until the Ca^{2+} deficit has been balanced by a calcitriol-mediated increase in Ca^{2+} absorption from the gut. If too little calcitriol is available, skeletal demineralization will lead to **osteomalacia** in adults and **rickets** in children. **Vitamin D deficiencies** are caused by inadequate dietary intake, reduced absorption (fat maldigestion), insufficient UV light exposure, and/or reduced 1-cahydroxylation (renal insufficiency). Skeletal demineralization mostly occurs due to the prolonged increase in parathyroid hormone secretion associated with chronic hypocalcemia (*compensatory hyperparathyroidism*).



Biosynthesis of Steroid Hormones

Cholesterol is the precursor of steroid hormones $(\rightarrow A)$. Cholesterol is mainly synthesized in the **liver**. It arises from acetylcoenzyme A (acetyl-CoA) via a number of intermediates (e.g., squalene, lanosterol) and is transported to the endocrine glands by lipoproteins $(\rightarrow p. 256)$. Cholesterol can be synthesized de novo also in the adrenal cortex, but not in the placenta $(\rightarrow p. 304)$. Since only small quantities of steroid hormones are stored in the organs of origin, i.e., the adrenal cortex, ovaries, testes and placenta $(\rightarrow p. 304)$, they must be synthesized from the cellular cholesterol pool as needed.

Cholesterol contains 27 carbon atoms. Preq**nenolone** (21 C atoms; \rightarrow **A, a**), the precursor of steroid hormones, arises from cholesterol via numerous intermediates. Pregnenolone also vields **progesterone** $(\rightarrow A. b)$, which is not only a potent hormone itself (female sex hormone; \rightarrow p. 298ff.), but can act as the precursor of all other steroid hormones, i.e., (1) the adrenocortical hormones with 21 carbon atoms ($\rightarrow A$, yellow and orange fields); (2) male sex hormones (androgens, 19 carbon atoms) synthesized in the testes ($\rightarrow p.306$), ovaries and adrenal cortex ($\rightarrow A$, green and blue fields); and (3) female sex hormones (estrogens, 18 carbon atoms; \rightarrow p. 29 B ff.) synthe sized in the ovaries (\rightarrow A, red zones).

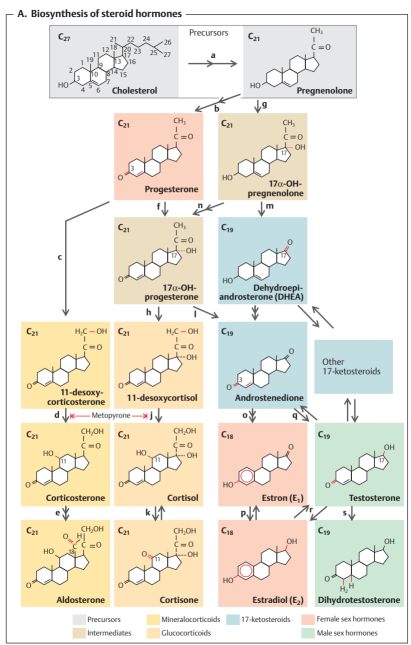
The precursors for steroid hormone synthesis are present in all steroid hormone glands. The type of hormone produced and the site of hormone synthesis depend on (1) the type of receptors available for the superordinate control hormones (ACTH, FSH, LH, etc.) and (2) the dominant enzyme responsible for changing the structure of the steroid molecule in the hormone-producing cells of the gland in question. The adrenal cortex contains 11-, 17- and 21-hydroxylases-enzymes that introduce an OH group at position C21, C17 or C11, respectively, of the steroid molecule ($\rightarrow A$, top left panel for numerical order). Hydroxylation at C21 (\rightarrow **A, c**)—as realized in the glomerular zone of the adrenal cortex-makes the steroid insensitive to the effects of 17-hydroxylase. As a result, only mineralocorticoids like corticosterone and aldosterone (A, $\mathbf{d} \Rightarrow \mathbf{e}$; see also

p. 182) can be synthesized. Initial hydroxylation at C17 (\rightarrow A, f or g) results in the synthesis of **glucocorticoids**—realized mainly in the fascicular zone of the adrenal cortex (\rightarrow A, h \Rightarrow j \Rightarrow k)—and 17-ketosteroids, steroids with a keto group at C17 (\rightarrow A, I and m). Glucocorticoids and 17-ketosteroids can therefore be synthesized from 17 α -hydroxypregnenolone without the aid of progesterone (\rightarrow A, n \Rightarrow h \Rightarrow j).

The **estrogens** $(\rightarrow p.302)$ *estrone* and *estradiol* can be directly or indirectly synthesized from 17-ketosteroids $(\rightarrow A, o \Rightarrow p)$; they are produced indirectly by way of testosterone $(\rightarrow A, q \Rightarrow r \Rightarrow p)$. The true active substance of certain target cells for androgens (e.g., in the prostate) is either *dihydrotestosterone* or *estradiol*; both are synthesized from testosterone $(\rightarrow A, s \text{ and } A, r, \text{ respectively})$.

17-ketosteroids are synthesized by the gonads (testes and ovaries) and adrenal cortex. Since they are found in the urine, the *metyrapone test* of pituitary function is used to assess the *ACTH reserve* based on urinary 17-ketosteroids levels. ACTH secretion is normally subject to feedback control by glucocorticoids (→ p. 296). Metyrapone inhibits 11-hydroxylase activity (→ **A**, **d** and **j**), which leaves ACTH unsuppressed in healthy subjects. Urinary 17-ketosteroid levels should therefore increase after metyrapone administration. An abnormality of ACTH secretion can be assumed when this does not occur in patients with a healthy adrenal cortex.

Degradation of steroid hormones occurs mainly in the liver. Their OH groups are usually linked to sulfate or glucuronic acid molecules and are ultimately excreted in the bile or urine $(\rightarrow pp. 160, 183 \text{ and } 250)$. The chief urinary metabolite of the estrogens is estriol, while that of the gestagens (mainly progesterone and 17α-hydroxyprogesterone) is pregnanediol (→ p. 304). Pregnanediol levels in urine can be measured to confirm or exclude pregnancy test (pregnanediol test). Chronically increased estrogen levels due, for example, to decreased estrogen degradation secondary to liver damage, can lead to breast development (gynecomastia) in the male, among other things. For normal estrogen ranges, see table on p. 302.



Adrenal Cortex and Glucocorticoid Synthesis

The **mineralocortico(stero)ids** aldosterone, corticosterone and 11-desoxycorticosterone (\rightarrow pp. 182ff. and 294) are synthesized in the glomerular zone of the **adrenal cortex** (\rightarrow **A1**), whereas the **glucocortico(stero)ids** cortisol (hydrocortisone) and cortisone (\rightarrow p. 294, small quantities) are synthesized in the *fascicular zone* (\rightarrow **A2**). **Androgens** are synthesized in the *reticular zone* of the adrenal cortex (\rightarrow **A3**). One of the androgens is *dehydroepiandrosterone* (DHEA), which is used (partly in its sulfated form, DHEA-S) to synthesize various sex hormones in other tissues (\rightarrow p. 304).

Cortisol transport. Most of the plasma cortisol is bound to *transcortin*, or *cortisol-binding globulin* (*CBG*), a specific transport protein with a high-affinity binding site for cortisol. Cortisol is released in response to conformational changes of CBG due to inflammation etc.

CRH and **ACTH** regulate cortisol synthesis and secretion (\rightarrow **A4**, **A5**; see also p. 270). ACTH ensures also structural preservation of the adrenal cortex and supplies cortisol precursors, e.g., by forming cholesterol from its esters, by de novo synthesis of cholesterol and by converting it to progesterone and 17 α -hydroxyprogesterone (\rightarrow pp. 256 and 294). *ACTH secretion* is stimulated by *CRH* and epinephrine and inhibited (negative feedback control) by *cortisol* with or without the aid of CRH (\rightarrow **A**; see also p. 273 A).

A **circadian rhythm** of CRH secretion and thus of ACTH and cortisol secretion can be observed. The peak secretion is in the morning (\rightarrow **B**, mean values). Continuous hormone conc. sampling at short intervals have shown that ACTH and cortisol are secreted in 2–3-hour *episodes* (\rightarrow **B**).

Receptor proteins (\rightarrow p. 278) for glucocorticoids can be found in virtually every cell. Glucocorticoids are vital hormones that exert numerous **effects**, the most important of which are listed below.

Carbohydrate and amino acid (AA) metabolism (see also pp. 283 A and 285 C): Cortisol uses AA derived from protein degradation to increase the plasma glucose concentration (gluconeogenesis), which can lead to the so-

called steroid diabetes in extreme cases. Thus, cortisol has a catabolic effect (degrades proteins) that results in the increased excretion of urea.

Cardiovascular function: Glucocorticoids increase myocardial contractility and vasoconstriction due to enhancement of catecholamine effects (\rightarrow pp. 194 and 214). These are described as *permissive effects* of cortisol. Cortisol increases the synthesis of epinephrine in the adrenal medulla (\rightarrow **A6**) and of angiotensinogen in the liver (\rightarrow p. 184).

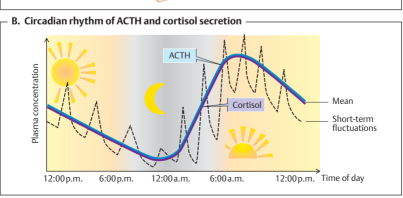
Especially when administered at high doses, glucocorticoids induce anti-inflammatory and anti-allergic effects because they stabilize lymphokine synthesis and histamine release (\rightarrow p. 100). On the other hand, interleukin-1, interleukin-2 and TNF- α (e.g., in severe infection) leads to increased secretion of CRH and high cortisol conc. (see below).

Renal function: Glucocorticoids delay the *excretion* of water and help to maintain a normal glomerular filtration rate. They can react also with aldosterone receptors but are converted to cortisone by 11β -hydroxysteroid oxidoreductase in aldosterone target cells. Normal cortisol conc. are therefore ineffective at the aldosterone receptor. High conc., however, have the same effect as aldosterone (\rightarrow p. 182).

Gastric function: Glucocorticoids weaken the protective mechanisms of the gastric mucosa. Thus, high-dose glucocorticoids or stress (see below) increase the risk of gastric ulcers (→ p. 242).

Cerebral function: High glucocorticoid conc. change hypothalamic $(\rightarrow A)$ and electrical brain activity (EEG) and lead to psychic abnormalities.

Stress: Physical or mental stress increases cortisol secretion as a result of increased CRH secretion and increased sympathetic tone $(\rightarrow A)$. Many of the aforementioned effects of cortisol therefore play a role in the body's response to stress (activation of energy metabolism, increase in cardiac performance, etc.). In severe physical (e.g., sepsis) or mental stress (e.g., depression), the cortisol plasma conc. remains at a very high level (up to 10 times the normal value) throughout the day.



Oogenesis and the Menstrual Cycle

Oogenesis. The development of the female gametes (ova) extends from the oogonium stage to the primary oocyte stage (in the primordial follicle), starting long before birth. Oogenesis therefore occurs much sooner than the corresponding stages of spermatogenesis (\rightarrow p. 306). The fetal phase of oogenesis is completed by the first week of gestation; these oocytes remain latent until puberty. In the sexually mature female, a fertilizable ovum develops in the graafian follicles approximately every 28 days.

Menstrual cycle. After the start of sexual maturation, a woman starts to secrete the following hormones in a cyclic (approx.) 28-day rhythm $(\rightarrow A1, A2)$. Gonadoliberin (= gonadotropin-releasing hormone, Gn-RH) and dopamine (PIH) are secreted by the hypothalamus, Folliclestimulating hormone (FSH), luteinizing hormone (LH) and prolactin (PRL) are released by the anterior pituitary. Progesterone, estrogens (chiefly estradiol, E2) and inhibin are secreted by the ovaries. Gn-RH controls the pulsatile secretion of FSH and LH (\rightarrow p. 300), which in turn regulate the secretion of estradiol and progesterone. The female sex functions are controlled by the periodic release of hormones, the purpose of which is to produce a fertilizable egg in the ovaries each month $(\rightarrow A4)$ and produce an environment suitable for sperm reception (fertilization) and implantation of the fertilized ovum (*nidation*) (\rightarrow **A5**). This cyclic activity is reflected by the monthly menses (menstruation) which, by definition, marks the start of the menstrual cycle.

Girls in Central Europe usually have their first menstrual period (menarche) around the age of 13. By about age 40, the cycle becomes increasingly irregular over a period of up to 10 years (climacteric) as the end of the reproductive period nears. The last menses (menopause) generally occurs around the age of 48–52.

A menstrual cycle can last 21–35 days. The second half of the cycle (**luteal phase** = secretory phase) usually lasts 14 days, while the first half (**follicular phase** = proliferative phase) lasts 7–21 days. **Ovulation** separates the two phases (→ A). If the cycle length varies by more than 2–3 days, ovulation generally does not occur. Such *anovulatory cycles* account for 20% of all cycles in healthy females.

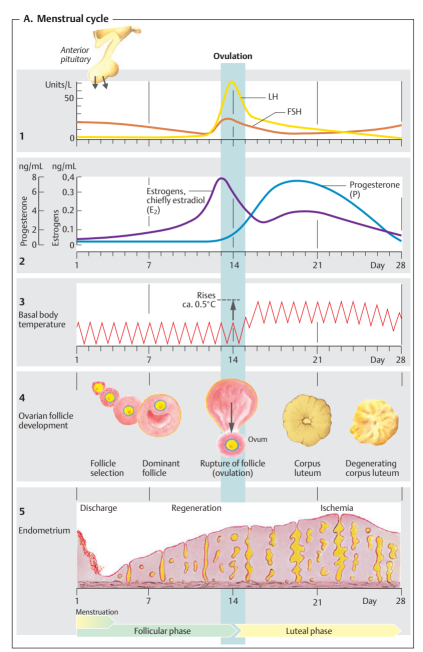
In addition to general changes in the body and mood, the following changes occur in the ovaries, uterus and cervix during the menstrual cycle (\rightarrow A):

Day 1: Start of menstruation (lasting about 2–6 days).

Days 1–14 (variable, see above): The follicular phase starts on the first day of menstruation. The endometrium thickens to become prepared for the implantation of the fertilized ovum during the luteal phase (\rightarrow A5), and about 20 ovarian follicles mature under the influence of FSH. One of these becomes the *dominant follicle*, which produces increasing quantities of *estrogens* (\rightarrow A4 and p. 300). The small cervical os is blocked by a viscous mucous plug.

Day 14 (variable, see above): Ovulation. The amount of *estrogens* produced by the follicle increases rapidly between day 12 and 13 (→ A2). The increased secretion of LH in response to higher levels of estrogen leads to ovulation (→ A1, A4; see also p. 300). The *basal body temperature* (measured on an empty stomach before rising in the morning) rises about 0.5°C about 1–2 days later and remains elevated until the end of the cycle (→ A3). This temperature rise generally indicates that ovulation has occurred. During ovulation, the *cervical mucus* is less viscous (it can be stretched into long threads—*spinnbarkeit*) and the cervical os opens slightly to allow the sperm to enter.

Days 14-28: The luteal phase is characterized by the development of a corpus luteum $(\rightarrow A4)$, which secretes progesterone, $(\rightarrow A2)$; an increase in mucoid secretion from the uterine glands also occurs (\rightarrow **A5**). The endometrium is most responsive to progesterone around the 22nd day of the cycle, which is when nidation should occur if the ovum has been fertilized. Otherwise, progesterone and estrogens now inhibit Gn-RH secretion $(\rightarrow p.300)$, resulting in degeneration of the corpus luteum. The subsequent rapid decrease in the plasma concentrations of estrogens and progesterone (\rightarrow A2) results in constriction of endometrial blood vessels and ischemia. This ultimately leads to the breakdown and discharge of the uterine lining and to bleeding, i.e., menstruation (\rightarrow **A5**).



In sexually mature women, **gonadoliberin** or **gonadotropin-releasing hormone** (**Gn-RH**) is secreted in one-minute pulses every 60–90 min in response to signals from various neurotransmitters. This, in turn, induces the pulsatile secretion of FSH and LH from the anterior pituitary. If the rhythm of Gn-RH secretion is much faster or continuous, less FSH and LH will be secreted, which can result in infertility. The LH:FSH secretion ratio changes during the course of the menstrual cycle. Their release must be therefore subject to additional factors besides Gn-RH.

The secretion of LH and FSH is, for example, subject to **central nervous effects** (psychogenic factors, stress) mediated by various **transmitters** circulating in the portal blood in the hypothalamic region, e.g., norepinephrine (NE) and neuropeptide Y (NPY) as well as by ovarian hormones, i.e., by *estrogens* (estrone, estradiol, estriol, etc.), *progesterone* and *inhibin*. Ovarian hormones affect Gn-RH secretion indirectly by stimulating central nerve cells that activate Gn-RH-secreting neurons by way of neurotransmitters such as norepinephrine and NPY and inhibit Gn-RH secretion by way of GABA and opioids.

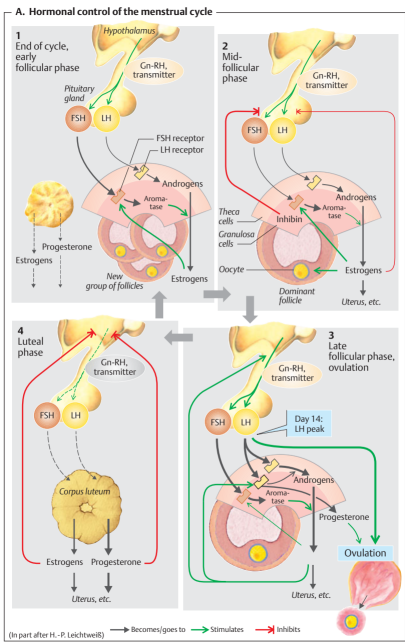
FSH production again increases toward the end of the luteal phase (\rightarrow p. 299, A1). In the early follicular phase (→A1), FSH induces the proliferation of the stratum granulosum in about 20 follicles and stimulates the secretion of aromatase in their granulosa cells. Aromatase catalyzes the conversion of the androgens testosterone and androstenedione to estradiol (E₂) and estrone (E₁) (\rightarrow p. 295 A, steps r and o). Estrogens are synthesized in theca cells and absorbed by granulosa cells. Although relatively small amounts of LH are secreted (\rightarrow A1 and p. 299 A1), this is enough to activate theca cell-based enzymes (17β-hydroxysteroid dehydrogenase and C17/C20lyase) that help to produce the androgens needed for estrogen synthesis. The folliclebased estrogens increase their own FSH receptor density. The follicle with the highest estrogen content is therefore the most sensitive to FSH. This loop has a self-amplifying effect, and the follicle in question is selected as the dominant follicle around the 6th day of the cycle $(\rightarrow A2)$. In the mid-follicular phase, estrogens restrict FSH and LH secretion (via negative feedback control and with the aid of *inhibin*; \rightarrow A2) but later stimulate LH receptor production in granulosa cells. These cells now also start to produce progesterone (start of *luteinization*), which is absorbed by the theca cells (\rightarrow A3) and used as precursor for further increase in androgen synthesis (\rightarrow p. 295 A, steps f and f).

Inhibin and estrogens secreted by the dominant follicle increasingly inhibit FSH secretion, thereby decreasing the estrogen production in other follicles. This leads to an androgen build-up in and apoptosis of the unselected follicles.

Increasing quantities of LH and FSH are released in the late follicular phase ($\rightarrow A3$), causing a sharp rise in their plasma concentrations. The FSH peak occurring around day 13 of the cycle induces the first meiotic division of the ovum. Estrogens increase the LH secretion (mainly via the hypothalamus), resulting in the increased production of androgens and estrogens (positive feedback) and a rapid rise in the LH conc. (LH surge). The LH peak occurs around day 14 (\rightarrow A2). The follicle ruptures and discharges its ovum about 10 hours later (ovulation). Ovulation does not take place if the LH surge does not occur or is too slow. Pregnancy is not possible in the absence of ovulation.

Luteal phase (\rightarrow **A4**). LH, FSH and estrogens transform the ovarian follicle into a corpus luteum. It actively secretes large quantities of progesterone (progestational hormone). marking the beginning of the luteal phase $(\rightarrow A)$. Estrogens and progesterone now inhibit the secretion of FSH and LH directly and indirectly (e.g., through inhibition of Gn-RH; see above), causing a rapid drop in their plasma conc. This negative feedback leads to a marked drop in the plasma conc. of estrogens and progesterone towards the end of the menstrual cycle (approx. day 26), thereby triggering the menses (\rightarrow p. 299, A2). FSH secretion starts to rise just before the start of menstruation $(\rightarrow A4)$.

Combined administration of estrogens and gestagens during the first half of the menstrual cycle prevents ovulation. Since ovulation does not occur, pregnancy cannot take place. Most **contraceptives** work according to this principle.



Estrogens

Estrogens are steroid hormones with 18 carbon atoms. Estrogens are primarily synthesized from the 17-ketosteroid *androstene-dione*, but testosterone can also be a precursor (\rightarrow p. 295 A). The *ovaries*, (granulosa and theca cells), *placenta* (\rightarrow p. 304), *adrenal cortex*, and in *Leydig's cells* (interstitial cells) of the *testes* (\rightarrow p. 306) are the physiological **sites of estrogen synthesis.**. In some target cells for testosterone, it must first be converted to estradiol to become active.

Estradiol (E₂) is the most potent estrogen (E). The potencies of **estrone (E₁)** and **estriol (E₃)** are relatively low in comparison (E₂: E₁: E₃ = 10:5:1). Most estrogens (and testosterone) circulating in the blood are bound to *sex hormone-binding globulin (SHBG)*. Estriol (E₃) is the chief degradation product of estradiol (E₂).

Oral administration of estradiol has virtually no effect because almost all of the hormone is removed from the bloodstream during its first pass through the liver. Therefore, other estrogens (with a different chemical structure) must be used for effective **oral estrogen therapy**.

Actions. Although estrogens play a role in the development of female sex characteristics, they are not nearly as important as the androgens for male sexual development (\rightarrow p. 306). The preparatory action of estrogen is often required for optimal progesterone effects (e.g., in the uterus; see below). Other important effects of estrogens in human females are as follows.

 Menstrual cycle. Estrogens accelerate maturation of the ovarian follicle during the menstrual cycle (\rightarrow p. 298 and table). In the uterus. estrogen promotes the proliferation (thickening) of the endometrium and increases uterine muscle contraction. In the vagina, estrogen thickens the mucosal lining, leading to the increased discharge of glycogen-containing epithelial cells. The liberated glycogen is used for an increased production of lactic acid by Döderlein's bacillus. This lowers the vaginal pH to 3.5-5.5, thereby reducing the risk of vaginal infection. In the cervix, the mucous plug sealing the cervical os functions as a barrier that prevents sperm from entering the uterus. Estrogens change the consistency of the cervical mucus, making it more conducive to sperm penetration and survival, especially around the time of ovulation.

- Fertilization. In the female body, estrogens prepare the sperm to penetrate and fertilize the ovum (capacitation) and regulate the speed at which the ovum travels in the fallopian tube.
- Extragonadal effects of estrogen. During puberty, estrogens stimulate breast development, induces changes in the vagina and in the distribution of subcutaneous fat, and (together with androgens) stimulate the growth of pubic and axillary hair. Since estrogens increase the coagulability of the blood, the administration of estrogens (e.g., in contraceptives) increases the risk of thrombosis and leads renal salt and water retention. Estrogens slow longitudinal bone growth, accelerate epiphyseal closure (in men and women) and increase osteoblast activity. Estrogen deficiencies in menopause consequently lead to the loss of bone mass (osteoporosis). Estrogens induce a decrease in LDL and a rise in VLDL and HDL concentrations $(\rightarrow p, 254ff.)$, which is why arteriosclerosis is less common in premenopausal women than in men. Estrogen also makes the skin thinner and softer, reduces the sebaceous glands, and increases fat deposits in subcutaneous tissue. Lastly, estrogen influences a number of central nervous functions, e.g., sexual response, social behavior, and mood.

Plasma concentrations of estradiol and progesterone (ng/mL)

Phase	Estradiol	Progesteron
Women		
Early follicular phase	0.06	0.3
Mid- and late follicular	$0.1 \Rightarrow 0.4$	1.0
phase		
Ovulation	0.4	2.0
Mid-luteal phase	0.2	8-16
Pregnancy	7–14	$40 \Rightarrow 130$
Day 1 after parturition		20
Men	0.05	0.3

Progesterone

Progesterone, the most potent progestational (pregnancy-sustaining) hormone, is a steroid hormone (21 C atoms) synthesized from

cholesterol via pregnenolone (\rightarrow p. 295). It is produced in the corpus luteum, ovarian follicles and placenta (\rightarrow p. 304) of the female, and in the adrenal cortex of males and females. Like cortisol, most circulating progesterone is bound to cortisol-binding globulin (CBG = transcortin). Like estradiol (E₂), most progesterone is broken down during its first pass through the liver, so oral doses of progesterone are almost completely ineffective. *Pregnane-diol* is the most important degradation product of progesterone.

Actions of progesterone. The main functions of progesterone are to prepare the female genital tract for implantation and maturation of the fertilized ovum and to sustain pregnancy (→ see table). Progesterone counteracts many of the effects induced by estrogens, but various effects of progesterone depend on the preparatory activity or simultaneous action of estrogens. During the follicular phase, for example, estrogens increases the density of progesterone receptors, while simultaneous estrogen activity is needed to induce mammary growth (see below).

- The **uterus** is the chief target organ of progesterone. Once estrogen induces endometrial thickening, progesterone stimulates growth of the uterine muscle (myometrium), restructures the endometrial glands ($\rightarrow p.298$), alters the blood supply to the endometrium, and changes the glycogen content. This represents the transformation from a proliferative endometrium to a secretory endometrium, with a peak occurring around day 22 of the cycle. Progesterone later plays an important role in the potential implantation (nidation) of the fertilized ovum because it reduces myometrial activity (important during pregnancy), narrows the cervical os, and changes the consistency of the cervical mucous plug so that it becomes virtually impregnable to sperm.
- ◆ Progesterone inhibits the release of LH during the luteal phase. The administration of gestagens like progesterone during the follicular phase inhibits ovulation. Together with its effects on the cervix (see above) and its inhibitory effect on capacitation (→p.302), progesterone can therefore have a contraceptive effect ("mini pill").
- High levels of progesterone have an anesthetic effect on the central nervous system.
 Progesterone also increases the susceptibility

to epileptic fits and exerts thermogenic action, i.e., it raises the basal body temperature (→ p. 298). In addition, a decrease in the progesterone concentration is also believed to be responsible for the mood changes and depression observed before menstruation (*premenstrual syndrome*, *PMS*) and after pregnancy (*postpartum depression*).

• In the **kidneys**, progesterone slightly inhibits the effects aldosterone, thereby inducing increased NaCl excretion.

Prolactin and Oxytocin

The secretion of prolactin (PRL) is inhibited by prolactin-inhibiting hormone (PIH = dopamine) and stimulated by thyroliberin (TRH) (\rightarrow p. 270). Prolactin increases the hypothalamic secretion of PIH in both men and women (negative feedback control). Conversely, estradiol (E2) and progesterone inhibit PIH secretion (indirectly via transmitters, as observed with Gn-RH: see above). Consequently, prolactin secretion rises significantly during the second half of the menstrual cycle and during pregnancy. Prolactin (together with estrogens, progesterone, glucocorticoids and insulin) stimulate breast enlargement during pregnancy and lactogenesis after parturition. In breast-feeding, stimulation of the nerve endings in the nipples by the suckling infant stimulates the secretion of prolactin (lactation reflex). This also increases release of oxytocin which triggers milk ejection and increases uterine contractions, thereby increasing lochia discharge after birth. When the mother stops breast-feeding, the prolactin levels drop, leading to the rapid stoppage of milk production.

Hyperprolactinemia. Stress and certain drugs inhibit the secretion of PIH, causing an increase in prolactin secretion. *Hypothyroidism* (\rightarrow p. 288) can also lead to hyperprolactinemia, because the associated increase in TRH stimulates the release of prolactin. Hyperprolactinemia inhibits ovulation and leads to *galactorrhea*, i.e., the secretion of milk irrespective of pregnancy. Some women utilize the anti-ovulatory effect of nursing as a natural method of birth control, which is often but not always effective.

Hormonal Control of Pregnancy and Birth

Beside its other functions, the **placenta** produces most of the hormones needed during pregnancy (\rightarrow p. 220). *Ovarian hormones* also play a role, especially at the start of pregnancy (\rightarrow **A**).

Placental hormones. The primary hormones produced by the placenta are human chorionic gonadotropin (hCG), corticotropin-releasing hormone (CRH). estrogens. progesterone. human placental lactogen (hPL), and proopiomelanocortin (POMC; \rightarrow p. 280). hCG is the predominant hormone during the first trimester of pregnancy (3-month period calculated from the beginning of the last menses). Maternal conc. of hPL and CRH-controlled estrogens rise sharply during the third trimester $(\rightarrow B)$. Placental hormones are distributed to mother and fetus. Because of the close connection between maternal, fetal and placental hormone synthesis, they are jointly referred to as the **fetoplacental unit** $(\rightarrow A)$.

Human chorionic gonadotropin (hCG) (a) stimulates the synthesis of steroids like DHEA and DHEA-S by the fetal adrenal cortex (see below); (b) suppresses follicle maturation in the maternal ovaries, and (c) maintains the production of progesterone and estrogen in the corpus luteum (\rightarrow A1) until the 6th week of gestation, i.e., until the placenta is able to produce sufficient quantities of the hormones.

Most **pregnancy tests** are based on the fact that hCG is detectable in the urine about 6–8 days after conception. Since the levels of estrogen and progesterone greatly increase during pregnancy (see table on p. 302), larger quantities of these hormones and their metabolites *estriol* and *pregnanediol* are excreted in the urine. Therefore, their conc. can also be measured to test for pregnancy.

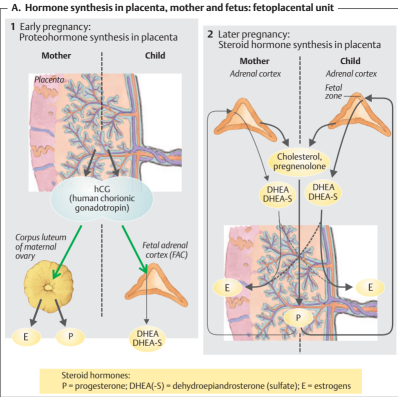
In contrast to other endocrine organs, the placenta has to receive the appropriate precursors (cholesterol or androgens, \rightarrow p. 294) from the maternal and fetal adrenal cortex, respectively, before it can synthesize progesterone and estrogen (\rightarrow A2). The fetal adrenal cortex (FAC) is sometimes larger than the kidneys and consists of a *fetal zone* and an *adult zone*. The placenta takes up cholesterol and pregnenolone and uses them to synthesize progesterone. It is transported to the fetal zone of

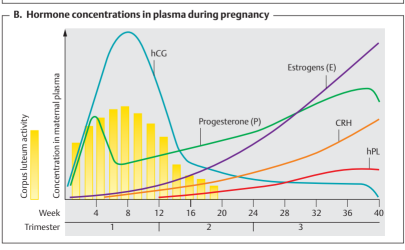
the FAC, where it is converted to dehydroepiandrosterone (**DHEA**) and dehydroepiandrosterone sulfate (**DHEA-S**). DHEA and DHEA-S pass to the **placenta**, where they are used for estrogen synthesis. Progesterone is converted to testosterone in the testes of the male fetus.

Human placental lactogen (hPL = human chorionic somatomammotropin, HCS) levels rise steadily during pregnancy. Like prolactin (\rightarrow p. 303), hPL stimulates mammary enlargement and lactogenesis in particular and, like GH (\rightarrow p. 280), stimulates physical growth and development in general. hPL also seems to increase maternal plasma glucose conc.

Corticotropin-releasing hormone (CRH) secreted by the placenta seems to play a key role in the hormonal regulation of birth. The plasma levels of maternal CRH increase exponentially from the 12th week of gestation on. This rise is more rapid in premature births and slower in post-term births. In other words. the rate at which the CRH concentration rises seems to determine the duration of the pregnancy. Placental CRH stimulates the release of ACTH by the fetal pituitary, resulting in increased cortisol production in the adult zone of FAC; this again stimulates the release of CRH (positive feedback). CRH also stimulates lung development and the production of DHEA and DHEA-S in the fetal zone of FAC.

The maternal estrogen conc. rises sharply towards the **end of the pregnancy**, thereby counteracting the actions of progesterone, including its pregnancy-sustaining effect. Estrogens induce oxytocin receptors (\rightarrow p. 303), α_1 adrenoceptors (\rightarrow p. 84ff.), and gap junctions in the uterine musculature ($\rightarrow p$, 16ff.), and uterine cells are depolarized. All these effects increase the responsiveness of the uterine musculature. The simultaneous increase in progesterone synthesis triggers the production of collagenases that soften the taut cervix. Stretch receptors in the uterus respond to the increase in size and movement of the fetus. Nerve fibers relay these signals to the hypothalamus, which responds by secreting larger quantities of oxytocin which, in turn, increases uterine contractions (positive feedback). The gap junctions conduct the spontaneous impulses from individual pacemaker cells in the fundus across the entire myometrium at a rate of approximately $2 \text{ cm/s} (\rightarrow \text{p. } 70)$.





Androgens and Testicular Function

Androgens (male sex hormones) are steroid hormones with 19 C atoms. This group includes potent hormones like *testosterone* (**T**) and 5α -dihydrotestosterone (**DHT**) and less potent 17-ketosteroids (17-KS) such as DHEA (\rightarrow p. 294). In males, up to 95% of testosterone is synthesized by the *testes* (\rightarrow **A2**) and 5% by the *adrenal cortex* (\rightarrow **AI**). The *ovaries* and adrenal cortex synthesize testosterone in females. The plasma testosterone conc. in males is about 15 times higher than in females, but decreases with age. Up to 98% of testosterone circulating in blood is bound to plasma proteins (albumin and sex hormone-binding globulin, SHBG; \rightarrow **A2**).

The testes secrete also small quantities of **DHT** and **estradiol** (**E**₂). Larger quantities of DHT (via 5-\alpha-reductase) and estradiol are synthesized from testosterone (via *aromatase*) by their respective target cells. A portion of this supply is released into the plasma. DHT and testosterone bind to the same intracellular receptor. Estradiol influences many functions in the male, e.g., epiphyseal cartilage and ejaculate formation and pituitary and hypothalamic activity.

Testosterone secretion is **regulated** by luteinizing hormone (=LH, also called ICSH, \rightarrow p. 269), the pulsatile secretion of which is controlled by Gn-RH at 1.5- to 2-hourly intervals, as in the female. LH stimulates the release of testosterone from Leydig's cells (interstitial cells) in the testes (\rightarrow A2), whereas testosterone and estradiol inhibit LH and Gn-RH secretion (negative feedback).

Gn-RH also induces the release of **FSH**, which stimulates the secretion of **inhibin** and induces the expression of *androgen-binding protein* (**ABP**) in Sertoli cells of the testes (\rightarrow **A3**). Testosterone cannot induce spermatogenesis without the help of ABP (see below). FSH also induces the formation of LH receptors in the interstitial cells of Leydig. Testosterone, DHT, estradiol and inhibin inhibit the secretion of FSH (negative feedback; \rightarrow **A**). **Activin**, the physiological significance of which is still unclear, inhibits FSH secretion.

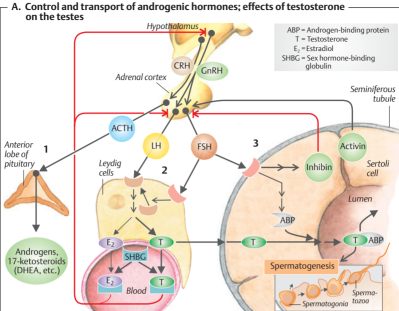
Apart from the important **effects of testosterone** on male sexual differentiation, spermatogenesis and sperm growth as well as on the functions of the genitalia, prostate and seminal vesicle (see below), testosterone also induces the *secondary sex characteristics* that occur in males around the time of puberty, i.e., body hair distribution, physique, laryngeal size (voice change), acne, etc. In addition, testosterone is necessary for normal sex drive (*libido*), procreative capacity (*fertility*) and coital capacity (*potentia coeundi*) in the male. Testosterone also stimulates *hematopoiesis* and has *anabolic properties*, leading to increased muscle mass in males. It also has central nervous effects and can influence behavior—e.g., cause aggressiveness.

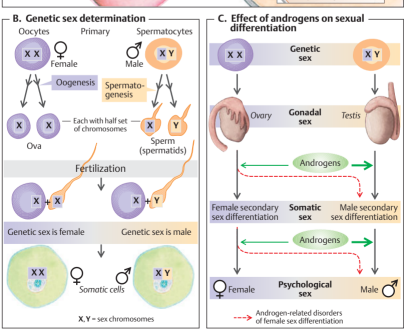
Sexual development and differentiation. The genetic $sex (\rightarrow B)$ determines the development of the sex-specific gonads (gamete-producing glands). The germ cells (spermatogonia; see below) then migrate into the gonads. The somatic sex is female when the subsequent somatic sex development and sex differentiation occurs in the absence of testosterone ($\rightarrow C$). Male development requires the presence of testosterone in both steps ($\rightarrow C$) with or without the aid of additional factors (e.g., calcitonin gene-related peptide, CGRP?) in certain stages of development (e.g., descent of testes into scrotum). High conc. of testosterone, either natural or synthetic (anabolic steroids), lead to masculinization (virilization) of the female ($\rightarrow C$).

Testicular function. *Spermatogenesis* occurs in several stages in the testes (target organ of testosterone) and produces *sperm* (*spermatozoa*) (→ A3). Sperm are produced in the *seminiferous tubules* (total length, ca. 300 m), the epithelium of which consists of *germ cells* and *Sertoli cells* that support and nourish the spermatogenic cells. The seminiferous tubules are strictly separated from other testicular tissues by a *blood–testis barrier*. The testosterone required for sperm maturation and semen production (→ p. 308) must be bound to androgen-binding protein (ABP) to cross the barrier.

Spermatogonia (→B) are primitive sex cells. At puberty, a spermatogonium divides mitotically to form two daughter cells. One of these is kept as a lifetime stem cell reservoir (in contrast to oogonia in the female; →p. 298). The other undergoes several divisions to form a primary spermatocyte. It undergoes a first meiotic division (MD1) to produce two secondary spermatocytes, each of which undergoes a second meiotic division (MD2), producing a total of four spermatids, which ultimately differentiate into spermatozoa. After MD1, the spermatocytes have a single (haploid) set of chromosomes.

306





Sexual Response, Intercourse and Fertilization

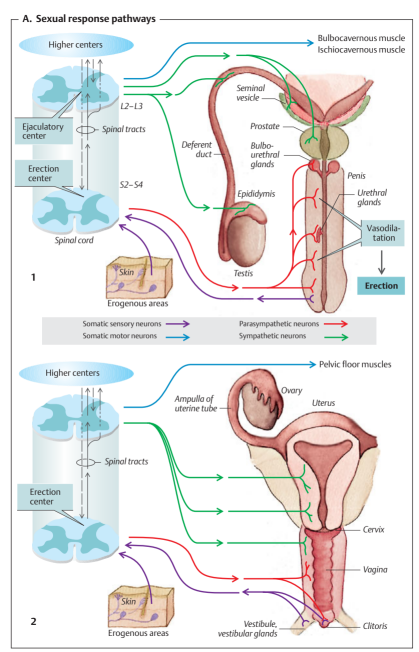
Sexual response in the male $(\rightarrow A1)$. Impulses from tactile receptors on the skin in the genital region (especially the glans penis) and other parts of the body (erogenous areas) are transmitted to the erection center in the sacral spinal cord (S2-S4), which conducts them to parasympathetic neurons of the pelvic splanchnic nerves, thereby triggering sexual arousal. Sexual arousal is decisively influenced by stimulatory or inhibitory impulses from the brain triggered by sensual perceptions, imagination and other factors. Via nitric oxide (→ p. 278), efferent impulses lead to dilatation of deep penile artery branches (helicine arteries) in the erectile body (corpus cavernosum). while the veins are compressed to restrict the drainage of blood. The resulting high pressure (> 1000 mmHg) in the erectile body causes the penis to stiffen and rise (erection). The ejaculatory center in the spinal cord (L2-L3) is activated when arousal reaches a certain threshold (\rightarrow A2). Immediately prior to ejaculation, efferent sympathetic impulses trigger the partial evacuation of the prostate gland and the emission of semen from the vas deferens to the posterior part of the urethra. This triggers the eiaculation reflex and is accompanied by orgasm, the apex of sexual excitement. The effects of orgasm can be felt throughout the entire body, which is reflected by perspiration and an increase in respiratory rate, heart rate, blood pressure, and skeletal muscle tone. During ejaculation, the internal sphincter muscle closes off the urinary bladder while the vas deferens, seminal vesicles and bulbocavernous and ischiocavernous muscles contract rhythmically to propel the semen out of the urethra.

Semen. The fluid expelled during ejaculation (2–6 mL) contains 35–200 million sperm in a nutrient fluid (*seminal plasma*) composed of various substances, such as prostaglandins (from the prostate) that stimulate uterine contraction. Once semen enters the vagina during **intercourse**, the alkaline seminal plasma increase the vaginal pH to increase sperm motility. At least one sperm cell must reach the ovum for fertilization to occur.

Sexual response in the female (\rightarrow A2). Due to impulses similar to those in the male, the erectile tissues of the clitoris and vestibule of the vagina engorge with blood during the erection phase. Sexual arousal triggers the release of secretions from glands in the labia minora and transudates from the vaginal wall, both of which lubricate the vagina, and the nipples become erect. On continued stimulation, afferent impulses are transmitted to the lumbar spinal cord, where sympathetic impulses trigger orgasm (climax). The vaginal walls contract rhythmically (orgasmic cuff), the vagina lengthens and widens, and the uterus becomes erect, thereby creating a space for the semen. The cervical os also widens and remains open for about a half an hour after orgasm. Uterine contractions begin shortly after orgasm (and are probably induced locally by oxytocin). Although the accompanying physical reactions are similar to those in the male (see above). there is a wide range of variation in the orgasmic phase of the female. Erection and orgasm are not essential for conception.

Fertilization. The fusion of sperm and egg usually occurs in the *ampulla* of the fallopian tube. Only a small percentage of the sperm expelled during ejaculation ($1000-10\,000$ out of 10^7 to 10^8 sperm) reach the fallopian tubes (*sperm ascension*). To do so, the sperm must penetrate the mucous plug sealing the cervix, which also acts as a sperm reservoir for a few days. In the time required for them to reach the ampullary portion of the fallopian tube (about 5 hours), the sperm must undergo certain changes to be able to fertilize an ovum; this is referred to as **capacitation** (\rightarrow p. 302).

After ovulation (\rightarrow p. 298ff.) the ovum enters the tube to the uterus (oviduct) via the abdominal cavity. When a sperm makes contact with the egg (via chemotaxis), species-specific sperm-binding receptors on the ovum are exposed and the proteolytic enzyme acrosin is thereby activated (acrosomal reaction). Acrosin allows the sperm to penetrate the cells surrounding the egg (corona radiata). The sperm bind to receptors on the envelope surrounding the ovum (zona pellucida) and enters the eaa. The membranes of both cells then fuse. The ovum now undergoes a second meiotic division, which concludes the act of fertilization. Rapid proteolytic changes in the receptors on the ovum (zona pellucida reaction) prevent other sperm from entering the egg. Fertilization usually takes place on the first day after intercourse and is only possible within 24 hours after ovulation.



Central Nervous System and Senses

Central Nervous System

The brain and spinal cord make up the central nervous system (CNS) (\rightarrow A). The spinal cord is divided into similar segments, but is 30% shorter than the spinal column. The spinal nerves exit the spinal canal at the level of their respective vertebrae and contains the afferent somatic and visceral fibers of the dorsal root, which project to the spinal cord, and the efferent somatic (and partly autonomic) fibers of the anterior root, which project to the periphery. Thus, a nerve is a bundle of nerve fibers that has different functions and conducts impulses in different directions (\rightarrow p. 42).

Spinal cord (\rightarrow **A**). Viewed in cross-section, the spinal cord has a dark, butterfly-shaped inner area (*gray matter*) surrounded by a lighter outer area (*white matter*). The four wings of the gray matter are called *horns* (cross-section) or columns (longitudinal section). The *anterior horn* contains motoneurons (projecting to the muscles), the *posterior horn* contains interneurons. The cell bodies of most afferent fibers lie within the **spinal ganglion** outside the spinal cord. The white matter contains the axons of ascending and descending tracts.

Brain (\rightarrow **D**). The main parts of the brain are the *medulla oblongata* (\rightarrow **D7**) *pons* (\rightarrow **D6**), *mesencephalon* (\rightarrow **D5**), *cerebellum* (\rightarrow **E**), *diencephalon* and *telencephalon* (\rightarrow **E**). The medulla, pons and mesencephalon are collectively called the **brain stem**. It is structurally similar to the spinal cord but also contains cell bodies (nuclei) of *cranial nerves*, neurons controlling *respiration* and *circulation* (\rightarrow **p**). 132 and 212ff.) etc. The **cerebellum** is an important control center for motor function (\rightarrow **p**, 326ff.).

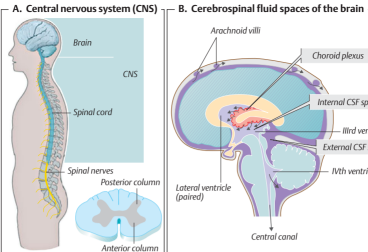
Diencephalon. The *thalamus* (\rightarrow **C6**) of the diencephalon functions as a relay station for most afferents, e.g., from the eyes, ears and skin as well as from other parts of the brain. The *hypothalamus* (\rightarrow **C9**) is a higher autonomic center (\rightarrow p. 330), but it also plays a dominant role in endocrine function (\rightarrow p. 266ff.) as it controls the release of hormones from the adjacent *hypophysis* (\rightarrow **D4**).

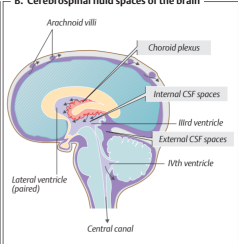
The **telencephalon** consists of the cortex and nuclei important for motor function, the

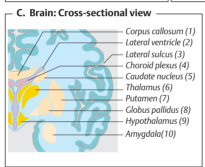
basal ganglia, i.e. *caudate nucleus* (\rightarrow **C5**), *putamen* (\rightarrow **C7**), *globus pallidus* (\rightarrow **C8**), and parts of the *amygdala* (\rightarrow **C10**). The amygdaloid nucleus and *cingulate gyrus* (\rightarrow **D2**) belong to the **limbic system** (\rightarrow **p.** 330). The **cerebral cortex** consists of four *lobes* divided by fissures (sulci), e.g., the *central sulcus* (\rightarrow **D1**, **E**) and *lateral sulcus* (\rightarrow **C3**, **E**). According to *Brodmann's map*, the cerebral cortex is divided into histologically distinct regions (\rightarrow **E**, italic letters) that generally have different functions (\rightarrow **E**). The *hemispheres* of the brain are closely connected by nerve fibers of the *corpus callosum* (\rightarrow **C1**, **D3**).

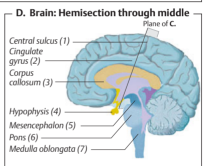
Cerebrospinal Fluid

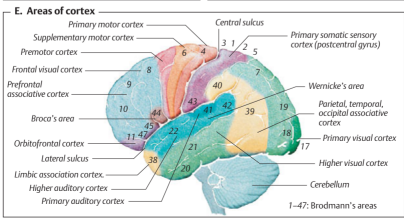
The brain is surrounded by external and internal cerebrospinal fluid (CSF) spaces (\rightarrow **B**). The internal CSF spaces are called ventricles. The two lateral ventricles, I and II, $(\rightarrow B, C2)$ are connected to the IIIrd and IVth ventricle and to the central canal of the spinal cord $(\rightarrow B)$. Approximately 650 mL of CSF forms in the choroid plexus $(\rightarrow B, C4)$ and drains through the arachnoid villi each day (\rightarrow **B**). Lesions that obstruct the drainage of CSF (e.g., brain tumors) result in cerebral compression; in children, they lead to fluid accumulation (hydrocephalus). The blood-brain barrier and the blood-CSF barrier prevents the passage of most substances except CO₂, O₂, water and lipophilic substances. (As an exception, the circumventricular organs of the brain such as the organum vasculosum laminae terminalis (OVLT; \rightarrow p. 280) and the area postrema (\rightarrow p. 238) have a less tight blood-brain barrier.) Certain substances like glucose and amino acids can cross the blood-brain barrier with the aid of carriers, whereas proteins cannot. The ability or inability of a drug to cross the blood-brain barrier is an important factor in pharmacotherapeutics.











Stimulus Reception and Processing

With our **senses**, we receive huge quantities of **information** from the surroundings (10^9 bits/s) . Only a small portion of it is consciously perceived $(10^1-10^2 \text{ bits/s})$; the rest is either subconsciously processed or not at all. Conversely, we transmit ca. 10^7 bits/s of information to the environment through speech and motor activity. especially facial expression $(\rightarrow A)$.

A **bit** (binary digit) is a single unit of information (1 byte = 8 bits). The average page of a book contains roughly 1000 bits, and TV images convey more than 10^6 bits/s.

Stimuli reach the body in different forms of energy, e.g., electromagnetic (visual stimuli) or mechanical energy (e.g., tactile stimuli). Various sensory receptors or sensors for these stimuli are located in the five "classic" sense organs (eye, ear, skin, tongue, nose) at the body surface as well as inside the body (e.g., propriosensors, vestibular organ). (In this book, sensory receptors are called sensors to distinguish them from binding sites for hormones and transmitters.) The sensory system extracts four stimulatory elements: modality, intensity, duration, and localization. Each type of sensor is specific for a unique or adequate stimulus that evokes specific sensory **modalities** such as sight, sound, touch, vibration, temperature, pain, taste, smell, as well as the body's position and movement, etc. Each modality has several submodalities, e.g., taste can be sweet or bitter. etc.

In **secondary sensors** (e.g., gustatory and auditory sensors), sensor and afferent fibers are separated by a synapse, whereas **primary sensors** (e.g., olfactory sensors and nocisensors) have their own afferent fibers.

A stimulus induces a change in *sensor potential* (**transduction**), which results in depolarization of the sensor cell (in most types; \rightarrow **B1**) or hyperpolarization as in retinal sensors. The stronger the stimulus, the greater the *amplitude* of the sensor potential (\rightarrow **C1**). Once the sensor potential exceeds a certain threshold, it is **transformed** into an **action potential**, **AP** (\rightarrow **B1**; p. 46ff.).

Coding of signals. The stimulus is encoded in *AP frequency* (impulses/s = Hz), i.e., the

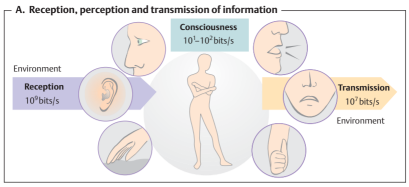
higher the sensor potential, the higher the AP frequency (\rightarrow C2). This information is decoded at the next synapse: The higher the frequency of arriving APs, the higher the *excitatory post-synaptic potential* (EPSP; \rightarrow 50ff.). New APs are fired by the postsynaptic neuron when the EPSP exceeds a certain threshold (\rightarrow B2).

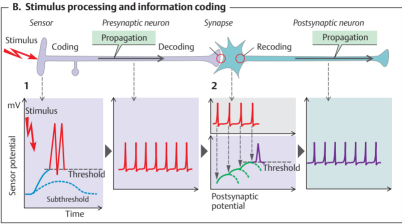
Frequency coding of APs is a more reliable way of transmitting information over long distances than amplitude coding because the latter is much more susceptible to change (and falsification of its information content). At the synapse, however, the signal must be amplified or attenuated (by other neurons), which is better achieved by amplitude coding.

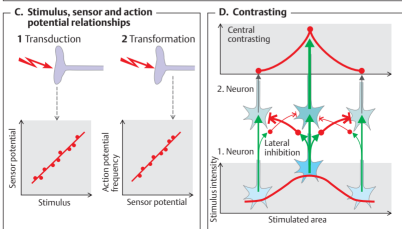
Adaptation. At constant stimulation, most sensors adapt, i.e., their potential decreases. The potential of slowly adapting sensors becomes **proportional** to stimulus intensity (*Psensors* or *tonic sensors*). Fast-adapting sensors respond only at the onset and end of a stimulus. They sense **differential** changes in the stimulus intensity (*Dsensors* or *phasic sensors*). *PD sensors* have both characteristics (\rightarrow p. 314).

Central processing. In a first phase, inhibitory and stimulatory impulses conducted to the CNS are integrated-e.g., to increase the *contrast* of stimuli (\rightarrow **D**; see also p. 354). In this case, stimulatory impulses originating from adjacent sensor are attenuated in the process (lateral inhibition). In a second step, a sensory impression of the stimuli (e.g. "green" or "sweet") takes form in low-level areas of the sensory cortex. This is the first step of subjective sensory physiology. Consciousness is a prerequisite for this process. Sensory impressions are followed by their interpretation. The result of it is called perception, which is based on experience and reason, and is subject to ininterpretation. The impression "green," for example, can evoke the perception "There is a tree" or "This is a meadow."

Absolute threshold (\rightarrow pp. 340ff., 352, 362), difference threshold (\rightarrow pp. 340ff., 352, 368), spatial and temporal summation (\rightarrow pp. 52, 352), receptive field (\rightarrow p. 354), habituation and sensitization are other important concepts of sensory physiology. The latter two mechanisms play an important role in learning processes (\rightarrow p. 336).







Sensory Functions of the Skin

Somatovisceral sensibility is the collective term for all sensory input from receptors or *sensors* of the body (as opposed to the sensory organs of the head). It includes the areas of proprioception $(\rightarrow p.316)$, nociception $(\rightarrow p.318)$, and skin or surface sensitivity.

The **sense of touch** (taction) is essential for perception of *form*, *shape*, and *spatial nature* of objects (*stereognosis*). Tactile sensors are located predominantly in the palm, especially in the fingertips, and in the tongue and oral cavity. Stereognostic perception of an object requires that the CNS integrate signals from adjacent receptors into a spatial pattern and coordinate them with **tactile motor function**.

Mechanosensors. *Hairless areas* of the skin contain the following mechanosensors (\rightarrow **A**), which are afferently innervated by myelinated nerve fibers of class II/A β (\rightarrow p. 49 C):

- The spindle-shaped Ruffini's corpuscle (→ A3) partly encapsulates the afferent axon branches. This unit is a slowly adapting (SA) pressosensor of the SA2 type. They are P sensors (→ p. 312). Thus, the greater the pressure on the skin (depth of indentation or weight of an object), the higher the AP frequency (→ B1).
 Merkel's cells (→ A2) are in synaptic contact to meniscus-shaped axon terminals. These complexes are pressure-sensitive SA1 sensors. They are PD sensors (combination of B1 and B2) since their AP frequency is not only dependent on the pressure intensity but also on
- ♦ Meissner's corpuscles (→ A1) are composed of lamellar cell layers between which clubshaped axons terminate. This unit represents a rapidly adapting pressure sensor (RA sensor) that responds only to pressure changes, dp/dt (pure *D sensor* or velocity sensor). The RA sensors are specific for touch (skin indentation of 10–100 μm) and low-frequency vibration (10–100 Hz). Hair follicle receptors (→ A5), which respond to bending of the hairs, assume these functions in hairy areas of the skin.

the rate of its change (dp/dt; \rightarrow p. 312).

◆ Pacinian corpuscles (→ **A4**) are innervated by a centrally situated axon. They *adapt very rapidly* and therefore respond to changes in pressure change velocity, i.e. to *acceleration* (d^2p/dt^2), and sense high-frequency vibration (100–400 Hz; indentation depths < 3 µm). The

AP frequency is proportional to the vibration frequency (\rightarrow **B3**).

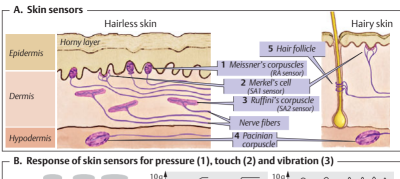
Resolution. RA and SA1 sensors are densely distributed in the mouth, lips and fingertips, especially in the index and middle finger (about 100/cm²). They can distinguish closely adjacent stimuli as separate, i.e., each afferent axon has a *narrow receptive field*. Since the signals do not converge as they travel to the CNS, the ability of these sensors in the mouth, lips and fingertips to distinguish between two closely adjacent tactile stimuli, i.e. their *resolution*, is very high.

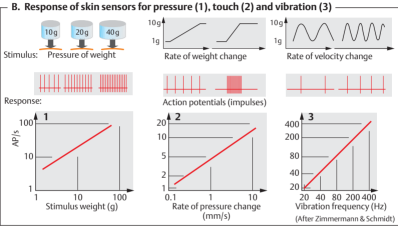
The spatial threshold for two-point discrimination, i.e., the distance at which two simultaneous stimuli can be perceived as separate, is used as a measure of tactile resolution. The spatial thresholds are roughly 1 mm on the fingers, lips and tip of the tongue, 4 mm on the palm of the hand, 15 mm on the arm, and over 60 mm on the back.

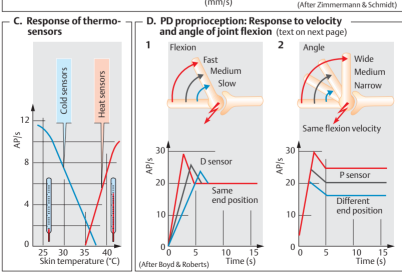
SA2 receptors and pacinian corpuscles have a broad receptive field (the exact function of SA2 receptors is not known). Pacinian corpuscles are therefore well adapted to detect vibrations, e.g., earth tremors.

Two types of **thermosensors** are located in the skin: cold sensors for temperatures < 36 °C and warm sensors for those > 36 °C. The lower the temperature (in the 20-36°C range), the higher the AP frequency of the cold receptors. The reverse applies to warm receptors in the 36–43 °C range (\rightarrow **C**). Temperatures ranging from 20° to 40°C are subject to rapid adaptation of thermosensation (PD characteristics). Water warmed, for example, to 25 °C initially feels cold. More extreme temperatures, on the other hand, are persistently perceived as cold or hot (this helps to maintain a constant core temperature and prevent skin damage). The density of these cold and warm sensors in most skin areas is low as compared to the much higher densities in the mouth and lips. (That is why the lips or cheeks are used for temperature testing.)

Different sensors are responsible for thermoception at temperatures exceeding 45 °C. These **heat sensors** are also used for the perception of pungent substances such as *capsaicin*, the active constituent of hot chili peppers. Stimulation of VR1 receptors (vanilloid receptor type 1) for capsaicin mediates the opening of cation channels in nociceptive nerve endings, which leads to their depolarization.







Proprioception, Stretch Reflex

Proprioception is the mechanism by which we sense the *strength* that our muscles develop as well as the *position* and *movement* of our body and limbs. The *vestibular organ* (\rightarrow p. 342) and *cutaneous mechanosensors* (\rightarrow p. 314) assist the *propriosensors* in muscle spindles, joints and tendons. Sensors of *Golgi tendon organs* are located near muscle—tendon iunctions.

Muscle spindles $(\rightarrow A1)$ contain intensity (P) and differential (D) sensors for monitoring of joint position and movement. The velocity of position change is reflected by a transient rise in impulse frequency (D sensor; \rightarrow p. 315 D1, spike), and the final joint position is expressed as a constant impulse frequency (P-sensor, → p. 315 D2, plateau). Muscle spindles function to regulate muscle length. They lie parallel to the skeletal muscle fibers (extrafusal muscle fibers) and contain their own muscle fibers (intrafusal muscle fibers). There are two types of intrafusal muscle fibers: (1) nuclear chain fibers (P sensors) and (2) nuclear bag fibers (D sensors). The endings of type Ia afferent neurons coil around both types, whereas type II neurons wind around the nuclear chain fibers only (neuron types described on p. 49 C). These annulospiral endings detect longitudinal stretching of intrafusal muscle fibers and report their length (type Ia and II afferents) and changes in length (Ia afferents) to the spinal cord. The efferent γ motoneurons (fusimotor fibers) innervate both intrafusal fiber types, allowing variation of their length and stretch sensitivity (\rightarrow A1, B1).

Golgi tendon organs (\rightarrow **A2**) are arranged in series with the muscle and respond to the contraction of only a few motor units or more. Their primary function is to regulate muscle tension. Impulses from Golgi tendon organs (conveyed by type Ib afferents), the skin and joints, and muscle spindles (some of which are type Ia and II afferent fibers), as well as descending impulses, are jointly integrated in type Ib interneurons of the spinal cord; this is referred to as **multimodal integration** $(\rightarrow D2)$. Type Ib interneurons inhibit α motoneurons of the muscle from which the Ib afferent input originated (autogenous inhibition) and activate antagonistic muscles via excitatory interneurons (\rightarrow **D5**).

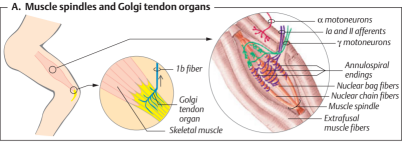
Monosynaptic stretch reflex (\rightarrow C). Muscles spindles are also affected by sudden stretching of a skeletal muscle, e.g. due to a tap on the tendon attaching it. Stretching of the muscle spindles triggers the activation of type Ia afferent impulses (\rightarrow **B2. C**), which enter the spinal cord via the dorsal root and terminate in the ventral horn at the α motoneurons of the same muscle. This type Ia afferent input therefore induces contraction of the same muscle by only one synaptic connection. The reflex time for this monosynaptic stretch reflex is therefore very short (ca. 30 ms). This is classified as a proprioceptive reflex, since the stimulation and response arise in the same organ. The monosynaptic stretch reflex functions to rapidly correct "involuntary" changes in muscle length and joint position.

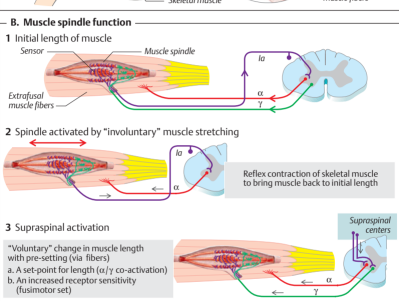
Supraspinal activation $(\rightarrow$ B3). Voluntary muscle contractions are characterized by *coactivation* of α and γ neurons. The latter adjust the muscle spindles (length sensors) to a certain set-point of length. Any deviations from this set-point due, for example, to unexpected shifting of weight, are compensated for by readjusting the α -innervation (load compensation reflex). Expected changes in muscle length, especially during complex movements, can also be more precisely controlled by (centrally regulated) γ fiber activity by increasing the preload and stretch sensitivity of the intrafusal muscle fibers (fusimotor set).

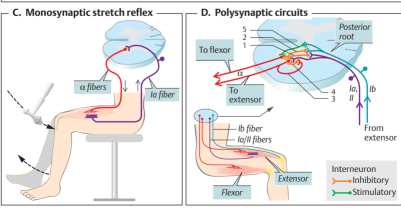
Hoffmann's reflex can be also used to test the stretch reflex pathway. This can be done by positioning electrodes on the skin over (mixed) muscle nerves and subsequently recording the muscle contraction induced by electrical stimuli of different intensity.

Polysynaptic circuits, also arising from type II afferents complement the stretch reflex. If a stretch reflex (e.g., knee-jerk reflex) occurs in an extensor muscle, the α motoneurons of the antagonistic flexor muscle must be inhibited via *inhibitory la interneurons* to achieve efficient extension (\rightarrow **D1**).

Deactivation of stretch reflex is achieved by inhibiting muscle contraction as follows: 1) The muscle spindles relax, thereby allowing the deactivation of type la fibers; 2) the Golgi tendon organs inhibit the α motoneurons via type lb interneurons (\rightarrow **D2**); 3) the α motoneurons are inhibited by the interneurons (*Renshaw cells*; \rightarrow **D4**) that they themselves stimulated via axon collaterals (*recurrent inhibition*; \rightarrow **D3**; p. 321 C1).







Pain is an unpleasant sensory experience associated with discomfort. It is protective insofar as it signals that the body is being threatened by an injury (noxa). Nociception is the perception of noxae via nocisensors, neural conduction and central processing. The pain that is ultimately felt is a subjective experience. Pain can also occur without stimulation of nocisensors, and excitation of nocisensors does not always evoke pain.

All body tissues except the brain and liver contain sensors for pain, i.e., **nocisensors** or **nociceptors** (\rightarrow **A**). Nocisensors are bead-like endings of peripheral axons, the somata of which are located in dorsal root ganglia and in nuclei of the trigeminal nerve. Most of these fibers are slowly conducting C fibers (< 1 m/s); the rest are myelinated A δ fibers (5–30 m/s; fiber types described on p. 49 C).

When an injury occurs, one first senses sharp "fast pain" (Aδ fibers) before feeling the dull "slow pain" (C fibers), which is felt longer and over a broader area. Since nocisensors do not adapt, the pain can last for days. Sensitization can even lower the stimulus threshold.

Most nocisensors are **polymodal sensors** (C fibers) activated by mechanical stimuli, chemical mediators of inflammation, and high-intensity heat or cold stimuli. **Unimodal nociceptors**, the less common type, consist of thermal nocisensors (A δ fibers), mechanical nocisensors (A δ fibers), and "dormant nocisensors." Thermal nocisensors are activated by extremely hot (> 45 °C) or cold (< 5 °C) stimuli (\rightarrow p. 314). Dormant nocisensors are chiefly located in internal organs and are "awakened" after prolonged exposure (sensitization) to a stimulus, e.g., inflammation.

Nocisensors can be inhibited by opioids (**desensitization**) and stimulated by prostaglandin E_2 or bradykinin, which is released in response to inflammation (**sensitization**; \rightarrow **A**). Endogenous opioids (e.g., dynorphin, enkephalin, endorphin) and exogenous opioids (e.g., morphium) as well as inhibitors of prostaglandin synthesis (\rightarrow p. 269) are therefore able to alleviate pain (*analgesic action*).

Inflammatory sensitization (e.g., sunburn) lowers the threshold for noxious stimuli, leading to excessive sensitivity (hyperalgesia) and additional pain resulting from non-noxious stimuli to the skin (allodynia), e.g., touch or warm water (37 °C). Once the

nocisensors are stimulated, they start to release *neu-ropeptides* such as substance P or CGRP (calcitonin gene-related peptide) that cause inflammation of the surrounding vessels (**neurogenic inflammation**).

Projected pain. Damage to nociceptive fibers causes pain (neurogenic or neuropathic) that is often projected to and perceived as arising from the periphery. A prolapsed disk compressing a spinal nerve can, for example, cause leg pain. Nociceptive fibers can be blocked by cold or local anesthesia.

Nociceptive tracts (→ C1). The central axons of nociceptive somatic neurons and nociceptive afferents of internal organs end on neurons of the dorsal horn of the spinal cord. In many cases, they terminate on the same neurons as the skin afferents.

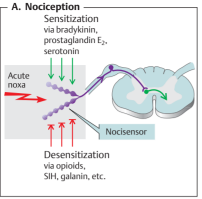
Referred pain (\rightarrow B). Convergence of somatic and visceral nociceptive afferents is probably the main cause of referred pain. In this type of pain, noxious visceral stimuli cause a perception of pain in certain skin areas called **Head's zones**. That for the heart, for example, is located mainly in the chest region. Myocardial ischemia is therefore perceived as pain on the surface of the chest wall (angina pectoris) and often also in the lower arm and upper abdominal region.

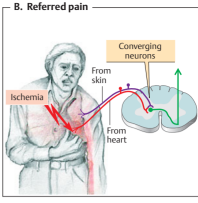
In the spinal cord, the neuroceptive afferents cross to the opposite side (decussation) and are conducted in the **tracts of the anterolateral funiculus**—mainly in the spinothalamic tract—and continue centrally via the brain stem where they join nociceptive afferents from the head (mainly trigeminal nerve) to the **thalamus** (\rightarrow **C1**). From the ventrolateral thalamus, sensory aspects of pain are projected to S1 and S2 areas of the **cortex**. Tracts from the medial thalamic nuclei project to the limbic system and other centers.

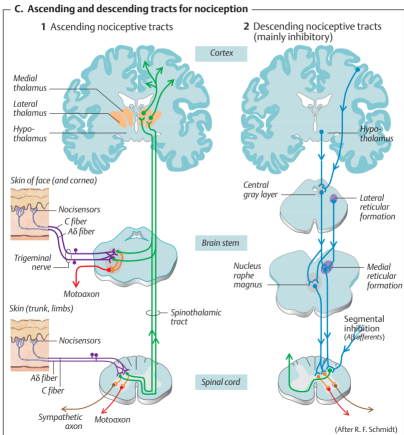
Components of pain. Pain has a sensory component including the conscious perception of site, duration and intensity of pain; a motor component (e.g., defensive posture and withdrawal reflex; \rightarrow p. 320), an autonomic component (e.g., tachycardia), and an affective component (e.g., aversion). In addition, pain assessments based on the memory of a previous pain experience can lead to pain-related behavior (e.g., moaning).

In the thalamus and spinal cord, nociception can be **inhibited via descending tracts** with the aid of various transmitters (mainly opioids). The nuclei of these tracts (\rightarrow **C2**, blue) are located in the *brain stem* and are mainly activated via the nociceptive spinoreticular tract (negative feedback loop).

318







Polysynaptic Reflexes

Unlike proprioceptive reflexes (\rightarrow p. 316), polysynaptic reflexes are activated by sensors that are spatially separate from the effector organ. This type of reflex is called polysynaptic. since the reflex arc involves many synapses in series. This results in a relatively long reflex time. The intensity of the response is dependent on the duration and intensity of stimulus, which is temporally and spatially summated in the CNS (\rightarrow p. 52). Example: itching sensation in nose _⇒ sneezing. The response spreads when the stimulus intensity increases (e.g., coughing ⇒ choking cough). Protective reflexes (e.g., withdrawal reflex, corneal and lacrimal reflexes, coughing and sneezing), nutrition reflexes (e.g., swallowing, sucking reflexes), locomotor reflexes, and the various autonomic reflexes are polysynaptic reflexes. Certain reflexes, e.g., plantar reflex, cremasteric reflex and abdominal reflex, are used as diagnostic tests.

Withdrawal reflex $(\rightarrow A)$. Example: A painful stimulus in the sole of the right foot (e.g., stepping on a tack) leads to flexion of all joints of that leg (flexion reflex). Nociceptive afferents $(\rightarrow p.318)$ are conducted via stimulatory interneurons $(\rightarrow A1)$ in the spinal cord to motoneurons of ipsilateral flexors and via inhibitory interneurons $(\rightarrow A2)$ to motoneurons of ipsilateral extensors (→ A3), leading to their relaxation; this is called antagonistic inhibition. One part of the response is the crossed extensor reflex, which promotes the withdrawal from the injurious stimulus by increasing the distance between the nociceptive stimulus (e.g. the tack) and the nocisensor and helps to support the body. It consists of contraction of extensor muscles (\rightarrow A5) and relaxation of the flexor muscles in the contralateral leg ($\rightarrow A4$, **A6**). Nociceptive afferents are also conducted to other segments of the spinal cord (ascending and descending; \rightarrow A7, A8) because different extensors and flexors are innervated by different segments. A noxious stimulus can also trigger flexion of the ipsilateral arm and extension of the contralateral arm (double crossed extensor reflex). The noxious stimulus produces the perception of pain in the brain $(\to p.316)$.

Unlike monosynaptic stretch reflexes, polysynaptic reflexes occur through the *co-activation* of α and γ motoneurons (\rightarrow p. 316). The reflex excitability of α motoneurons is largely controlled by supraspinal centers via multiple interneurons (\rightarrow p. 324). The brain can therefore shorten the reflex time of spinal cord reflexes when a noxious stimulus is anticipated.

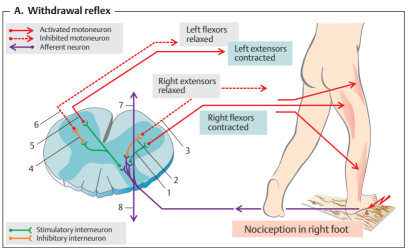
Supraspinal lesions or interruption of descending tracts (e.g., in paraplegics) can lead to exaggeration of reflexes (**hyperreflexia**) and stereotypic reflexes. The absence of reflexes (**areflexia**) corresponds to specific disorders of the spinal cord or peripheral nerve.

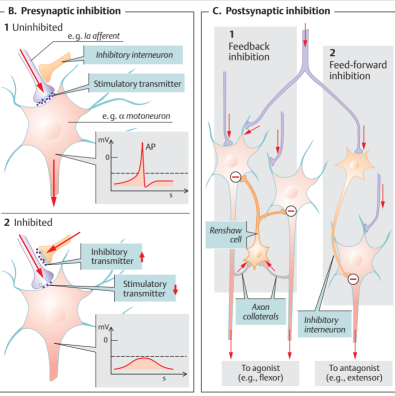
Synaptic Inhibition

GABA (γ-aminobutyric acid) and glycine $(\rightarrow p.55f.)$ function as inhibitory transmitters in the spinal cord. **Presynaptic inhibition** $(\rightarrow B)$ occurs frequently in the CNS, for example, at synapses between type Ia afferents and α motoneurons, and involves axoaxonic synapses of GABAergic interneurons at presynaptic nerve endings, GABA exerts inhibitory effects at the nerve endings by increasing the membrane conductance to Cl- (GABAA receptors) and K+ (GABA_R receptors) and by decreasing the conductance to Ca2+ (GABAB receptors). This decreases the release of transmitters from the nerve ending of the target neuron $(\rightarrow B2)$, thereby lowering the amplitude of its postsynaptic EPSP (\rightarrow p. 50). The purpose of presynaptic inhibition is to reduce certain influences on the motoneuron without reducing the overall excitability of the cell.

In **postsynaptic inhibition** (\rightarrow C), an *inhibitory interneuron* increases the membrane conductance of the postsynaptic neuron to Cl $^-$ or K $^+$, especially near the axon hillock, thereby short-circuiting the depolarizing electrical currents from excitatory EPSPs (\rightarrow p. 54 D).

The interneuron responsible for postsynaptic inhibition is either activated by feedback from axonal collaterals of the target neurons (**recurrent inhibition** of motoneurons via glycinergic Renshaw cells; \rightarrow C1) or is directly activated by another neuron via feed-forward control (\rightarrow C2). Inhibition of the ipsilateral extensor (\rightarrow A2, A3) in the flexor reflex is an example of feed-forward inhibition.





SD12.6
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Central Conduction of Sensory Input

The posterior funiculus-lemniscus system $(\rightarrow \mathbf{C}, \text{ green})$ is the principal route by which the somatosensory cortex S1 (postcentral gyrus) receives sensory input from skin sensors and propriosensors. Messages from the skin (superficial sensibility) and locomotor system (proprioceptive sensibility) reach the spinal cord via the dorsal roots. Part of these primarily afferent fibers project in tracts of the posterior funiculus without synapses to the posterior funicular nuclei of the caudal medulla oblongata (nuclei cuneatus and gracilis). The tracts of the posterior funiculi exhibit a somatotopic arrangement, i.e., the further cranial the origin of the fibers the more lateral their location. At the medial lemniscus, the secondary afferent somatosensory fibers cross to the contralateral side (decussate) and continue to the posterolateral ventral nucleus (PLVN) of the thalamus, where they are also somatotopically arranged. The secondary afferent trigeminal fibers (lemniscus trigeminalis) end in the posteromedial ventral nucleus (PMVN) of the thalamus. The tertiary afferent somatosensory fibers end at the quaternary somatosensory neurons in the somatosensory cortex \$1. The main function of the posterior funiculus-lemniscus pathway is to relay information about tactile stimuli (pressure, touch, vibration) and joint position and movement (proprioception) to the brain cortex via its predominantly rapidly conducting fibers with a high degree of spatial and temporal resolution.

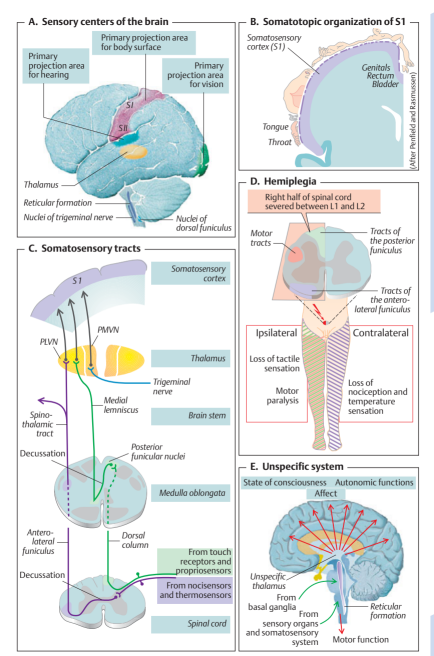
As in the motor cortex (\rightarrow p. 325 B), each body part is assigned to a corresponding projection area in the **somatosensory cortex S1** (\rightarrow A) following a *somatotopic arrangement* (\rightarrow B). Three features of the organization of S1 are (1) that one hemisphere of the brain receives the information from the contralateral side of the body (tracts decussate in the medial lemniscus; \rightarrow C); (2) that most neurons in S1 receive afferent signals from tactile sensors in the fingers and mouth (\rightarrow p. 314); and (3) that the afferent signals are processed in columns of the cortex (\rightarrow p. 333 A) that are activated by specific types of stimuli (e.g., touch).

Anterolateral spinothalamic pathway (\rightarrow **C**; violet). Afferent signals from nocisensors, thermosensors and the second part of pressure and touch afferent neurons are already relayed (partly via interneurons) at various levels of the *spinal cord*. The secondary neurons cross to the opposite side at the corresponding segment of the spinal cord, form the lateral and ventral *spinothalamic tract* in the anterolateral funiculus, and project to the thalamus.

Descending tracts (from the cortex) can inhibit the flow of sensory input to the cortex at all relay stations (spinal cord, medulla oblongata, thalamus). The main function of these tracts is to modify the receptive field and adjust stimulus thresholds. When impulses from different sources are conducted in a common afferent, they also help to suppress unimportant sensory input and selectively process more important and interesting sensory modalities and stimuli (e.g., eavesdropping).

Hemiplegia. (\rightarrow **D**) Brown–Séquard syndrome occurs due to hemisection of the spinal cord, resulting in ipsilateral paralysis and loss of various functions below the lesion. The injured side exhibits motor paralysis (initially flaccid, later spastic) and loss of tactile sensation (e.g., impaired two-point discrimination, \rightarrow p. 314). An additional loss of pain and temperature sensation occurs on the contralateral side (*dissociated paralysis*).

Reticular activating system. (\rightarrow E) The sensory input described above as well as the input from the sensory organs are specific, whereas the reticular activating system (RAS) is an unspecific system. The RAS is a complex processing and integrating system of cells of the reticular formation of the brainstem. These cells receive sensory input from all sensory organs and ascending spinal cord pathways (e.g., eyes, ears, surface sensitivity, nociception), basal ganglia, etc. Cholinergic and adrenergic output from the RAS is conducted along descending pathways to the spinal cord and along ascending "unspecific" thalamic nuclei and "unspecific" thalamocortical tracts to almost all cortical regions (\rightarrow p. 333 A), the limbic system and the hypothalamus. The ascending RAS or ARAS controls the state of consciousness and the degree of wakefulness (arousal activity).



Motor System

Coordinated muscular movements (walking, grasping, throwing, etc.) are functionally dependent on the *postural motor system*, which is responsible for maintaining upright posture, balance, and spatial integration of body movement. Since control of postural motor function and muscle coordination requires the simultaneous and uninterrupted flow of sensory impulses from the periphery, this is also referred to as sensorimator function.

 α **motoneurons** in the anterior horn of the spinal cord and in cranial nerve nuclei are the terminal tracts for skeletal muscle activation. Only certain parts of the corticospinal tract and type la afferents connect to α motoneurons monosynaptically. Other afferents from the periphery (propriosensors, nocisensors, mechanosensors), other spinal cord segments, the motor cortex, cerebellum, and motor centers of the brain stem connect to α motoneurons via hundreds of inhibitory and stimulatory interneurons per motoneuron.

Voluntary motor function. Voluntary movement requires a series of actions: decision to move ⇒ programming (recall of stored subprograms) ⇒ command to move ⇒ execution of movement (→ **A1-4**). Feedback from afferents (re-afferents) from motor subsystems and information from the periphery is constantly integrated in the process. This allows for adjustments before and while executing voluntary movement.

The neuronal activity associated with the first two phases of voluntary movement activates numerous motor areas of the cortex. This electrical brain activity is reflected as a negative **cortical expectancy potential**, which can best be measured in association areas and the vertex. The more complex the movement, the higher the expectancy potential and the earlier its onset (roughly 0.3–3 s).

The **motor cortex** consists of three main areas (\rightarrow C, top; \rightarrow see p. 311 E for area numbers): (a) *primary motor area*, M1 (area 4), (b) *premotor area*, PMA (lateral area 6); and (c) *supplementary motor area*, SMA (medial area 6). The motor areas of the cortex exhibit somatopic organization with respect to the target muscles of their fibers (shown for M1 in B) and their mutual connections.

Cortical afferents. The cortex receives motor input from (a) the *body periphery* (via thalamus \Rightarrow S1 [\rightarrow p. 323 A] \Rightarrow sensory association cortex \Rightarrow PMA); (b) the *basal ganglia* (via thalamus \Rightarrow M1, PMA, SMA [\rightarrow A2] \Rightarrow prefrontal association cortex); (c) the *cerebellum* (via thalamus \Rightarrow M1, PMA; \rightarrow A2); and (d) sensory and posterior parietal areas of the *cortex* (areas 1–3 and 5–7, respectively).

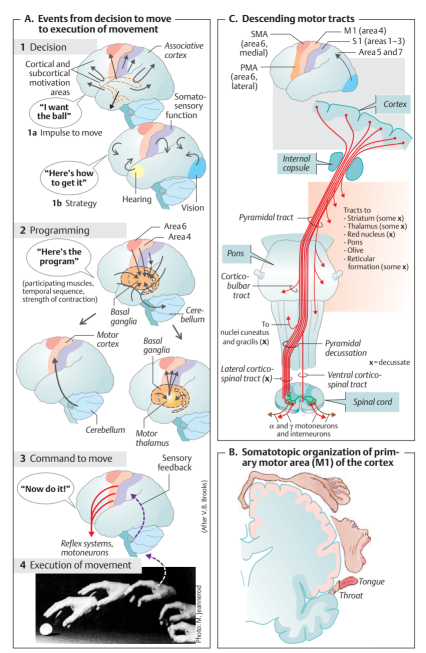
Cortical efferents. (\rightarrow **C**, **D**, **E**, **F**) Motor output from the cortex is mainly projected to (a) the spinal cord, (b) subcortical motor centers (see below and p.328), and (c) the contralateral cortex via commissural pathways.

The **pyramidal tract** includes the *corticospinal tract* and part of the *corticobulbar tract*. Over 90% of the pyramidal tract consists of thin fibers, but little is known about their function. The thick, rapidly conducting corticospinal tract (\rightarrow **C**) project to the spinal cord from areas 4 and 6 and from areas 1–3 of the sensory cortex. Some of the fibers connect monosynaptically to α and γ motoneurons responsible for finger movement (precision grasping). The majority synapse with interneurons of the spinal cord, where they influence input from peripheral afferents as well as motor output (via Renshaw's cells) and thereby spinal reflexes.

Function of the Basal Ganglia

Circuitry. The basal ganglia are part of multiple parallel corticocortical signal loops. Associative loops arising in the frontal and limbic cortex play a role in mental activities such as assessment of sensory information, adaptation of behavior to emotional context, motivation, and long-term action planning. The function of the skeletomotor and oculomotor loops (see below) is to coordinate and control the velocity of movement sequences. Efferent projections of the basal ganglia control thalamocortical signal conduction by (a) attenuating the inhibition (disinhibiting effect. direct mode) of the thalamic motor nuclei and the superior colliculus, respectively, or (b) by intensifying their inhibition (indirect mode).

The principal **input** to the basal ganglia comes from the putamen and caudate nucleus, which are collectively referred to as the *striatum*. Neurons of the striatum are activated by



tracts from the entire cortex and use glutamate as their transmitter $(\rightarrow \mathbf{D})$. Once activated, neurons of the striatum release an inhibitory transmitter (GABA) and a co-transmittereither substance P (SP) or enkephalin (Enk., \rightarrow **D**: \rightarrow p. 55). The principal **output** of the basal ganglia runs through the pars reticularis of the substantia nigra (SNr) and the pars interna of the globus pallidus (GPi), both of which are inhibited by SP/GABAergic neurons of the striatum $(\rightarrow \mathbf{D})$.

Both SNr and GPi inhibit (by GABA) the ventrolateral thalamus with a high level of spontaneous activity. Activation of the striatum therefore leads to disinhibition of the thalamus by this direct pathway. If, however, enkephalin/GABA-releasing neurons of the striatum are activated, then they inhibit the pars externa of the alobus pallidus (GPe) which, in turn, inhibits (by GABA) the subthalamic nucleus. The subthalamic nucleus induces glutamatergic activation of SNr and GPi. The ultimate effect of this indirect pathway is increased thalamic inhibition. Since the thalamus projects to motor and prefrontal cortex, a corticothalamocortical loop that influences skeletal muscle movement (skeletomotor loop) via the putamen runs through the basal ganglia. An oculomotor loop projects through the caudate nucleus, pars reticularis and superior colliculus and is involved in the control of eve movement ($\rightarrow pp.342$. 360). Descending tracts from the SNr project to the tectum and nucleus pedunculus pontinus.

The fact that the pars compacta of the substantia nigra (SNc) showers the entire striatum with dopamine (dopaminergic neurons) is of pathophysiological importance $(\rightarrow \mathbf{D})$. On the one hand, dopamine binds to D1 receptors (rising cAMP levels), thereby activating SP/GABAergic neurons of the striatum: this is the direct route (see above). On the other hand, dopamine also reacts with D2 receptors (decreasing cAMP levels), thereby inhibiting enkephalin/GABAergic neurons; this is the indirect route. These effects of dopamine are essential for normal striatum function. Degeneration of more than 70% of the dopaminergic neurons of the pars compacta results in excessive inhibition of the motor areas of the thalamus, thereby impairing voluntary motor function. This occurs in Parkinson's disease and can be due genetic predisposition, trauma (e.g., boxing), cerebral infection and other causes. The characteristic symptoms of disease include poverty of movement (akinesia), slowness of movement (bradykinesia), a festinating gait, small handwriting (micrography), masklike facial expression, muscular hypertonia (rigor), bent posture, and a tremor of resting muscles ("money-counting" movement of thumb and fingers).

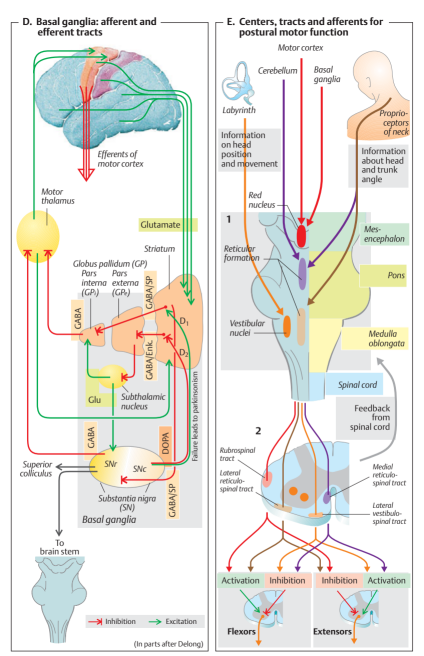
Function of the Cerebellum

The cerebellum contains as many neurons as the rest of the brain combined. It is an important control center for motor function that has afferent and efferent connections to the cortex and periphery (\rightarrow **F**, **top** panel). The cerebellum is involved in the planning, execution and control of movement and is responsible for adaptation to new movement motor sequences (motor learning). It is also cooperates with higher centers to control attention. etc.

Anatomy (\rightarrow **F, top**). The archeocerebellum (flocculonodular lobe) and paleocerebellum (pyramids, uvula, paraflocculus and parts of the anterior lobe) are the phylogenetically older parts of the cerebellum. These structures and the pars intermedia form the median cerebellum. The neocerebellum (posterior lobe of the body of the cerebellum) is the phylogenetically younger part of the cerebellum and forms the lateral cerebellum. Based on the origin of their principal efferents, the archicerebellum and vermis are sometimes referred to as the vestibulocerebellum, the paleocerebellum as the spinocerebellum, and the neocerebellum as the pontocerebellum. The cerebellar cortex is the folded (fissured) superficial gray matter of the cerebellum consisting of an outer molecular layer of Purkinje cell dendrites and their afferents, a middle layer of Purkinje cells (Purkinje somata), and an inner layer of granular cells. The outer surface of the cerebellum exhibits small, parallel convolutions called folia.

The median cerebellum and pars intermedia of the cerebellum mainly control postural and supportive motor function (\rightarrow **F1.2**) and oculomotor function (\rightarrow pp. 342 and 360). **Input:** The median cerebellum receives afference copies of spinal, vestibular and ocular origin and efference copies of descending motor signals to the skeletal muscles. Output from the median cerebellum flows through the intracerebellar fastigial, globose, and emboliform nuclei to motor centers of the spinal cord and brain stem and to extracerebellar vestibular nuclei (mainly Deiter's nucleus). These centers control oculomotor function and influence locomotor and postural/supportive motor function via the vestibulospinal tract.

The lateral cerebellum (hemispheres) mainly takes part in programmed movement $(\rightarrow F3)$, but its plasticity also permits motor adaptation and the learning of motor sequences. The hemispheres have two-way



connections to the cortex. **Input**: **a.** Via the pontine nuclei and mossy fibers, the lateral cerebellum receives input from cortical centers for movement planning (e.g., parietal, prefrontal and premotor *association cortex*; sensorimotor and visual areas). **b.** It also receives input from cortical and subcortical motor centers via the inferior olive and climbing fibers (see below). **Output** from the lateral cerebellum projects across motor areas of the thalamus from the dentate nucleus to motor areas of the cortex.

Lesions of the median cerebellum lead to disturbances of balance and oculomotor control (vertigo, nausea, pendular nystagmus) and cause trunk and gait ataxia. Lesions of the lateral cerebellum lead to disturbances of initiation, coordination and termination of goal-directed movement and impair the rapid reprogramming of diametrically opposing movement (diadochokinesia). The typical patient exhibits tremor when attempting voluntary coordinated movement (intention tremor), difficulty in measuring the distances during muscular movement (dysmetria), pendular rebound motion after stopping a movement (rebound phenomenon), and inability to perform rapid alternating movements (adiadochokinesia).

The cerebellar cortex exhibits a uniform neural **ultrastructure** and **circuitry**. All **output** from the cerebellar cortex is conducted via neurites of approximately 15×10^6 Purkinje cells. These GABAergic cells project to and *inhibit* neurons of the fastigial, emboliform, dentate, and lateral vestibular nuclei (Deiter's nucleus; \rightarrow **F**, **right** panel).

Input and circuitry: Input from the spinal cord (spinocerebellar tracts) is relayed by the inferior olive and projected via stimulatory (1:15 diverging) climbing fibers that terminate on a band of Purkinje cells extending across the folia of the cerebellum, forming the sagittal excitatory foci. The climbing fibers use aspartate as their transmitter. Serotoninergic fibers from the raphe nuclei and noradrenergic fibers from the locus caeruleus terminate also on the excitatory foci. Mossy fibers (pontine, reticular and spinal afferents) excite the granular cells. Their axons form T-shaped branches (parallel fibers). In the molecular layer, they densely converge (ca. 10⁵:1) on strips of Purkinje cells that run alongside the folium; these are called longitudinal excitatory foci. It is assumed that the climbing fiber system (at the "crossing points" of the perpendicular excitatory foci) amplify the relatively weak signals of mossy fiber afferents to Purkinje cells. Numerous interneurons (Golgi, stellate and basket cells) heighten the contrast of the excitatory pattern on the cerebellar cortex by lateral and recurrent inhibition.

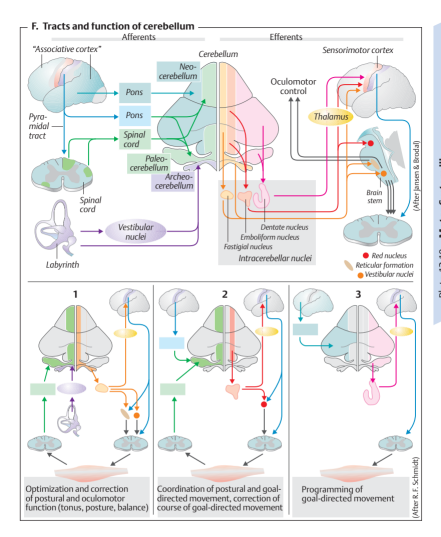
Postural Motor Control

Simple stretch reflexes (\rightarrow p.316)) as well as the more complicated flexor reflexes and crossed extensor reflexes (\rightarrow p.320) are controlled at the level of the **spinal cord**.

Spinal cord transection (paraplegia) leads to an initial loss of peripheral reflexes below the lesion (areflexia, spinal shock), but the reflexes can later be provoked in spite of continued transection.

The spinal reflexes are mainly subordinate to supraspinal centers $(\rightarrow E)$. Postural motor function is chiefly controlled by motor centers of the brain stem $(\rightarrow E1)$, i.e., the red nucleus, vestibular nuclei (mainly lateral vestibular nucleus), and parts of the reticular formation. These centers function as relay stations that pass along information pertaining to postural and labyrinthine postural reflexes required to maintain posture and balance (involuntary). Postural reflexes function to regulate muscle tone and eve adaptation movements $(\rightarrow p)$. 343 C). **Input** is received from the equilibrium organ (tonic labyrinthine reflexes) and from propriosensors in the neck (tonic neck reflexes). The same afferents are involved in postural reflexes (labyrinthine and neck reflexes) that help to maintain the body in its normal position. The trunk is first brought to its normal position in response to inflow from neck proprioceptors. Afferents projecting from the cerebellum, cerebral motor cortex $(\rightarrow C)$, eyes, ears, and olfactory organ as well as skin receptors also influence postural reflexes. Statokinetic reflexes also play an important role in the control of body posture and position. They play a role e.g. in startle reflexes and nystagmus (\rightarrow p. 360).

Descending tracts to the spinal cord arising from the red nucleus and medullary reticular formation (rubrospinal and lateral reticulospinal tracts) have a generally inhibitory effect on α and γ motoneurons (\rightarrow p. 316) of extensor muscles and an excitatory effect on flexor muscles (\rightarrow **E2**). Conversely, the tracts from Deiter's nucleus and the pontine areas of the reticular formation (vestibulospinal and medial reticulospinal tracts) inhibit the flexors and excite the α and γ fibers of the extensors. Transection of the brain stem below the red nucleus leads to decerebrate rigid-



ity because the extensor effect of Deiter's nucleus predominates.

The integrating and coordinating function of the sensorimotor system can be illustrated in two tennis players. When one player serves, the body of the other player moves to meet the ball (goal-directed movement) while using the right leg for support and the left arm for balance (postural motor control). The player keeps his eye on the ball (oculomotor control) and the visual area of the cortex assesses the trajectory and velocity of the ball. The associative cere-

bral cortex initiates the movement of returning the ball while taking the ball, net, other side of the court, and position of the opponent into consideration. Positional adjustments may be necessary when returning the ball. Using the movement concept programmed in the cerebellum and basal ganglia, the motor cortex subsequently executes the directed movement of returning the ball. In doing so, the player may "slice" the ball to give it an additional spinning motion (acquired rapid directed movement).

Hypothalamus, Limbic System

The **hypothalamus** coordinates all autonomic and most endocrine processes (\rightarrow p. 266ff.) and integrates signals for control of internal milieu, sleep–wake cycle, growth, mental/physical development, reproduction and other functions. The hypothalamus receives numerous sensory and humoral signals (\rightarrow A). Peptide hormones can circumvent the bloodbrain barrier by way of the *circumventricular organs* (\rightarrow p. 280).

Afferents. Thermosensors for control of body temperature (→ p. 224), osmosensors for regulation of osmolality and water balance (→ p. 168), and alucose sensors for maintenance of a minimum glucose concentration are located within the hypothalamus. Information about the current status of the internal milieu is neuronally projected to the hypothalamus from distant sensors, e.g., thermosensors in the skin, osmosensors in the liver (→ p. 170), and stretch sensors in the cardiac atria (\rightarrow p. 214ff.). The hypothalamus/circumventricular organs also contain receptors for various hormones (e.g., cortisol and angiotensin II), some of which form part of control loops for energy metabolism and metabolic homeostasis (e.g., receptors for cortisol, ACTH, CRH, leptin, and CCK). For functions related to growth and reproduction, the hypothalamus receives hormonal signals from the gonads and input from neuronal afferents that report cervical widening at the beginning of the birth process and breast stimulation (suckling reflexes), among other things.

The **limbic system** (→ **A**) and other areas of the brain influence hypothalamic function. The limbic system controls inborn and acquired behavior ("program selection") and is the seat of instinctive behavior, emotions and motivation ("inner world"). It controls the expression of emotions conveying important signals to the environment (e.g., fear, anger, wrath, discomfort, joy, happiness). Inversely, signals from the environment (e.g., odors) are closely associated to behavior.

The limbic system has cortical components (hippocampus, parahippocampal gyrus, cingulate gyrus, parts of olfactory brain) and subcortical components (amygdaloid body, septal nuclei, anterior thalamic nucleus). It has reciprocal connections to the lateral hypothalamus (chiefly used for recall of "programs", see below) and to the temporal and frontal cortex. Its connections to the cortex are primarily

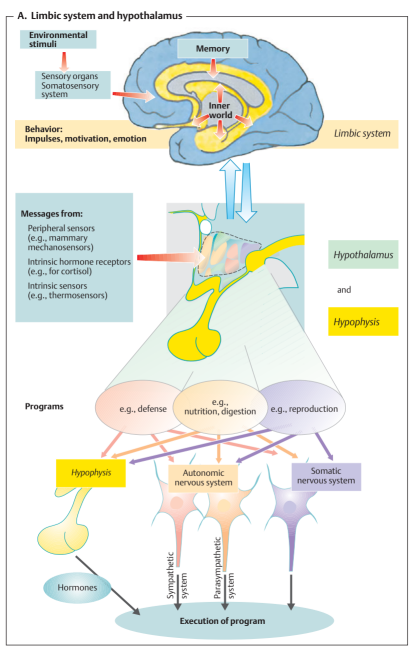
used to perceive and assess signals from the "outer world" and from memories. Processing of both types of input is important for behavior.

Programmed behavior (→**A**). The lateral hypothalamus has various programs to control lower hormonal, autonomic and motor processes. This is reflected internally by numerous autonomic and hormonal activities, and is reflected outwardly by different types of behavior.

Different programs exist for different behavioral reactions, for example:

- ◆ **Defensive behavior** ("fight or flight"). This program has somatic (repulsive facial expression and posture, flight or fight behavior), hormonal (epinephrine, cortisol) and autonomic (sympathetic nervous system) components. Its activation results in the release of energy-rich free fatty acids, the inhibition of insulin release, and a decrease in blood flow to the gastrointestinal tract as well as to rises in cardiac output, respiratory rate, and blood flow to the skeletal muscles.
- Physical exercise. The components of this program are similar to those of defensive behavior.
- Nutritive behavior, the purpose of which is to ensure an adequate supply, digestion and intake of foods and liquids. This includes searching for food, e.g. in the refrigerator, activation of the parasympathetic system with increased gastrointestinal secretion and motility in response to food intake, post-prandial reduction of skeletal muscle activity and similar activities.
- Reproductive behavior, e.g., courting a partner, neuronal mechanisms of sexual response, hormonal regulation of pregnancy (→ p. 304), etc.
- ◆ Thermoregulatory behavior, which enables us to maintain a relatively constant core temperature (→ p. 224), even in extreme ambient temperatures or at the high level of heat production during strenuous physical work.

Monoaminergic neuron systems contain neurons that release the monoamine neurotransmitters norepinephrine, epinephrine, dopamine, and serotonin. These neuron tracts extend from the brain stem to almost all parts of the brain and play an important role in the overall regulation of behavior. Experimental activation of noradrenergic neurons, for example, led to positive reinforcement (liking, rewards), whereas the serotoninergic neurons are thought to be associated with dislike. A number of psychotropic drugs target monoaminergic neuron systems.



Cerebral Cortex, Electroencephalogram (EEG)

Proper function of the cerebral cortex is essential for *conscious perception, planning, action*, and *voluntary movement* (\rightarrow p. 322ff.).

Cortical ultrastructure and **neuronal circuitry** (→ **A**). The cerebral cortex consists of six layers, I–VI, lying parallel to the brain surface. Vertically, it is divided into columns and modules (diameter 0.05–0.3 mm, depth 1.3–4.5 mm) that extend through all six layers.

Input from specific and unspecific areas of the thalamus terminate mainly on layers IV and on layers I and II, respectively (\rightarrow A3); those from other areas of the cortex terminate mainly on layer II (\rightarrow **A2**). The large and small pyramidal cells $(\rightarrow A1)$ comprise 80% of all cells in the cortex and are located in lavers V and III, respectively (glutamate generally serves as the transmitter, e.g., in the striatum; \rightarrow p. 325 D). The pyramidal cell axons leave the layer VI of their respective columns and are the sole source of output from the cortex. Most of the axons project to other areas of the ipsilateral cortex (association fibers) or to areas of the contralateral cortex (commissural fibers) $(\rightarrow A2)$; only a few extend to the periphery $(\rightarrow A4)$ and p. 325 C). Locally, the pyramidal cells are connected to each other by axon collaterals. The principal dendrite of a pyramidal cell projects to the upper layers of its column and has many thorn-like processes (spines) where many thalamocortical, commissural and association fibers terminate. The afferent fibers utilize various transmitters, e.g., norepinephrine, dopamine, serotonin, acetylcholine and histamine. Inside the cerebral cortex, information is processed by many morphologically variable **stellate cells** (\rightarrow A1), some of which have stimulatory effects (VIP, CCK and other peptide transmitters), while others have inhibitory effects (GABA). Dendrites of pyramidal and stellate cells project to neighboring columns, so the columns are connected by thousands of threads. Plasticity of pyramidal cell synapses — i.e., the fact that they can be modified in conformity with their activity pattern — is important for the learning process (\rightarrow p. 336).

Cortical potentials. Similar to electrocardiography, collective fluctuations of electrical potentials (brain waves) in the cerebral cortex can be recorded by **electroencephalography** using electrodes applied to the skin over the cranium (\rightarrow **B**). The EPSPs contribute the most to the electroencephalogram (**EEG**) whereas the share of the relatively low IPSPs (\rightarrow p. 50ff.) generated at the synapses of pyramidal cell

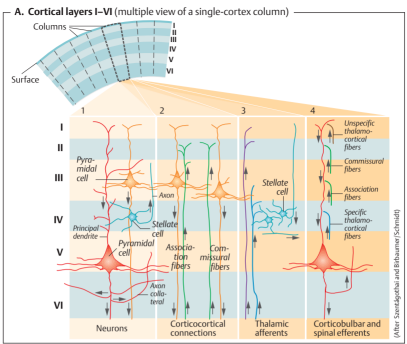
dendrites is small. Only a portion of the rhythms recorded in the EEG are produced directly in the cortex (α and γ waves in conscious perception; see below). Lower frequency waves from other parts of the brain, e.g. α waves from the thalamus and θ waves from the hippocampus, are "forced on" the cortex (*brain wave entrainment*).

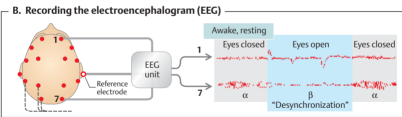
By convention, downward **deflections of the EEG** are positive. Generally speaking, depolarization (excitation) of deeper layers of the cortex and hyperpolarization of superficial layers cause downward deflection (+) and vice versa.

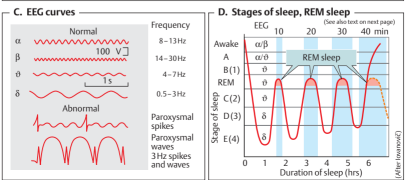
Brain wave types. The electrical activity level of the cortex is mainly determined by the degree of wakefulness and can be distinguished based on the amplitude (a) and frequency (f) of the waves (\rightarrow **B, C**). α **Waves** ($f \approx 10 \, \text{Hz}$; $a \approx 50 \, \mu \text{V}$), which predominate when an adult subject is awake and relaxed (with eyes closed), are generally detected in multiple electrodes (synchronized activity). When the eyes are opened, other sensory organs are stimulated, or the subject solves a math problem, the α waves subside (α blockade) and β waves appear ($f \approx 20 \,\text{Hz}$). The amplitude of β waves is lower than that of α waves, and they are chiefly found in occipital $(\rightarrow B)$ and parietal regions when they eyes are opened. The frequency and amplitude of β waves varies greatly in the different leads (desynchronization). B Waves reflect the increased attention and activity (arousal activity) of the ascending reticular activating system (ARAS; \rightarrow p. 322). γ Waves (> 30 Hz) appear during learning activity. Low-frequency θ waves appear when drowsiness descends to sleep (sleep stages A/B/C; \rightarrow **D**); they transform into even slower δ waves during deep sleep (\rightarrow **C. D**).

The EEG is used to diagnose epilepsy (localized or generalized paroxysmal waves and spikes; \rightarrow **C**), to assess the degree of brain maturation, monitor anesthesia, and to determine brain death (*isoelectric EEG*).

Magnetoencephalography (MEG), i.e. recording magnetic signals induced by cortical ion currents, can be combined with the EEG to precisely locate the site of cortical activity (resolution a few mm).







Sleep-Wake Cycle, Circadian Rhythms

Various stages of sleep can be identified in the EEG (\rightarrow p. 333 D). When a normal person who is awake, relaxed and has the eyes closed (α waves) starts to fall asleep, the level of consciousness first descends to sleep phase A (dozing), where only a few isolated α waves can be detected. Drowsiness further descends to sleep stage B (stage 1), where θ waves appear, then to stage C (stage 2), where burst of fast waves (sleep spindles) and isolated waves (K complexes) can be recorded, and ultimately to the stages of deen sleen (stages D/E = stages 3/4), characterized by the appearance of δ waves. Their amplitude increases while their frequency drops to a minimum in phase E $(\rightarrow p.333 D)$. This phase is therefore referred to as slow-wave sleep (SWS). The arousal threshold is highest about 1 hour after a person falls asleep. Sleep then becomes less deep and the first episode of rapid eye movement (REM) occurs. This completes the first sleep cycle. During REM sleep, most of the skeletal muscles become atonic (inhibition of motoneurons) while the breathing and heart rates increase. The face and fingers suddenly start to twitch, and penile erection and rapid eye movements occur. All other stages of sleep are collectively referred to as non-REM sleep (NREM). Sleepers aroused from REM sleep are more often able to describe their dreams than when aroused from NREM sleep. The sleep cycle normally lasts about 90 min and is repeated 4-5 times each night (\rightarrow p. 333 D). Towards morning, NREM sleep becomes shorter and more even. while the REM episodes increase from ca. 10 min to over 30 min.

Infants sleep longest (about 16 hours/day, 50% REM), 10-year-olds sleep an average 10 hours (20% REM), young adults sleep 7–8 hours a day, and adults over 50 sleep an average 6 hours or so (both 20% REM). The proportion of SWS clearly decreases in favor of stage C (stage 2) sleep.

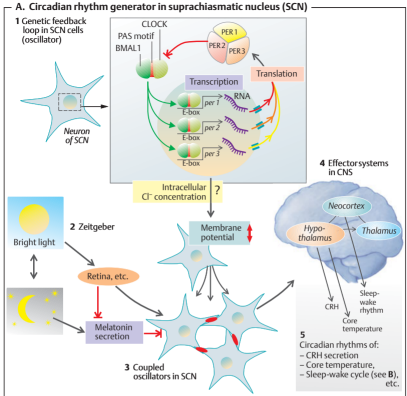
When a person is **deprived of REM sleep** (awakened during this phase), the duration of the next REM phase increases to compensate for the deficit. The first two to three sleep cycles (**core sleep**) are essential. Total **sleep deprivation** leads to death, but the reason is still unclear because too little is known about the physiological role of sleep.

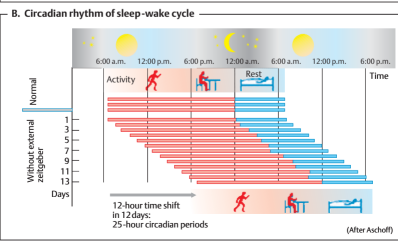
The daily sleep—wake cycle and other **circadian rhythms** (diurnal rhythms) are controlled by *endogenous rhythm generators*. The *central biological clock* (oscillator) that times these processes is located in the *suprachiasmatic nucleus* (**SCN**) of the hypothalamus (\rightarrow **A**). The endogenous circadian rhythm occurs in cycles of roughly 24–25 hours, but is unadulterated only when a person is completely isolated from the outside influences (e.g., in a windowless basement, dark cave, etc.). External *zeitgebers* (entraining signals) synchronize the biological clock to precise 24-hour cycles. It takes several days to "reset" the biological clock, e.g., after a long journey from east to west (*jet lag*).

Important genetic "cogwheels" of the central biological clock of mammals were recently discovered (→ A1). Neurons of the SCN contain specific proteins (CLOCK and BMAL1), the PAS domains of which bind to form heterodimers. The resulting CLOCK/BMAL1 complexes enter the cell nuclei, where their promoter sequences (E-box) bind to period (per) oscillator genes per1, per2, and per3, thereby activating their transcription. After a latency period, expression of the genes yields the proteins PER1, PER2, and PER3, which jointly function as a trimer to block the effect of CLOCK/BMAL1, thereby completing the negative feedback loop. The mechanism by which this cycle activates subsequent neuronal actions (membrane potentials) is still unclear.

The main **external zeitgeber** for 24-hour synchronization of the sleep-wake cycle is *bright light* (photic entrainment). Light stimuli are directly sensed by a small, *melanopsin*-containing fraction of retinal ganglion cells and conducted to the SCN via the retinohypothalamic tract (\rightarrow **A2,3**). The coupled cells of the SCN (\rightarrow **A3**) bring about circadian rhythms of hormone secretion, core temperature, and sleep-wake cycles (\rightarrow **A5, B**) by various effector systems of the CNS (\rightarrow **A4**).

The zeitgeber slows or accelerates the rhythm, depending on which phase it is in. Signals from the zeitgeber also reaches the epiphysis (pineal body, pineal gland) where it inhibits the secretion of melatonin which is high at night. Since it exerts its effects mainly on the SCN, administration of melatonin before retiring at night can greatly reduce the time required to "reset" the biological clock. The main reason is that it temporarily "deactivates" the SCN (via MT₂ receptors), thereby excluding most nocturnal neuronal input (except light stimuli).





Consciousness, Memory, Language

Consciousness. Selective attention, abstract thinking, the ability to verbalize experiences, the capacity to plan activities based on experience, self-awareness and the concept of values are some of the many characteristics of *consciousness*. Consciousness enables us to deal with difficult environmental conditions (adaptation). Little is known about the brain activity associated with consciousness and controlled attention (LCCS, see below), but we do know that subcortical activation systems such as the reticular formation (\rightarrow p. 322) and corticostriatal systems that inhibit the afferent signals to the cortex in the thalamus (\rightarrow p. 326) play an important role.

Attention. Sensory stimuli arriving in the sensory memory are evaluated and compared to the contents of the long-term memory within fractions of a second $(\rightarrow A)$. In routine situations such as driving in traffic, these stimuli are unconsciously processed (automated attention) and do not interfere with other reaction sequences such as conversation with a passenger. Our conscious, selective (controlled) attention is stimulated by novel or ambiguous stimuli, the reaction to which (e.g., the setting of priorities) is controlled by vast parts of the brain called the limited capacity control system (LCCS). Since our capacity for selective attention is limited, it normally is utilized only in stress situations.

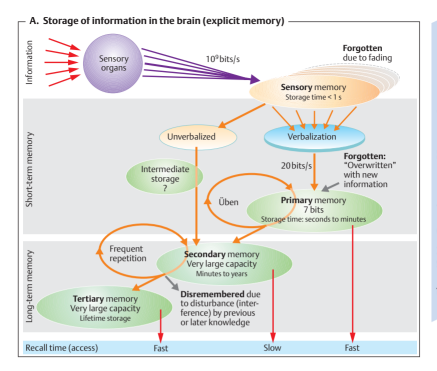
The **implicit memory** (procedural memory) stores skill-related information and information necessary for associative learning (conditioning of conditional reflexes; → p. 236) and non-associative learning (habituation and sensitization of reflex pathways). This type of unconscious memory involves the basal ganglia, cerebellum, motor cortex, amygdaloid body (emotional reactions) and other structures of the brain.

The **explicit memory** (declarative/knowledge memory) stores facts (semantic knowledge) and experiences (episodic knowledge, especially when experienced by selective attention) and *consciously* renders the data. Storage of information processed in the uniand polymodal association fields is the responsibility of the temporal lobe system (hip-

pocampus, perirhinal, entorhinal and parahippocampal cortex, etc.). It establishes the temporal and spatial *context* surrounding an experience and recurrently stores the information back into the *spines* of cortical dendrites in the association areas (\rightarrow p. 322). The recurrence of a portion of the experience then suffices to recall the contents of the memory.

Explicit learning $(\rightarrow A)$ starts in the sensorv memory, which holds the sensory impression automatically for less than 1 s. A small fraction of the information reaches the primary memory (short-term memory), which can retain about 7 units of information (e.g., groups of numbers) for a few seconds. In most cases. the information is also verbalized. Long-term storage of information in the secondary memory (long-term memory) is achieved by repetition (consolidation). The tertiary memory is the place where frequently repeated impressions are stored (e.g., reading, writing, one's own name); these things are never forgotten, and can be quickly recalled throughout one's lifetime. Impulses circulating in neuronal tracts are presumed to be the physiological correlative for short-term (primary) memory, whereas biochemical mechanisms are mainly responsible for long-term memory. Learning leads to long-term genomic changes. In addition, frequently repeated stimulation can lead to long-term potentiation (LTP) of synaptic connections that lasts for several hours to several days. The spines of dendrites in the cortex play an important role in LTP.

Mechanisms for LTP. Once receptors for AMPA (α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid) are activated by the presynaptic release of glutamate (→ p. 55 F), influxing Na⁺ depolarizes the postsynaptic membrane. Receptors for NMDA (Nmethyl-D-aspartic acid) are also activated, but the Ca²⁺ channels of the NMDA receptors are blocked by Mq2+, thereby inhibiting the influx of Ca2+ until the Mq2+ block is relieved by depolarization. The cytosolic Ca2+ concentration [Ca2+]; then rises. If this is repeated often enough, calmodulin mediates the autophosphorylation of CaM kinase II (\rightarrow p. 36), which persists even after the [Ca2+]i falls back to normal. CaM kinase II phosphorylates AMPA receptors (increases their conductivity) and promotes their insertion into the postsynaptic membrane, thereby enhancing synaptic transmission over longer periods of time (LTP).



Amnesia (memory loss). Retrograde amnesia (loss of memories of past events) is characterized by the loss of primary memory and (temporary) difficulty in recalling information from the secondary memory due to various causes (concussion, electroshock, etc.). Anterograde amnesia (inability to form new memories) is characterized by the inability to transfer new information from the primary memory to the secondary memory (Korsakoff's syndrome).

Language is a mode of *communication* used (1) to receive information through visual and aural channels (and through tactile channels in the blind) and (2) to transmit information in written and spoken form (see also p. 370). Language is also needed to form and verbalize *concepts and strategies* based on consciously processed sensory input. Memories can therefore be stored efficiently. The centers for formation and processing of concepts and language are unevenly distributed in the cerebral hemispheres. The left hemisphere is usually

the main center of speech in right-handed individuals ("dominant" hemisphere, large planum temporale), whereas the right hemisphere is dominant in 30–40% of all left-handers. The non-dominant hemisphere is important for word recognition, sentence melody, and numerous nonverbal capacities (e.g., music, spatial thinking, face recognition).

This can be illustrated using the example of patients in whom the two hemispheres are surgically disconnected due to conditions such as otherwise untreatable, severe epilepsy. If such a **split-brain patient** touches an object with the right hand (reported to the left hemisphere), he can name the object. If, however, he touches the object with the left hand (right hemisphere), he cannot name the object but can point to a picture of it. Since complete separation of the two hemispheres also causes many other severe disturbances, this type of surgery is used only in patients with otherwise unmanageable, extremely severe seizures.

Glia

The central nervous system contains around 10^{11} nerve cells and 10 times as many **glia cells** such as *oligodendrocytes*, *astrocytes*, *ependymal cells*, and *microglia* (\rightarrow **A**). **Oligodendrocytes** (ODC) form the myelin sheath that surrounds axons of the CNS (\rightarrow **A**).

Astrocytes (AC) are responsible for extracellular K+ and H+ homeostasis in the CNS. Neurons release K⁺ in response to high-frequency stimulation $(\rightarrow B)$. Astrocytes prevent an increase in the interstitial K+ concentration and thus an undesirable depolarization of neurons (see Nernst equation, Eq. 1.18, p. 32) by taking up K+, and intervene in a similar manner with H+ ions. Since AC are connected by gap junctions (\rightarrow p. 16ff.), they can transfer their K⁺ or H^+ load to nearby AC (\rightarrow **B**). In addition to forming a barrier that prevents transmitters from one synapse from being absorbed by another. AC also take transmitters up, e.g. glutamate (Glu). Intracellular Glu is converted to glutamine (GluNH₂), then transported out of the cell and taken up by the nerve cells, which convert it back to Glu (**transmitter recycling**; \rightarrow **B**).

Some AC have **receptors for transmitters** such as Glu, which triggers a Ca²⁺ wave from one AC to another. Astrocytes are also able to modify the Ca²⁺ concentration in the neuronal cytosol so that the two cell types can "communicate" with each other. AC also mediate the transport of materials between capillaries and neurons and play an important part in *energy homeostasis* of the neurons by mediating glycogen synthesis and breakdown.

During embryonal development, the long processes of AC serve as guiding structures that help undifferentiated nerve cells migrate to their target areas. Glia cells also play an important role in CNS development by helping to control gene expression in nerve cell clusters with or without the aid of growth factors such as NGF (nerve growth factor), BDGF (brain-derived growth factor), and GDNF (glial cell line-derived neurotropic factor). GDNF also serves as a trophic factor for all mature neurons. Cell division of glia cells can lead to in scarring (epileptic foci) and tumor formation (glioma).

Immunocompetent **microglia** (\rightarrow **A**) assume many functions of macrophages outside the CNS when CNS injuries or infections occur (\rightarrow p. 94ff.). **Ependymal cells** line internal hollow cavities of the CNS (\rightarrow **A**).

Sense of Taste

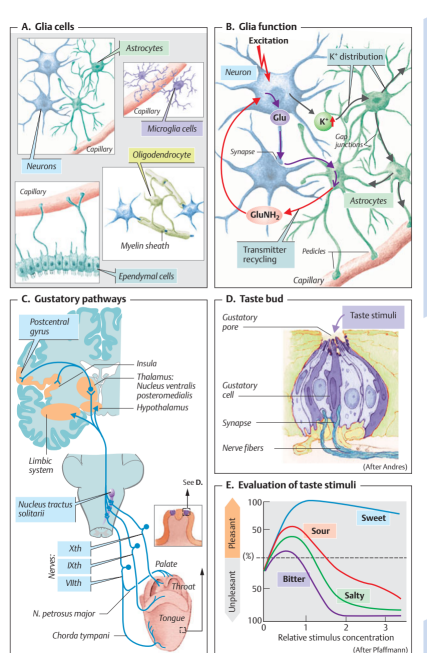
Gustatory pathways. The taste buds $(\rightarrow D)$ consist of clusters of 50–100 secondary sensory cells on the tongue (renewed in 2-week cycles); humans have around 5000 taste buds. Sensory stimuli from the taste buds are conducted to endings of the VIIth, IXth and Xth cranial nerves, relayed by the nucleus tractus solitarii, and converge at a high frequency on (a) the postcentral gyrus via the thalamus $(\rightarrow p. 323 \, B, \text{ "tongue"})$ and (b) the hypothalamus and limbic system via the pons $(\rightarrow C)$.

The **qualities of taste** distinguishable in humans are conventionally defined as *sweet*, *sour*, *salty*, and *bitter*. The specific taste sensor cells for these qualities are distributed over the whole tongue but differ with respect to their densities. *Umami*, the sensation caused by monosodium-L-glutamate (MSG), is now classified as a fifth quality of taste. MSG is chiefly found in protein-rich foods.

Taste sensor cells distinguish the types of taste as follows: Salty: Cations (Na+, K+, etc.) taste salty, but the presence of anions also plays a role. E.g., Na+ enters the taste sensor cell via Na+ channels and depolarizes the cell. Sour: H⁺ ions lead to a more frequent closure of K+ channels, which also has a depolarizing effect. Bitter: A family of > 50 genes codes for an battery of bitter sensors. A number of sensory proteins specific for a particular substance are expressed in a single taste sensor cell, making it sensitive to different bitter tastes. The sensory input is relaved by the G-protein α -austducin. No nuances but only the overall warning signal "bitter" is perceived. Umami: Certain taste sensor contain a metabotropic glutamate receptor, mGluR4, the stimulation of which leads to a drop in cAMP conc.

Taste thresholds. The threshold (mol/L) for recognition of taste stimuli applied to the tongue is roughly 10^{-5} for quinine sulfate and saccharin, 10^{-3} for HCl, and 10^{-2} for sucrose and NaCl. The relative *intensity differential threshold* $\Delta I/I$ (\rightarrow p. 352) is about 0.20. The concentration of the gustatory stimulus determines whether its taste will be perceived as pleasant or unpleasant (\rightarrow E). For the *adaptation* of the sense of taste, see p. 341 C.

Function of taste. The sense of taste has a protective function as spoiled or bitter-tasting food (low taste threshold) is often poisonous. Tasting substances also stimulate the secretion of saliva and gastric juices (→ pp. 236, 242).



Sense of Smell

The neuroepithelium of the olfactory region contains ca. 10^7 primary olfactory sensor cells (\rightarrow A1) which are bipolar neurons. Their dendrites branch to form 5–20 mucus-covered cilia, whereas the axons extend centrally in bundles called *fila olfactoria* (\rightarrow A1,2). Olfactory neurons are replenished by basal cell division in 30–60-day cycles. *Free nerve endings* (trigeminal nerve) in the nasal mucosa also react to certain aggressive odors (e.g., acid or ammonia vapors).

Olfactory sensors. Odorant molecules (Mr 15-300) are transported by the inhaled air to the olfactory region, where they first dissolve in the mucous lining before combining with **receptor proteins** in the *cilial membrane*. These are coded by a huge family of genes (500-750 genes distributed in most chromosomes), whereby probably one olfactory sensor cell only expresses one of these genes. Since only a part of the sequence of about 40% of these genes is expressed, humans have roughly 200-400 different sensor cell types. Olfactory receptors couple with G_s-proteins (G_{olf} proteins; → B and p. 274ff.) that increase the conductivity of the sensor cell membrane to cations, thereby increasing the influx of Na+ and Ca2+ and thus depolarizing the cell.

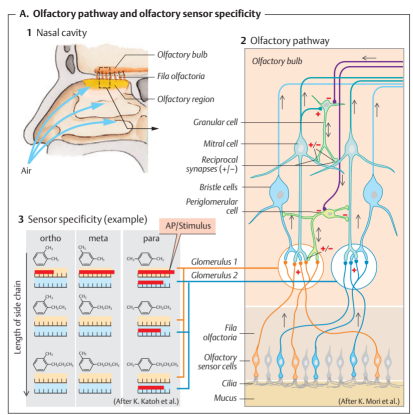
Sensor specificity (→ A3). Olfactory sensor cells recognize a very specific structural feature of the odorant molecules they are sensitive to. The cloned receptor 17 of the rat, for example, reacts with the aldehyde n-octanal but not to octanol, octanoic acid, or aldehydes which have two methyl groups more or less than n-octanal. In the case of aromatic compounds, one sensor recognizes whether the compound is ortho, meta or para-substituted, while another detects the length of the substituent regardless of where it is located on the ring. The different molecular moieties of an odorant molecule therefore activate different types of sensors (\rightarrow A3, top right). Jasmine leaves and wine contain several dozens and hundreds of odorants, respectively, so their overall scent is a more complex perception (integrated in the rhinencephalon).

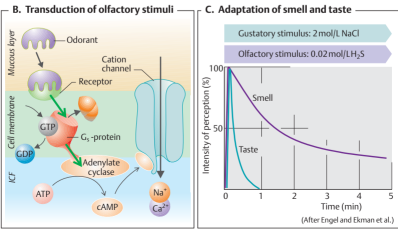
Olfactory pathway (\rightarrow A2). Axons of (ca. 10³) same-type sensors distributed over the olfactory epithelium synapse to dendrites of their respective *mitral cells* (MC) and *bristle cells* (BC) within the glomeruli olfactorii of the

olfactory bulb. The glomeruli therefore function as convergence centers that integrate and relay signals from the same sensor type. Their respective sensor protein also determines which glomerulus newly formed sensor axons will connect to. Periglomerular cells and granular cells connect and inhibit mitral and bristle cells (\rightarrow A2). Mitral cells act on the same reciprocal synapses ($\rightarrow A$. "+/-") in reverse direction to activate the periglomerular cells and granular cells which, on the other hand, are inhibited by efferents from the primary olfactory cortex and contralateral anterior olfactory nucleus (→ A2, violet tracts). These connections enable the cells to inhibit themselves or nearby cells (contrast), or they can be disinhibited by higher centers. The signals of the axons of mitral cells (1) reach the anterior olfactory nucleus. Its neurons cross over (in the anterior commissure) to the mitral cells of the contralateral bulb and (2) form the olfactory tract projecting to the primary olfactory cortex (prepiriform cortex, tuberculum olfactorium, nucleus corticalis amygdalae). The olfactory input processed there is relayed to the hypothalamus, limbic system (see also p. 330), and reticular formation; it is also relayed to the neocortex (insula, orbitofrontal area) either directly or by way of the thalamus.

Thresholds. It takes only 4×10^{-15} g of methylmercaptan (in garlic) per liter of air to trigger the vague sensation of smell (*perception* or *absolute threshold*). The odor can be properly identified when 2×10^{-13} g/L is present (*identification threshold*). Such thresholds are affected by air temperature and humidity; those for other substances can be 10^{10} times higher. The relative intensity differential threshold $\Delta I/I$ (0.25) is relatively high (\rightarrow p.352). *Adaptation* to smell is sensor-dependent (desensitization) and neuronal (\rightarrow C).

The sense of smell has various **functions**. Pleasant smells trigger the secretion of saliva and gastric juices, whereas unpleasant smells warn of potentially spoiled food. Body odor permits hygiene control (sweat, excrement), conveys social information (e.g., family, enemy; \rightarrow p. 330), and influences sexual behavior. Other aromas influence the emotional state.





Sense of Balance

Anatomy. Each of the three semicircular canals $(\rightarrow$ A1) is located in a plane about at right angles to the others. The ampulla of each canal contains a ridge-like structure called the *crista ampullaris* $(\rightarrow$ A2). It contains *hair cells* (secondary sensory cells), the *cilia* of which $(\rightarrow$ A3) project into a gelatinous membrane called the *cupula* $(\rightarrow$ A2). Each hair cell has a long *kinocilium* and ca. 80 streeocilia of variable length. Their tips are connected to longer adjacent cilia via the "tip links" $(\rightarrow$ A3).

Semicircular canals. When the cilia are in a resting state, the hair cells release a transmitter (glutamate) that triggers the firing of action potentials (AP) in the nerve fibers of the vestibular ganglion. When the head is turned, the semicircular canal automatically moves with it, but endolymph in the canal moves more sluggishly due to inertia. A brief pressure difference thus develops between the two sides of the cupula. The resultant vaulting of the cupula causes the stereocilia to bend $(\rightarrow A2)$ and shear against each other, thereby changing the cation conductance of the hair cell membrane. Bending of the stereocilia towards the kinocilium increases conductivity and allows the influx of K⁺ and Na⁺ along a high electrochemical gradient between the endolymph and hair cell interior (see also pp. 366 and 369 C). Thus, the hair cell becomes depolarized, Ca2+ channels open, more glutamate is released, and the AP frequency increases. The reverse occurs when the cilia bend in the other direction (away from the kinocilium). The semicircular canals function to detect angular (rotational) accelerations of the head in all planes (rotation, nodding, tilting sideways). Since normal head movements take less than $0.3 \, \text{s}$ (acceleration) \Rightarrow deceleration), stimulation of the semicircular canals usually reflects the rotational velocity.

The pressure difference across the cupula disappears when the **body rotates for longer periods** of time. Deceleration of the rotation causes a pressure gradient in the opposite direction. When bending of the cilia increased the AP frequency at the start of rotation, it decreases during deceleration and vice versa. Abrupt cessation of the rotation leads to **vertigo** and **nystagmus** (see below).

The **saccule** and **utricle** contain **maculae** (\rightarrow **A1**, **A4**) with cilia that project into a gelatinous

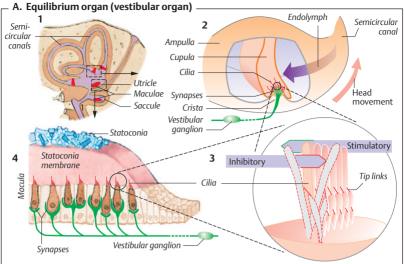
membrane (\rightarrow A4) with high density (\approx 3.0) calcite crystals called *statoconia*, *statoliths* or *otoliths*. They displace the membrane and thereby bend the embedded cilia (\rightarrow A4) due to changes of the direction of *gravity*, e.g. when the head position deviates from the perpendicular axis. The maculae respond also to other *linear* (*translational*) *accelerations*, e.g. of a car or an elevator.

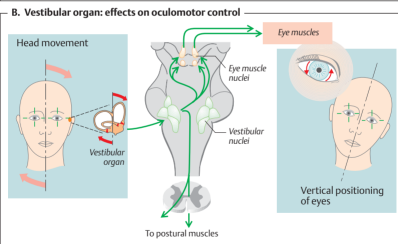
Central connections. The bipolar neurons of the vestibular ganglion synapse with the *vestibular nuclei* (\rightarrow **A**, **B**). Important tracts extend from there to the contralateral side and to *ocular muscle nuclei*, *cerebellum* (\rightarrow **p**. 326), *motoneurons* of the skeletal muscles, and to the *postcentral gyrus* (conscious spatial orientation). **Vestibular reflexes** (a) maintain the balance of the body (*postural motor function*, \rightarrow **p**. 328) and (b) keep the visual field in focus despite changes in head and body position (*oculomotor control*, \rightarrow **B** and p. 360).

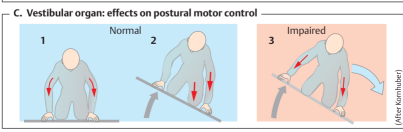
Example (\rightarrow C): If a support holding a test subject is tilted, the activated vestibular organ prompts the subject to extend the arm and thigh on the declining side and to bend the arm on the inclining side to maintain balance (\rightarrow C2). The patient with an impaired equilibrium organ fails to respond appropriately and topples over (\rightarrow C3).

Since the vestibular organ cannot determine whether the head alone or the entire body moves ($sense\ of\ movement$) or changed position (postural sense), the vestibular nuclei must also receive and process visual information and that from propriosensors in the neck muscles. Efferent fibers project bilaterally to the eye muscle nuclei, and any change in head position is immediately corrected by opposing $eye\ movement\ (\rightarrow B)$. This $vestibulo-ocular\ reflex\ maintains\ spatial\ orientation$.

Vestibular organ function can be assessed by testing oculomotor control. Secondary or postrotatory **nystagmus** occurs after abrupt cessation of prolonged rotation of the head around the vertical axis (e.g., in an office chair) due to activation of the horizontal semicircular canals. It is characterized by slow horizontal movement of the eyes in the direction of rotation and rapid return movement. Rightward rotation leads to left nystagmus and vice versa (\rightarrow p. 360). Caloric stimulation of the horizontal semicircular canal by instilling cold (30 °C) or warm water (44 °C) in the auditory canal leads to caloric nystagmus. This method can be used for unilateral testing.







Eye Structure, Tear Fluid, Aqueous Humor

Light entering the eve must pass through the cornea, aqueous humor, lens and vitreous body, which are collectively called the optical apparatus, before reaching the retina and its light-sensitive photosensors $(\rightarrow A)$. This produces a reduced and inverse image of the visual field on the retina. All parts of the apparatus must be transparent and have a stable shape and smooth surface to produce an undistorted image, which is the main purpose of tear fluid in case of the cornea. Tears are secreted by lacrimal glands located in the top outer portion of orbit and their mode of production is similar to that of saliva (\rightarrow p. 236). Tears are distributed by reflex blinking and then pass through the lacrimal puncta and lacrimal canaliculi (or ducts) of the upper and lower evelid into the lacrimal sac and finally drain into the nasal sinuses by way of the nasolacrimal duct. Tear fluid improves the optical characteristics of the cornea by smoothing uneven surfaces, washing away dust, protecting it from caustic vapors and chemicals, and protects it from drying out. Tears lubricate the eyelid movement and contain lysozyme and immunoglobulin A (\rightarrow pp. 96ff. and 232), which help ward off infections. In addition, tears are a well known mode of expressing emotions.

The entry of light into the eye is regulated by the **iris** (\rightarrow **A**; p. 353 C1), which contains annular and radial smooth muscle fibers. Cholinergic activation of the sphincter muscle of pupil leads to pupil contraction (*miosis*), and adrenergic activation of the dilator muscle of pupil results in pupil dilatation (*mydriasis*).

The **bulbus** (eyeball) maintains its shape due to its tough outer coat or **sclera** (\rightarrow **C**) and **intraocular pressure** which is normally 10–21 mmHg above the atmospheric pressure. The drainage of **aqueous humor** must balance its production to maintain a constant ocular pressure (\rightarrow **C**). Aqueous humor is produced in the *ciliary process* of the posterior ocular chamber with the aid of carbonic anhydrase and active ion transport. It flows through the pupil into the anterior ocular chamber and drains into the venous system by way of the trabecu-

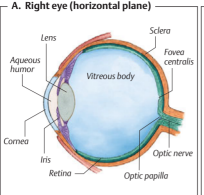
lar meshwork and Schlemm's canal. Aqueous humor is renewed once every hour or so.

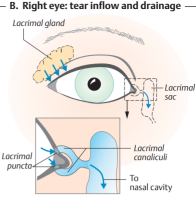
Glaucoma. Obstruction of humor drainage can occur due to chronic obliteration of the trabecular meshwork (open-angle glaucoma) or due to acute block of the anterior angle (angle-closure glaucoma) leading to elevated intraocular pressure, pain, retinal damage, and blindness. Drugs that decrease humor production (e.g. carbonic anhydrase inhibitors) and induce meiosis are used to treat glaucoma.

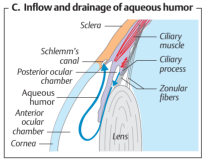
The **lens** is held in place by the *ciliary zonules* $(\rightarrow C)$. When the eye adjusts for *far vision*, the zonules are stretched and the lens becomes flatter, especially its anterior surface $(\rightarrow D,$ top). When looking at nearby objects (*near vision*), the zonules are relaxed due to contraction of the ciliary muscle, and the lens reassumes its original shape due to its elasticity $(\rightarrow D,$ bottom, and p. 346).

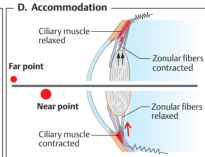
The **retina** lines the interior surface of the bulbus except the anterior surface and the site where the *optical nerve* (\rightarrow **A**) exits the bulbus via the *optic papilla* (\rightarrow **A**). The *fovea centralis* (\rightarrow **A**) forms a slight depression across from the pupillary opening. The retina consists of several layers, named from inside out as follows (\rightarrow **E**): pigmented epithelium, photosensors (rods and cones), Cajal's horizontal cells, bipolar cells, amacrine cells, and ganglion cells. The central processes of the ganglion cells ($n \approx 10^6$) exit the bulbus as the optical nerve (retinal circuitry; \rightarrow p. 355ff.).

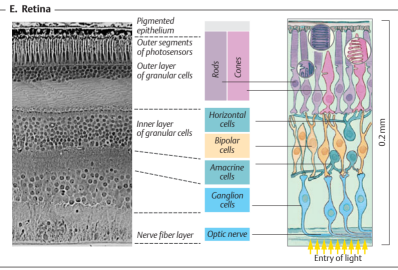
Photosensors. Retinal rods and cones have a light-sensitive outer segment, which is connected to a inner segment by a thin connecting part (\rightarrow p. 349 C1). The inner segment contains the normal cell organelles and establishes synaptic contact with the neighboring cells. The outer segment of the rod cells contains ca. 800 membranous disks, and the plasma membrane of the outer segment of the cones is folded. Visual pigments are stored in these disks and folds (\rightarrow p. 348). The outer segment is continuously regenerated; the old membranous disks at the tip of the cell are shed and replaced by new disks from the inner segment. The phagocytic cells of the pigmented epithelium engulf the disks shed by the rods in the morning, and those shed by the cones in the evening. Some ganglion cells contain a light-sensitive pigment (\rightarrow p. 334).











Optical Apparatus of the Eye

Physics. The production of an **optical image** is based on the refraction of light rays crossing a spherical interface between air and another medium. Such a **simple optical system** illustrated in plate **A** has an anterior focal point (F_a) in air, a posterior focal point (F_p), a principal point (P_p), and a nodal point (P_p). Light rays from a distant point (P_p) can be regarded as parallel. If they enter the system parallel to its optical axis, they will converge at $F_p \leftarrow A1$, red dot). If they enter at an angle to the axis, then they will form an image beside F_p but in the same focal plane ($\rightarrow A1$, violet dot). Light rays from a nearby point do not enter the system in parallel and form an image behind the focal plane ($\rightarrow A2$, green and brown dots).

The **optical apparatus** of the eye (\rightarrow p. 344) consists of multiple interfaces and media, and is therefore a *complex optical system*. It can, however, be treated as *a simple optical system*. Light rays from a focused object (O) pass through N and diverge at angle α until they reach the retina and form an image (I) there (\rightarrow A2).

Two points separated by a distance of 1.5 mm and located 5 m away from the eye (tan α = 1.5/5000; α = 0.0175 degrees \approx 1) will therefore be brought into focus 5 μ m apart on the retina. In a person with normal vision (\rightarrow p. 348), these two points can be distinguished as separate because 5 μ m corresponds to the diameter of three cones in the fovea (two are stimulated, the one in between is not).

Accommodation. When the eyes are adjusted for **far vision**, parallel light rays from a distant point meet at $F_p (\rightarrow B1, \text{red dot})$. Since the retina is also located at F_p , the distant point is clearly imaged there. The eye adjusted for far vision will not form a clear image of a nearby point (the light rays meet behind the retina, $\rightarrow B1$, green dot) until the eye has adjusted for **near vision**. In other words, the curvature of the lens (and its refractive power) increases and the image of the nearby point moves to the retinal plane ($\rightarrow B2$, green dot). Now, the distant point cannot not be sharply imaged since F_p does not lie in the retinal plane any more ($\rightarrow B2$).

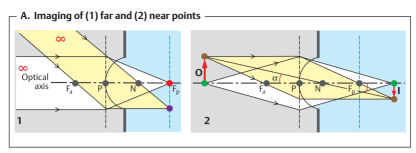
The **refractive power** around the edge of the optical apparatus is higher than near the optical axis. This *spherical aberration* can be minimized by narrowing the pupils. The refractive power of the eye is the reciprocal of the ante-

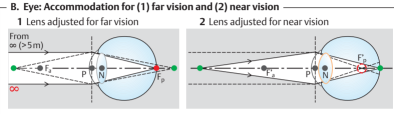
rior focal length in meters, and is measured in diopters (dpt). In accommodation for far vision, focal length = anterior focal point (F_a)-principal point (P) = 0.017 m (\rightarrow **B1**). Thus, the corresponding refractive power is 1/0.017 = 58.8dpt, which is mainly attributable to refraction at the air-cornea interface (43 dpt). In maximum accommodation for near vision in a young person with normal vision (emmetropia), the refractive power increases by around 10-14 dpt. This increase is called range of accommodation and is calculated as 1/near point - 1/far point $[m^{-1} = dpt]$. The **near point** is the closest distance to which the eye can accommodate; that of a young person with normal vision is 0.07–0.1 m. The far point is infinity (∞) in subjects with normal vision. The range of accommodation to a near point of 0.1 m is therefore 10 dpt since $1/\infty = 0$. It decreases as we grow older (to 1-3.5 dpt in 50-year-olds) due to the loss of elasticity of the lens. This visual impairment of aging, called presbyopia (\rightarrow C1-3), normally does not affect far vision, but convex lenses are generally required for near vision, e.g., reading.

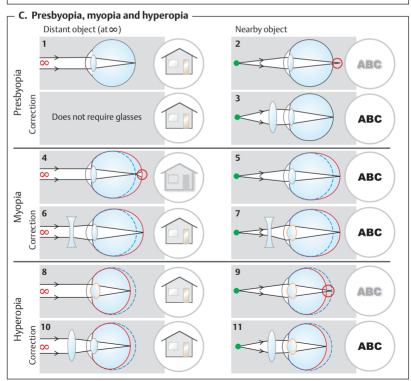
Cataract causes opacity of the lens of one or both eyes. When surgically treated, convex lenses (glasses or artificial intraocular lenses) of at least +15 dpt must be used to correct the vision.

In myopia (near-sightedness), rays of light entering the eye parallel to the optical axis are brought to focus in front of the retina because the eyeball is too long (\rightarrow C4). Distant objects are therefore seen as blurred because the far point is displaced towards the eyes $(\rightarrow C5)$. Myopia is corrected by concave lenses (negative dpt) that disperse the parallel light rays to the corresponding extent (\rightarrow **C6**). Example: When the far point = $0.5 \, \text{m}$, a lens of [-1/0.5] = [-2]dpt] will be required for correction (\rightarrow C7). In hyperopia (far-sightedness), on the other hand, the eyeball is too short. Since the accommodation mechanisms for near vision must then be already used to focus distant objects (\rightarrow C8), the range of accommodation no longer suffices to clearly focus nearby objects (→ C9). Hyperopia is corrected by convex lenses $(+ dpt) (\rightarrow C10-11).$

Astigmatism. In regular astigmatism, the corneal surface is more curved in one plane (usually the vertical: astigmatism with the rule) than the other, creating a difference in refraction between the two planes. A point source of light is therefore seen as a line or oval. Regular astigmatism is corrected by cylindrical lenses. Irregular astigmatism (caused by scars, etc.) can be corrected by contact lenses.







Visual Acuity, Photosensors

Visual acuity is an important measure of eye function. Under *well-lighted conditions*, the normal eye should be able to distinguish two points as separate when the light rays emitted by the point objects converge at an angle (α) of 1 min (1/60 degree) $(\rightarrow \mathbf{A} \text{ and p. 346})$. Visual acuity is calculated as $1/\alpha(\text{min}^{-1})$, and is 1/1 in subjects with normal vision.

Visual acuity testing is generally performed using charts with letters or other optotypes (e.g., Landolt rings) of various sizes used to simulate different distances to the test subject. The letters or rings are usually displayed at a distance of $5 \,\mathrm{m} \, (\rightarrow \, A)$. Visual acuity is normal (1/1) if the patient recognizes letters or ring openings seen at an angle of 1 min from a distance of $5 \,\mathrm{m} \, .Example$: It should be possible to identify the direction of the opening of the middle ring from a distance of $5 \,\mathrm{m} \, .A$). If only the opening of the left ring can be localized from the test distance of $5 \,\mathrm{m} \, .D$, the visual acuity is $5/8.5 \,\mathrm{m} \, .D$.

Photosensors or photoreceptors. The lightsensitive sensors of the eye consist of approximately $6 \cdot 10^6$ rods and 20 times as many cones (→p.345 E) distributed at variable densities throughout the retina (\rightarrow **B1**). (Certain ganglion cells also contain a light-sensitive pigment; \rightarrow p. 334). The fovea centralis is exclusively filled with cones, and their density rapidly decreases towards the periphery. Rods predominate 20-30 degrees away from the fovea centralis. Approaching the periphery of the retina, the density of the rods decreases continuously from $1.5 \times 10^5 / \text{mm}^2$ (maximum) to about one-third this value. No photosensors are present on the optic disk, which is therefore referred to as the **blind spot** in the visual field.

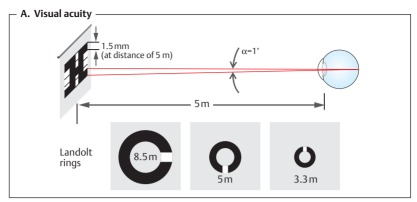
Clear visualization of an object in daylight requires that the gaze be fixed on it, i.e., that an image of the object is produced in the *fovea centralis*. Sudden motion in the periphery of the visual field triggers a **reflex saccade** (\rightarrow p. 360), which shifts the image of the object into the fovea centralis. Thereby, the retinal area with the highest visual acuity is selected (\rightarrow B2, yellow peak), which lies 5 degrees temporal to the optical axis. Visual acuity decreases rapidly when moving outward from the fovea (\rightarrow B2, yellow field), reflecting the decreasing density of cone distribution (\rightarrow B1,

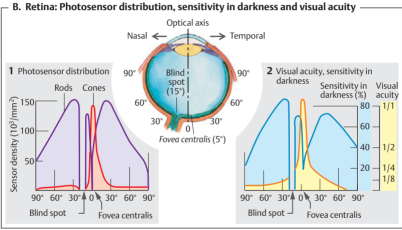
red curve). In a *dark-adapted eye*, on the other hand, the sensitivity of the retina (\rightarrow **B2**, blue curve) is completely dependent on the rod distribution (\rightarrow **B1**, purple curve). The color-sensitive cones are therefore used for visual perception in daylight or good lighting (*day vision*, *photopic vision*), while the black and whitesensitive cones are used to visualize objects in darkness (*dim-light vision*, *night vision*, *scotoptic vision*). The high light sensitivity in night vision is associated with a high loss of visual acuity (\rightarrow p. 354).

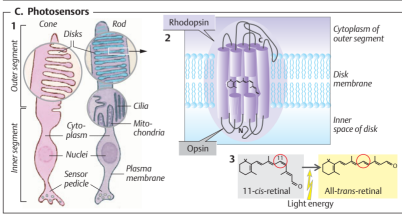
Photosensor Function

Light-absorbing visual pigments and a variety of enzymes and transmitters in retinal rods and cones $(\rightarrow C1)$ mediate the conversion of light stimuli into electrical stimuli: this is called photoelectric transduction. The membranous disks of the retinal rods contain rhodopsin $(\rightarrow C2)$, a photosensitive purple-red chromoprotein (visual purple). Rhodopsin consists of the integral membrane protein opsin and the aldehyde 11-cis-retinal. The latter is bound to a lysine residue of opsin which is embedded in this protein; it is stably kept in place by weak interactions with two other amino acid residues. Photic stimuli trigger a primary photochemical reaction in rhodopsin (duration, 2 · 10⁻¹⁴ s) in which 11-cis-retinal is converted to all-trans-retinal (\rightarrow C3). Even without continued photic stimulation, the reaction yields bathorhodopsin, the intermediates lumirhodopsin and metarhodopsin I, and finally metarhodopsin II within roughly 10^{-3} s (\rightarrow **D1**).

Metarhodopsin II (MR-II) reacts with a G_s -protein (→ p. 274) called transducin (G_t -protein), which breaks down into α_s and $\beta\gamma$ subunits once GDP has been replaced by GTP (→ D1). Activated α_s -GTP now binds the inhibitory subunit of cGMP phosphodiesterase (I_{PDE}) (→ D2). The consequently disinhibited phosphodiesterase (PDE) then lowers the cytosolic concentration of cyclic guanosine monophosphate (cGMP). The activation of a single retinal rhodopsin molecule by a quantum of light can induce the hydrolysis of up to 10^6 cGMP molecules per second. The reaction cascade therefore has tremendous amplifying power.







In darkness (\rightarrow D, left), cGMP is bound to cation channels (Na+, Ca2+) in the outer segment of the photosensor, thereby keeping them open. Na⁺ and Ca²⁺ can therefore enter the cell and depolarize it to about -40 mV (→ D3. D4). This darkness-induced influx into the outer segment is associated with the efflux of K+ from the inner segment of the sensor. The Ca2+ entering the outer segment is immediately transported out of the cell by a 3 Na⁺/Ca²⁺ exchanger (\rightarrow p. 36), so the cytosolic Ca²⁺ concentration [Ca 2+]; remains constant at ca. 350-500 nmol/L in darkness (\rightarrow **D6**). If the cytosolic cGMP concentration decreases in response to a **light stimulus** (\rightarrow **D2**), cGMP dissociates from the cation channels, allowing them to close. The photosensor then hyperpolarizes to ca. – 70 mV (sensor potential: → D, right). This inhibits the release of glutamate (transmitter) at the sensor pedicle (\rightarrow **D5**), which subsequently causes changes in the membrane potential in downstream retinal neurons (\rightarrow p. 354).

Deactivation of Photic Reactions and Regeneration Cycles

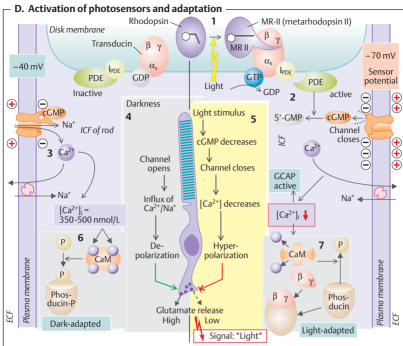
- ◆ Rhodopsin (→E2). Rhodopsin kinase (RK) competes with transducin for bindings sites on metarhodopsin II (MR-II); the concentration of transducin is 100 times higher (→E2, right). Binding of RK to MR-II leads to phosphorylation of MR-II. As a result, its affinity to transducin decreases while its affinity to another protein, arrestin, rises. Arrestin blocks the binding of further transducin molecules to MR-II. All-trans-retinal detaches from opsin, which is subsequently dephosphorylated and re-loaded with 11-cis-retinal.
- ◆ All-trans-retinal (→ E1) is transported out of the photosensor and into the pigmented epithelium, where it is reduced to all-transretinol, esterified and ultimately restored to 11-cis-retinal. After returning into the photosensor, it binds to opsin again, thereby completing the cycle (→ E2).
- **Note:** Retinol is vitamin A₁. A chronic deficiency of vitamin A₁ or its precursors (carotinoids) leads to impaired rhodopsin production and, ultimately, to *night blindness* (— p. 352).
- ♦ **Transducin** (→ **E3**). Since the GTPase activity of α_s -GTP breaks down GTP into GDP + P_i, the molecule deactivates itself. The α_s -GTP molecule and βγ subunit then reunite to trans-

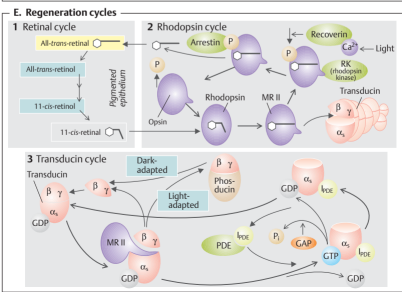
- ducin. GAP (GTPase-activating protein) accelerates the regeneration of transducin. *Phosducin*, another protein, is phosphorylated in the dark (\rightarrow **D6**) and dephosphorylated in light (\rightarrow **D7**). The latter form binds to the $\beta\gamma$ subunit (\rightarrow **D7**, **E3**), thereby blocking the regeneration of transducin. This plays a role in light adaptation (see below).
- Phosphodiesterase (PDE). In the course of transducin regeneration, the inhibitory subunit of cGMP phosphodiesterase (I_{PDE}) is released again and PDE is thus inactivated.
- ◆ cGMP. Since the 3 Na⁺/Ca²⁺ exchanger still functions even after photostimulation-induced closure of Ca²⁺ channels, the [Ca²⁺]_i starts to decrease. When a threshold of ca. 100 nmol/L is reached, the Ca²⁺-binding protein GCAP (guanylyl cyclase-activating protein) loses its 4 Ca²⁺ ions and stimulates guanylyl cyclase, thereby accelerating cGMP synthesis. Thus, the cGMP concentration rises, the cation channels re-open, and the sensor is ready to receive a new light stimulus. This Ca²⁺ cycle therefore mediates a negative feedback loop for cGMP production.

Ca2+ Ions and Adaptation (see also p. 352)

In the dark, the $[Ca^{2+}]_i$ is high, and calmodulinbound Ca^{2+} (\rightarrow p. 36) stimulates the phosphorylation of phosducin with the aid of cAMP and phosphokinase A (\rightarrow D6). In light, the $[Ca^{2+}]_i$ is low; phosducin is dephosphorylated and rapid regeneration of transducin is not possible (\rightarrow D7, E3). Moreover, Ca^{2+} accelerates the phosphorylation of MR-II in light with the aid of another Ca^{2+} binding protein, *recoverin* (\rightarrow E2). Ca^{2+} is therefore essential for the adaptation of photosensors (\rightarrow p. 352).

Although they contain similar enzymes and transmittes, the photosensitivity of the **cones** is about 100 times less than that of the rods. Thus, the cones are unable to detect a single quantum of light, presumably because photic reactions in the cones are deactivated too quickly. Compared to rhodopsin in the retinal rods, the three visual pigments in the three types of cones (11-cis-retinal with different opsin fractions) only absorb light in a narrow wavelength range (\rightarrow p. 357 E), which is a prerequisite for color vision (\rightarrow p. 356).





Adaptation of the Eye to Different Light Intensities

The eye is able to perceive a wide range of light intensities, from the extremely low intensity of a small star to the extremely strong intensity of the sun glaring on a glacier. The ability of the eye to process such a wide range of luminance (1:10¹¹) by adjusting to the prevailing light intensity is called adaptation. When going from normal daylight into a darkened room, the room will first appear black because its luminance value (measured in cd·m-2) is lower than the current ocular threshold. As the stimulus threshold decreases over the next few minutes, the furniture in the room gradually becomes identifiable. A longer period of adaptation is required to visualize stars. The maximum level of adaptation is reached in about 30 min (\rightarrow **A**). The minimum light intensity that can just be detected after maximum dark adaptation is the absolute visual threshold, which is defined as 1 in A and B.

The **retinal adaptation curve** exhibits a (Kohlrausch) break at roughly $2000 \times$ the absolute threshold (\rightarrow **A**, blue curve). This corresponds to the point where the excitation threshold of the cones is reached (threshold for day vision). The remainder of the curve is governed by the somewhat slower adaptation of the rods (\rightarrow **A**, violet curve). The isolated rod adaptation curve can be recorded in patients with complete color blindness (rod monochromatism), and the isolated cone adaptation curve can be observed in night blindness (hemeralopia, \rightarrow p. 350).

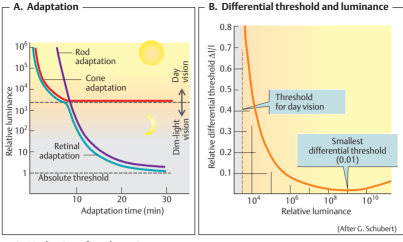
Differential threshold (or difference limen). The ability of the eye to distinguish the difference between two similar photic stimuli is an important prerequisite for proper eyesight. At the lowest limit of discriminative sensibility for two light intensities I and I', the absolute differential threshold (Δ I) is defined as I minus I'. The relative differential threshold is calculated as Δ I/I, and remains relatively constant in the median stimulus range (Weber's rule). Under optimal lighting conditions (approx. 10^9 times the absolute threshold; \rightarrow B), Δ I/I is very small (0.01). The relative differential threshold rises greatly in dark adaptation, but also rises in response to extremely bright light. Sun-

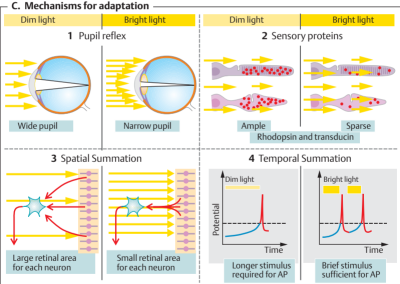
glasses decrease the differential threshold in the latter case.

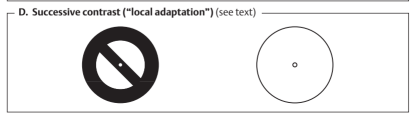
The **mechanisms for adaptation** of the eye are as follows $(\rightarrow C)$:

- ♦ **Pupil reflex** (\rightarrow **C1**). Through reflexive responses to light exposure of the retina (\rightarrow p. 359), the pupils can adjust the quantity of light entering the retina by a factor of 16. Thus, the pupils are larger in darkness than in daylight. The main function of the pupil reflex is to ensure rapid adaptation to sudden changes in light intensity.
- Chemical stimuli (→C2) help to adjust the sensitivity of photosensors to the prevailing light conditions. Large quantities of light lead to a prolonged decrease in the receptor's cytosolic Ca2+ concentration. This in conjunction with the activity of recoverin and phosducin reduces the availability of rhodopsin $(\rightarrow p. 348ff.)$. It therefore decreases the probability that a rhodopsin molecule will be struck by an incoming light ray (photon) or that a metarhodopsin II molecule will come in contact with a transducin molecule. When the light intensity is low, large concentrations of rhodopsin and transducin are available and the photosensors become very light-sensitive.
- ◆ Spatial summation (→ C3). Variation of retinal surface area (number of photosensors) exciting an optic nerve fiber causes a form of spatial summation that increases with darkness and decreases with brightness (→ p. 354).
- ◆ Temporal summation (→C4). Brief subthreshold stimuli can be raised above threshold by increasing the duration of stimulation (by staring at an object) long enough to trigger an action potential (AP). Thereby, the product of stimulus intensity times stimulus duration remains constant.

Successive contrast occurs due to "local adaptation." When a subject stares at the center of the black-and-white pattern $(\rightarrow \mathbf{D})$ for about 20 s and suddenly shifts the focus to the white circle, the previously dark areas appear to be brighter than the surroundings due to sensitization of the corresponding areas of the retina







Retinal Processing of Visual Stimuli

Light stimuli hyperpolarize the sensor potential of photosensors (\rightarrow A. left) from ca. – 40 mV to ca. - 70 mV (maximum) due to a decrease in conductance of the membrane of the outer sensor segment to Na⁺ and Ca²⁺ (\rightarrow p. 348ff.). The potential rises and falls much more sharply in the cones than in the rods. As in other sensory cells, the magnitude of the sensor potential is proportional to the logarithm of stimulus intensity divided by thresholdintensity (Fechner's law). Hyperpolarization decreases glutamate release from the receptor. When this signal is relaved within the retina, a distinction is made between "direct" signal flow for photopic vision and "lateral" signal flow for scotopic vision (see below). Action potentials (APs) can only be generated in ganglion cells (\rightarrow A, right), but stimulus-dependent amplitude changes of the potentials occur in the other retinal neurons ($\rightarrow A$, center). These are conducted electrotonically across the short spaces in the retina (\rightarrow p. 48ff.).

Direct signal flow from cones to bipolar cells is conducted via ON or OFF bipolar cells. Photostimulation leads to depolarization of ON bipolar cells (signal inversion) and activation of their respective ON ganglion cells (\rightarrow A). OFF bipolar cells, on the other hand, are hyperpolarized by photostimulation, which has an inhibitory effect on their OFF ganglion cells. **"Lateral" signal flow** can occur via the following pathway: rod \Rightarrow rod-bipolar cell \Rightarrow ON or OFF bipolar cell \Rightarrow ON or OFF ganglion cell. Both rod-bipolar cells and rod-amacrine cells are depolarized in response to light. Rod-amacrine cells inhibit OFF bipolar cells via a chemical synapse and stimulate ON bipolar cells via an electrical synapse (\rightarrow p. 50).

A light stimulus triggers the firing of an AP in ON ganglion cells (\rightarrow A, right). The AP frequency increases with the sensor potential amplitude. The APs of ON ganglion cells can be measured using microelectrodes. This data can be used to identify the retinal region in which the stimulatory and inhibitory effects on AP frequency originate. This region is called the **receptive field (RF)** of the ganglion cell. Retinal ganglion cells have concentric RFs comprising a central zone and a ringlike peripheral zone distinguishable during light adaptation (\rightarrow B). Photic stimulation of the center increases the

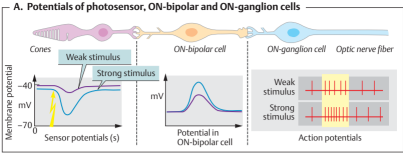
AP frequency of ON ganglion cells $(\rightarrow B1)$. Stimulation of the periphery, on the other hand, leads to a decrease in AP frequency, but excitation occurs when the light source is switched off (\rightarrow **B2**). This type of RF is referred to as an ON field (central field ON). The RF of OFF ganglion cells exhibits the reverse response and is referred to as an OFF field (central field OFF). Horizontal cells are responsible for the functional organization of the RFs $(\rightarrow p. 344)$. They invert the impulses from photosensors in the periphery of the RF and transmit them to the sensors of the center. The opposing central and peripheral responses lead to a stimulus contrast. At a light-dark interface, for example, the dark side appears darker and the light side brighter. If the entire RF is exposed to light, the impulses from the center usually predominate.

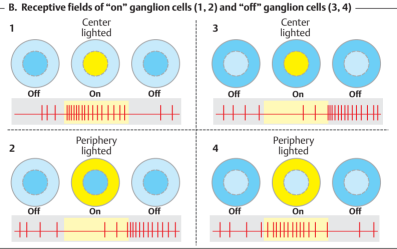
Simultaneous contrast. A solid gray circle appears darker in light surroundings than in dark surroundings (\rightarrow **C**, left). When a subject focuses on a blackand-white grid (\rightarrow **C**, right), the white grid lines appear to be darker at the cross-sections, black grid lines appear lighter because of reduced contrast in these areas. This effect can be attributed to a variable sum of stimuli within the RFs (\rightarrow **C**, center).

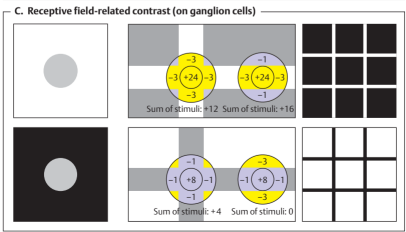
During **dark adaptation**, the center of the RFs increases in size at the expense of the periphery, which ultimately disappears. This leads to an increase in spatial summation (\rightarrow p. 353 C3), but to a simultaneous decrease in stimulus contrast and thus to a lower visual acuity (\rightarrow p. 349 B2).

Color opponency. Red and green light (or blue and yellow light) have opposing effects in the RFs of β ganglion cells (\rightarrow p. 358) and more centrally located cells of the optic tract (\rightarrow p. 357 E). These effects are explained by **Hering's opponent colors theory** and ensure contrast (increase color saturation; \rightarrow p. 356) in color vision. When a subject focuses on a color test pattern (\rightarrow p. 359 C) for about 30 min and then shifts the gaze to a neutral background, the complementary colors will be seen (*color successive contrast*).

RFs of **higher centers** of the optic tract (V1, V2; \rightarrow p. 358) can also be identified, but their characteristics change. Shape (striate or angular), length, axial direction and direction of movement of the photic stimuli play important roles.







Color Vision

White light passing through a prism is split into a color spectrum ranging from red to violet (colors of the rainbow). *Red* is perceived at a wavelength (λ) of 650–700 nm, and violet at around 400–420 nm (\rightarrow **A**). The eye is sensitive to waves in this λ range. Perception of white light does not require the presence of all colors of the visible spectrum. It is sufficient to have the additive effects (mixing) of only two **complementary colors**, e.g., orange (612 nm) and blue light (490 nm).

A **color triangle** $(\rightarrow B)$ or similar test panel can be used to illustrate this effect. The two upper limbs of the triangle show the visible spectrum, and a white dot is located inside the triangle. All straight lines passing through this dot intersect the upper limbs of the triangle at two complementary wavelengths (e.g., 612 and 490 nm). Additive color mixing $(\rightarrow \mathbf{C})$: The color vellow is obtained by mixing roughly equal parts of red and green. Orange is produced by using a higher red fraction, and yellowish green is obtained with a higher green fraction. These colors lie between red and green on the limbs of the color triangle. Similar rules apply when mixing green and violet $(\rightarrow B \text{ and } C)$. The combination of red with violet yields a shade of purple not contained in the spectrum (\rightarrow B). This means that all colors, including white, can be produced by varying the proportions of three colors-e.g. red (700 nm), green (546 nm) and blue (435 nm) because every possible pair of complementary colors can be obtained by mixing these three colors of the spectrum.

Subtractive color mixing is based on the opposite principle. This technique is applied when color paints and camera filters are used. Yellow paints or filters absorb ("subtract") the blue fraction of white light, leaving the complementary color yellow.

Light absorption. Photosensors must be able to absorb light to be photosensitive. **Rods** (\rightarrow p.348) contain **rhodopsin**, which is responsible for (achromatic) night vision. Rhodopsin absorbs light at wavelengths of ca. 400–600 nm; the maximum absorption value (λ_{max}) is 500 nm (\rightarrow E1). Relatively speaking, greenish blue light therefore appears brightest and red appears darkest at night. Wearing red glasses in daylight therefore leaves the rods adapted for darkness. Three types of color-sensitive **cones** are responsible for (chromatic) day vision (\rightarrow E1): (1) *S cones*, which absorb short-wave (S) blue-violet light (λ_{max} =

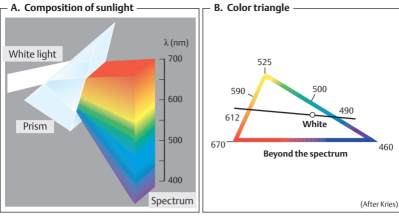
420 nm); (2) *M cones*, which absorb mediumwave (M) blue-green to yellow light (λ_{max} = 535 nm), and (3) *L cones*, which absorb longwave (L) yellow to red light (λ_{max} = 565 nm). (The physiological sensitivity curves shown in **E1** make allowances for light absorbed by the lens.) *Ultraviolet rays* (λ_{max} < 400 nm) and *infrared rays* (λ_{max} > 700 nm) are not visible.

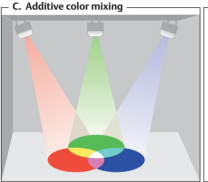
Sensory information relayed by the three types of cones (peripheral application of the *trichromatic theory of color vision*) and transduction of these visual impulses to *brightness* and *opponent color channels* (\rightarrow **E2** and p. 354) in the retina and lateral geniculate body (LGB) enables the visual cortex (\rightarrow p. 358) to recognize different **types of colors**. The human eye can distinguish 200 shades of color and different degrees of color saturation. The absolute *differential threshold for color vision* is 1–2 nm (\rightarrow **D**, "normal").

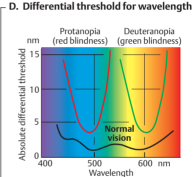
Color perception is more complex. White paper, for example, will look white in white light (sunlight), yellow light (light bulb) and red light. We also do not perceive the different shades of color in a house that is partially illuminated by sunlight and partially in the shade. This **color constancy** is the result of retinal and central processing of the retinal signal.

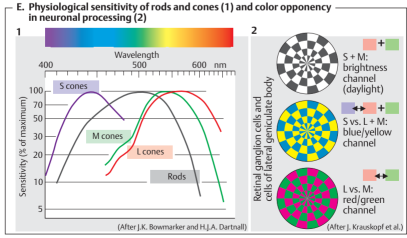
There is a similar **constancy of size and shape**: Although someone standing 200 meters away makes a much smaller image on the retina that at 2 meters' distance, we still recognize him or her as a person of normal height, and although a square table may appear rhomboid in shape when viewed from the side, we can still tell that it is square.

Color blindness occurs in 9% of all men and in 0.5% of women. The ability to distinguish certain colors is impaired or absent in these individuals, i.e., they have a high differential threshold for color $(\rightarrow \mathbf{D})$. Various types of color blindness are distinguished: protanopia (red blindness), deuteranopia (green blindness), and tritanopia (blue-violet blindness). Protanomaly, deuteranomaly and tritanomaly are characterized by decreased sensitivity of the cones to colored, green and blue, respectively. Color vision is tested using color perception charts or an anomaloscope. With the latter, the subject has to mix two color beams (e.g., red and green) with adjustable intensities until their additive mixture matches a specific shade of color (e.g. yellow, \rightarrow **C**) presented for comparison. A protanomal subject needs a too high red intensity, a deuteranomal person a too high green intensity. Protanopes perceive all colors with wavelengths over approx. 520 nm as yellow.









Visual Field, Visual Pathway, Central Processing of Visual Stimuli

The **visual field** $(\rightarrow A)$ is the area visualized by the *immobile* eye with the head fixed.

The visual field is examined by **perimetry**. The subject's eve is positioned in the center of the perimeter. which is a hollow hemispherical instrument. The subject must then report when laterally flashed points of light appear in or disappear from the visual field. An area of lost vision within the visual field is a scotoma. Lesions of the retina, or of the central visual pathway can cause scotoma. The **blind spot** $(\rightarrow A)$ is a normal scotoma occurring at 15 degrees temporal and is caused by nasal interruption of the retina by the optic disk (\rightarrow p. 349 B). In binocular vision (\rightarrow p. 361 A). the blind spot of one eye is compensated for by the other. The visual field for color stimuli is smaller than that for light-dark stimuli. If, for example, a red object is slowly moved into the visual field, the movement will be identified more quickly than the color of the object.

The retina contains more than 10^8 photosensors connected by retinal neurons (\rightarrow p. 354) to ca. 10^6 retinal ganglion cells. Their axons form the **optic nerve**. The convergence of so many sensors on only a few neurons is particularly evident in the retinal periphery (1000:1). In the fovea centralis, however, only one or a few cones are connected to one neuron. Due to the low convergence of impulses from the fovea, there is a high level of visual acuity with a low level of light sensitivity, whereas the high convergence of signals from the periphery produces the reverse effect (cf. spatial summation; \rightarrow p. 353 C3).

Ganglion cells. Three types of ganglion cells can be found in the retina: (1) 10% are \mathbf{M} (or α or Y) cells of the magnocellular system; their fast-conducting axons emit short phasic impulses in response to light and are very sensitive to movement; (2) 80% are the \mathbf{P} (or β or X) cells of the parvicellular system; their thin axons have small receptive fields (high spatial resolution), persistently react to constant light (tonic reaction), and therefore permit pattern and color analysis. Both types have equal densities of ON and OFF cells (\rightarrow p. 354). (3) 10% are γ (or W) cells of the coniocellular system; their very thin axons project to the mesencephalon and regulate pupil diameter (see below) and reflex saccades (\rightarrow pp. 348, 360).

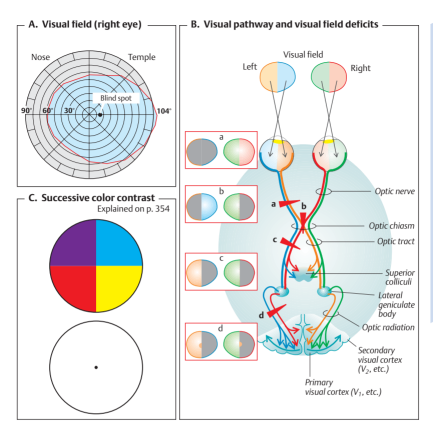
Objects located in the nasal half of the visual field of each eye (\rightarrow **B**, blue and green) are imaged in the temporal half of each retina and vice versa. Along the **visual pathway**, fibers of the **optic nerve** from the temporal half of each retina remain on the same side (\rightarrow **B**, blue and green), whereas the fibers from the nasal half of each retina decussate at the **optic chiasm** (\rightarrow **B**, orange and red). Fibers from the fovea centralis are present on both sides.

Lesions of the left optic *nerve* for instance cause deficits in the entire left visual field $(\rightarrow \mathbf{B}, \mathbf{a})$, whereas lesions of the left optic *tract* produce deficits in the right halves of both visual fields $(\rightarrow \mathbf{B}, \mathbf{c})$. Damage to the median optic *chiasm* results in bilateral temporal deficits, i.e., *bitemporal hemianopia* $(\rightarrow \mathbf{B}, \mathbf{b})$.

Fibers of the optic tract extend to the lateral **geniculate body** $(\rightarrow B)$ of the thalamus, the six layers of which are organized in a retinotopic manner. Axons of the ipsilateral eye terminate on layers 2, 3 and 5, while those of the contralateral eye terminate on layers 1, 4 and 6. The M cell axons communicate with cells of magnocellular layers 1 and 2, which serve as a relay station for motion-related stimuli that are rapidly conducted to the motor cortex. The P cell axons project to the parvocellular layers 3-6, the main function of which is to process colors and shapes. The neurons of all layers then project further through the optic radiation (arranged also retinotopically) to the primary visual cortex (V₁) and, after decussating, to further areas of the visual cortex (V2-5) including pathways to the parietal and temporal cortex. Magnocellular input reaches the parietal cortex also via the superior colliculi (see below) and the pulvinar.

The **primary visual cortex** (V_1) is divided depth-wise (x-axis) into six retinotopic layers numbered I to VI (\rightarrow p. 333 A). Cells of the primary visual cortex are arranged as three-dimensional modular **hyper-columns** ($3 \times 1 \times 1$ mm) representing modules for analysis of all sensory information from corresponding areas of both retinas (\rightarrow p. 360). Adjacent hypercolumns represent neighboring retinal regions. Hypercolumns contain *ocular dominance columns* (y-axis), *orientation columns* (z-axis), and *cylinders* (x-axis). The dominance columns receive alternating input from the right and left eye, orientation columns process direction of stimulus movement and cylinders receive information of colors.

359



Color, high-resolution stationary shapes, movement, and stereoscopic depth are processed in some subcortical visual pathways, and from V_1 onward in separate information channels. These individual aspects must be integrated to achieve visual perception. In diurnally active primates like humans, over half of the cortex is involved in processing visual information. On a simplified scale, the parietal cortex analyzes the "where" and involves motor systems, and the temporal cortex takes care of the "what" of visual input comparing it with memory.

Axons of the optic tract (especially those of M and γ cells) also project to **subcortical regions** of the brain such as the *pretectal region*, which regulates the diameter of the pupils (see below); the *superior colliculi* (\rightarrow **B**), which are involved in oculomotor function (\rightarrow p. 360);

the *hypothalamus*, which is responsible for circadian rhythms (\rightarrow p. 334).

The **pupillary reflex** is induced by sudden exposure of the retina to light (\rightarrow p. 350). The signal is relayed to the pretectal region; from here, a parasympathetic signal flows via the Edinger–Westphal nucleus, the ciliary ganglion and the oculomotor nerve, and induces narrowing of the pupils (*miosis*) within less than 1 s. Since both pupils respond simultaneously even if the light stimulus is unilateral, this is called a *consensual light response*. Meiosis also occurs when the eyes adjust for near vision (*near-vision response* \rightarrow p. 360).

The **corneal reflex** protects the eye. An object touching the cornea (afferent: trigeminal nerve) or approaching the eye (afferent: optic nerve) results in reflex closure of the eyelids.

Eye Movements, Stereoscopic Vision, Depth Perception

Conjugated movement of the eyes occurs when the external eye muscles move the eyes in the same direction (e.g., from left to right), whereas vergence movement is characterized by opposing (divergent or convergent) eye movement. The axes of the eyes are parallel when gazing into the distance. Fixation of the gaze on a nearby object results in convergence of the visual axes. In addition, the pupil contracts (to increase the depth of focus) and accommodation of the lens occurs (\rightarrow p. 346). The three reactions are called **near-vision response** or convergence response.

Strabismus. A greater power of accommodation for near vision is required in hyperopia than in normal vision. Since accommodation is always linked with a convergence impulse, hyperopia is often associated with squinting. If the visual axes wander too far apart, vision in one eye will be suppressed to avoid double vision (diplopia). This type of visual impairment, called *strabismic amblyopia*, can be either temporary or chronic.

Saccades. When scanning the visual field, the eyes make jerky movements when changing the point of fixation, e.g., when scanning a line of print. These quick movements that last 10–80 ms are called *saccades*. Displacement of the image is centrally suppressed during the eye due to *saccadic suppression*. A person looking at both of his or her eyes alternately in a mirror cannot perceive the movement of his or her own eyes, but an independent observer can. The small, darting saccades function to keep an object in focus.

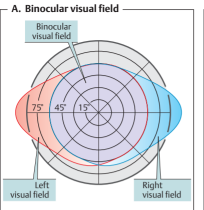
Objects entering the field of vision are reflexively imaged in the fovea centralis (\rightarrow p. 348). Slow **pursuit movements** of the eyes function to maintain the gaze on moving objects. **Nystagmus** is characterized by a combination of these slow and rapid (saccade-like) opposing eye movements. The direction of nystagmus (right or left) is classified according to the type of rapid phase, e.g., secondary nystagmus (\rightarrow p. 342). *Optokinetic nystagmus* occurs when viewing an object passing across the field of vision, e.g., when looking at a tree from inside a moving train. Once the eyes have returned to the normal position (return sac-

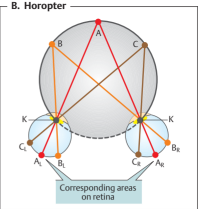
cade), a new object can be brought into focus. Damage to the cerebellum or organ of balance $(\rightarrow p.342)$ can result in *pathological nystagmus*.

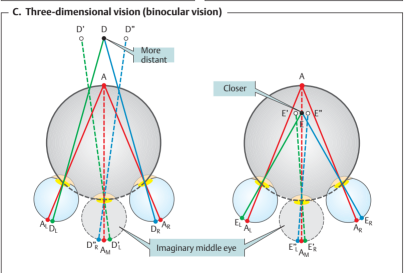
The brain stem is the main center responsible for **programming** of eye movements. Rapid horizontal (conjugated) movements such as saccades and rapid nystagmus movement are programmed in the pons, whereas vertical and torsion movements are programmed in the mesencephalon. The cerebellum provides the necessary fine tuning (\rightarrow p. 326). Neurons in the region of the Edinger–Westphal nucleus are responsible for vergence movements.

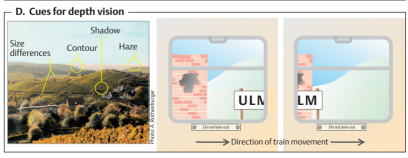
In near vision, depth vision and three-dimensional vision are primarily achieved through the coordinated efforts of both eves and are therefore limited to the binocular field of vision $(\rightarrow A)$. If both eyes focus on point A $(\rightarrow B)$, an image of the fixation point is projected on both foveae (A_I, A_R), i.e., on the corresponding areas of the retina. The same applies for points B and C $(\rightarrow B)$ since they both lie on a circle that intersects fixation point A and nodal points N (\rightarrow p. 347 B) of the two eyes (Vieth-Müller horopter). If there were an imaginary middle eye in which the two retinal regions (in the cortex) precisely overlapped, the retinal sites would correspond to a central point Ac \triangle $A_L + A_R (\rightarrow C)$. Assuming there is a point D outside the horopter $(\rightarrow C$, left), the middle eye would see a double image (D', D") instead of point D, where D' is from the left eye (D_L) . If D and A are not too far apart, central processing of the double image creates the perception that D is located behind D, i.e., depth perception occurs. A similar effect occurs when a point E $(\rightarrow C$, right) is closer than A; in this case, the E' image will arise in the right eye (E'_R) and E will be perceived as being closer.

Depth perception from a distance. When viewing objects from great distances or with only one eye, contour overlap, haze, shadows, size differences, etc. are cues for depth perception $(\rightarrow \mathbf{D})$. A nearby object moves across the field of vision more quickly than a distant object, e.g., in the case of the sign compared to the wall in plate \mathbf{D}). In addition, the moon appears to migrate with the moving car, while the mountains disappear from sight.









Sound waves are the adequate stimulus for the organ of hearing. They arise from a sound source such as a gong $(\rightarrow A1)$ and are conducted in gases, liquids, and solids. The air is the main carrier of sound.

The air pressure rises and falls rhythmically at the sound source. These pressure waves (sound waves) travel at a characteristic sound velocity (c) in different materials, e.g., at 332 m/s in air of 0 °C. A graphic recording of sound waves $(\rightarrow A1)$ will produce waveform curves. The wavelength (λ) is the distance between the top of one wave and the identical phase of the succeeding one, and the maximum deviation of pressure from baseline is the amplitude (a) (\rightarrow A1). Enlargement (reduction) of wavelength will lower (raise) the tone, whereas a fall (rise) in amplitude will produce a quieter (louder) tone (→A1). The **pitch** of a tone is defined by its frequency (f), i.e., the number of sound pressure oscillations per unit time. Frequency is measured in **hertz** (Hz = s^{-1}). Frequency, wavelength and the sound velocity are related:

$$f(Hz) \cdot \lambda(m) = c(m \cdot s^{-1}).$$
 [12.1]

A **pure tone** has a simple sinus waveform. The tones emanating from most sound sources (e.g., musical instrument, voice) are mixtures of different frequencies and amplitudes that result in complex periodic vibrations referred to as **sound** (\rightarrow **A2**). The fundamental (lowest) tone in the complex determines the *pitch* of the sound, and the higher ones determine its *timbre* (overtones). An a¹ (440 Hz) sung by a tenor or played on a harp has a different sound than one produced on an organ or piano. The overlap of two very similar tones produces a distinct effect characterized by a *beat tone* of a much lower frequency (\rightarrow **A3**, blue/red).

Audibility limits. Healthy young persons can hear sounds ranging in frequency from 16 to 20000 Hz. The upper limit of audibility can drop to 5000 Hz due to aging (presbycusis). At 1000 Hz, the absolute auditory threshold or lowest sound pressure perceived as sound is $3 \cdot 10^{-5}$ Pa. The threshold of sound is frequency-dependent (\rightarrow **B**, green curve). The threshold of hearing for a tone rises tremendously when other tones are heard simultaneously. This phenomenon called masking is the reason why it is so difficult to carry on a conversation against loud background noise. The ear is overwhelmed by sound pressures over 60 Pa, which corresponds to $2 \cdot 10^6$ times the sound pressure of the limit of audibility at

1000 Hz. Sounds above this level induce the sensation of pain (\rightarrow **B**, red curve).

For practical reasons, the *decibel* (*dB*) is used as a logarithmic measure of the **sound pressure level (SPL)**. Given an arbitrary reference sound pressure of $p_0 = 2 \cdot 10^{-5}$ Pa, the sound pressure level (SPL) can be calculated as follows:

$$SPL(dB) = 20 \cdot log(p_x/p_o)$$
 [12.2]

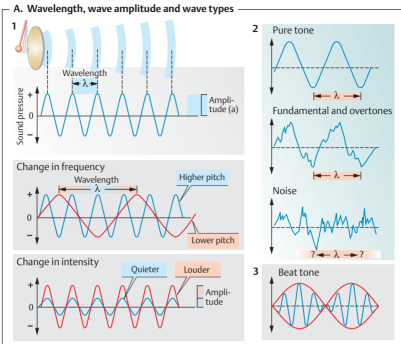
where p_x is the actual sound pressure. A tenfold increase in the sound pressure therefore means that the SPL rises by 20 dB.

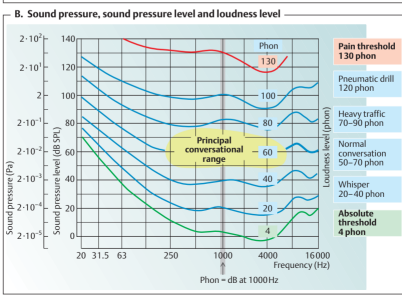
The **sound intensity (I)** is the amount of sound energy passing through a given unit of area per unit of time $(W \cdot m^2)$. The sound intensity is proportional to the *square* of p_x . Therefore, dB values cannot be calculated on a simple linear basis. If, for example, two loudspeakers produce 70 dB each $(p_x = 6.3 \cdot 10^{-2} \text{ Pa})$, they do *not* produce 140 dB together, but a mere 73 dB because p_x only increases by a factor of $\sqrt{2}$ when the intensity level doubles. Thus, $\sqrt{2} \cdot 6.3 \cdot 10^{-2}$ Pa has to be inserted for p_x into Eq. 12.2.

Sound waves with different frequencies but equal sound pressures are not subjectively perceived as equally loud. A 63 Hz tone is only perceived to be as loud as a 20 dB/1000 Hz reference tone if the sound pressure of the 63 Hz tone is 30-fold higher (+29 dB). In this case, the sound pressure level of the reference tone (20 dB/1000 Hz) gives the loudness level of the 63 Hz tone in **phon** (20 phon) as at a frequency of 1000 Hz, the phon scale is numerically equals the dB SPL scale (\rightarrow **B**). Equal loudness contours or isophones can be obtained by plotting the subjective values of equal loudness for test frequencies over the whole audible range $(\rightarrow B$, blue curves). The absolute auditory threshold is also an isophone (4 phons; \rightarrow **B**, green curve). Human hearing is most sensitive in the 2000–5000 Hz range (\rightarrow **B**).

Note: Another unit is used to describe how a tone of constant frequency is subjectively perceived as louder or less loud. *Sone* is the unit of this type of loudness, where 1 sone = 40 phons at 1000 Hz. 2 sones equal twice the reference loudness, and 0.5 sone is $^{1}/_{2}$ the reference loudness.

The **auditory** area in diagram **B** is limited by the highest and lowest audible frequencies on the one side, and by isophones of the thresholds of hearing and pain on the other. The green area in plate **B** represents the range of frequencies and intensities required for comprehension of ordinary speech $(\rightarrow B)$.





Sound waves are transmitted to the organ of hearing via the *external ear* and the *auditory canal*, which terminates at the *tympanic membrane* or *eardrum*. The sound waves are conducted through the air (**air conduction**) and set the eardrum in vibration. These are transmitted via the *auditory ossicles* of the tympanic cavity (*middle ear*) to the membrane of the *oval window* (\rightarrow A 1,2), where the *internal* or *inner ear* (labyrinth) begins.

In the **middle ear**, the *malleus*, *incus* and *stapes* conduct the vibrations of the tympanic membrane to the oval window. Their job is to conduct the sound from the low wave resistance/impedance in air to the high resistance in fluid with as little loss of energy as possible. This **impedance transformation** occurs at f < 2400 Hz and is based on a 22-fold pressure amplification (tympanic membrane area/oval window area is 17:1, and leverage arm action of the auditory ossicles amplifies force by a factor of 1.3). Impairment of impedance transforming capacity due, e.g., to destruction of the ossicles, causes roughly 20 dB of hearing loss (*conduction deafness*).

Muscles of the middle ear. The middle ear contains two small muscles—the *tensor tympani* (insertion: manubrium of malleus) and the *stapedius* (insertion: stapes)—that can slightly attenuate low-frequency sound. The main functions of the inner ear muscles are to maintain a constant sound intensity level, protect the ear from loud sounds, and to reduce distracting noises produced by the listener.

Bone conduction. Sound sets the skull in vibration, and these bone-borne vibrations are conducted directly to the cochlea. Bone conduction is fairly insignificant for physiological function, but is useful for testing the hearing. In Weber's test, a vibrating tuning fork (a1) is placed in the middle of the head. A person with normal hearing can determine the location of the tuning fork because of the symmetrical conduction of sound waves. A patient with unilateral conduction deafness will perceive the sound as coming from the affected side (lateralization) because of the lack of masking of environmental noises in that ear (bone conduction). A person with sensorineural deafness, on the other hand, will perceive the sound as coming from the healthy ear because of sound attenuation in the affected internal ear. In Rinne's test. the handle of a tuning fork is placed on one mastoid process (bony process behind the ear) of the patient (bone conduction). If the tone is no longer heard, the

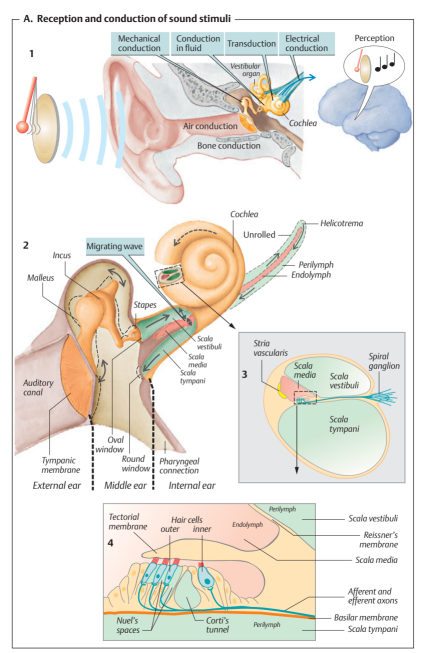
tines of the tuning fork are placed in front of the ear (air conduction). Individuals with normal hearing or sensorineural deafness can hear the turning fork in the latter position anew (positive test result), whereas those with conduction deafness cannot (test negative).

The **internal ear** consists of the *equilibrium* organ $(\rightarrow p. 342)$ and the **cochlea**, a spiraling bony tube that is 3-4 cm in length. Inside the cochlea is an endolymph-filled duct called the scala media (cochlear duct): the ductus reuniens connects the base of the cochlear duct to the endolymph-filled part of the equilibrium organ. The scala media is accompanied on either side by two perilymph-filled cavities: the scala vestibuli and scala tympani. These cavities merge at the apex of the cochlea to form the helicotrema. The scala vestibuli arises from the oval window, and the scala tympani terminates on the membrane of the round window (\rightarrow **A2**). The composition of perilymph is similar to that of plasma water (→ p. 93 C), and the composition of endolymph is similar to that of the cytosol (see below). Perilymph circulates in Corti's tunnel and Nuel's spaces (\rightarrow **A4**).

Organ of Corti. The (secondary) sensory cells of the hearing organ consist of approximately $10\,000-12\,000$ external **hair cells (HCs)** and 3500 internal hair cells that sit upon the basilar membrane (\rightarrow **A4**). Their structure is very similar to that of the vestibular organ (\rightarrow p. 342) with the main difference being that the kinocilia are absent or rudimentary.

There are three rows of slender, cylindrical outer hair cells, each of which contains approximately 100 cilia (actually microvilli) which touch the tectorial membrane. The bases of the hair cells are firmly attached to the basilar membrane by supporting cells, and their cell bodies float in perilymph of Nuel's spaces (\rightarrow A4). The outer hair cells are principally innervated by efferent, mostly cholinergic neurons from the spiral ganglion (N_M-cholinoceptors; → p. 82). The inner hair cells are pear-shaped and completely surrounded by supporting cells. Their cilia project freely into the endolymph. The inner hair cells are arranged in a single row and synapse with over 90% of the afferent fibers of the spiral ganglion. Efferent axons from the nucleus olivaris superior lateralis synapse with the afferent endings.

Sound conduction in the inner ear. The stapes moves against the membrane of the oval window membrane, causing it to vibrate. These are transmitted via the perilymph to the membrane of the round window (\rightarrow A2). The walls of the endolymph-filled cochlear duct, i.e. Reissner's membrane and the basilar mem-



brane $(\rightarrow D1)$ give against the pressure wave (migrating wave, \rightarrow B and C). It can therefore take a "short cut" to reach the round window without crossing the helicotrema. Since the cochlear duct is deformed in waves. Reissner's membrane and the basilar membrane vibrate alternately towards the scala vestibuli and scala tympani (\rightarrow **D1,2**). The velocity and wavelength of the migrating wave that started at the oval window decrease continuously $(\rightarrow B)$, while their amplitude increases to a maximum and then quickly subsides (\rightarrow B, envelope curve). (The wave velocity is not equal to the velocity of sound, but is much slower.) The site of the maximum excursion of the cochlear duct is characteristic of the wavelength of the stimulating sound. The higher the frequency of the sound, the closer the site is to the stapes $(\rightarrow \mathbb{C})$.

Outer hair cells. Vibration of the cochlear duct causes a discrete shearing (of roughly 0.3 nm) of the tectorial membrane against the basilar membrane, causing bending of the cilia of the outer hair cells (\rightarrow D3). This exerts also a shearing force between the rows of cilia of the individual external hair cell. Probably via the "tip links" (\rightarrow p. 342), cation channels in the ciliary membranes open (mechanosensitive transduction channels), allowing cations (K+, Na⁺, Ca²⁺) to enter and depolarize the outer hair cells. This causes the outer hair cells to shorten in sync with stimulation (\rightarrow **D3**). The successive shearing force on the cilia bends them in the opposite direction. This leads to hyperpolarization (opening of K⁺ channels) and extension of the outer hair cells.

The mechanism for this extremely fast **electromotility** (up to $20 \, \text{kHz}$ or $2 \cdot 10^4$ times per second) is unclear, but it seems to be related to the high turgor of outer hair cells ($128 \, \text{mmHg}$) and the unusual structure of their cell walls.

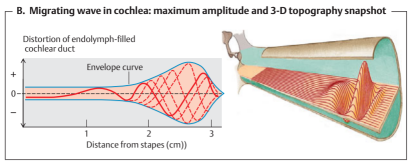
These outer hair cell electromotility contributes to the **cochlear amplification** (ca. 100-fold or 40 dB amplification), which occurs before sound waves reach the actual sound sensors, i.e. inner hair cells. This explains the very low threshold within the very narrow location (0.5 nm) and thus within a very small frequency range. The electromotility causes endolymph waves in the subtectorial space which exert *shearing forces on the* **inner hair**

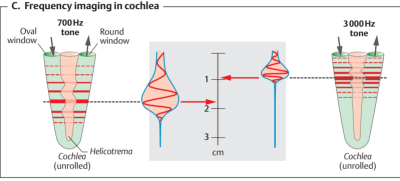
cell *cilia* at the site of maximum reaction to the sound frequency (\rightarrow **D4**), resulting in opening of transduction channels and depolarization of the cells (**sensor potential**). This leads to *transmitter* release (glutamate coupling to AMPA receptors; \rightarrow p. 55 F) by internal hair cells and the subsequent conduction of impulses to the CNS.

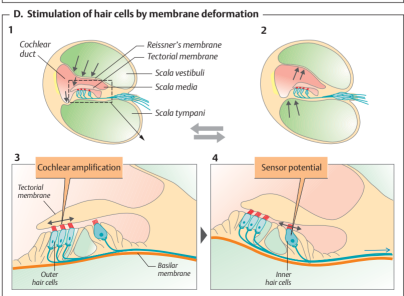
Vibrations in the internal ear set off an outward emission of sound. These **evoked otoacoustic emissions** can be measured by placing a microphone in front of the tympanic membrane, e.g., to test internal ear function in infants and other individuals incapable of reporting their hearing sensations.

Inner ear potentials (\rightarrow p. 369 C). On the cilia side, the hair cells border with the endolymphfilled space, which has a potential difference (endocochlear potential) of ca. +80 to +110 mV relative to perilymph (\rightarrow p. 369 C). This potential difference is maintained by an active transport mechanism in the stria vascularis. Since the cell potential of outer (inner) hair cells is -70 mV (-40 mV), a potential difference of roughly 150-180 mV (120-150 mV) prevails across the cell membrane occupied by cilia (cell interior negative). Since the K⁺ conc. in the endolymph and hair cells are roughly equal (≈ 140 mmol/L), the prevailing K⁺ equilibrium potential is ca. $0 \, \text{mV} \ (\rightarrow \text{p.} 32)$. These high potentials provide the driving forces for the influx not only of Ca2+ and Na+, but also of K+, prerequisites for provoking the sensor potential.

Hearing tests are performed using an **audiometer**. The patient is presented sounds of various frequencies and routes of conduction (bone, air). The sound pressure is initially set at a level under the threshold of hearing and is raised in increments until the patient is able to hear the presented sound (threshold audiogram). If the patient is unable to hear the sounds at normal levels, he or she has an hearing loss, which is quantitated in decibels (dB). In audiometry, all frequencies at the normal threshold of hearing are assigned the value of 0 dB (unlike the diagram on p. 363 B, green curve). Hearing loss can be caused by presbycusis (→ p. 362), middle ear infection (impaired air conduction), and damage to the internal ear (impaired air and bone conduction) caused, for example, by prolonged exposure to excessive sound pressure (> 90 dB, e.g. disco music, pneumatic drill etc.).







Central Processing of Acoustic Information

Various qualities of sound must be coded for signal transmission in the acoustic pathway. These include the frequency, intensity and direction of sound waves as well as the distance of the sound source from the listener.

Frequency imaging. Tones of various frequencies are "imaged" along the cochlea. conducted in separate fibers of the auditory pathway and centrally identified. Assuming that a tone of 1000 Hz can just be distinguished from one of 1003 Hz (resembling true conditions), the frequency difference of 3 Hz corresponds to a relative frequency differential threshold of 0.003 (\rightarrow p. 352). This fine differential capacity is mainly due to frequency imaging in the cochlea, amplification by its outer hair cells (\rightarrow p. 366), and neuronal contrast along the auditory pathway (\rightarrow p. 313 D). This fine tuning ensures that a certain frequency has a particularly low threshold at its "imaging" site. Adjacent fibers are not recruited until higher sound pressures are encountered.

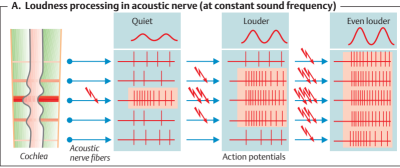
Intensity. Higher intensity levels result in higher action potential frequencies in afferent nerve fibers and recruitment of neighboring nerve fibers (\rightarrow **A**). The relative *intensity differential threshold* is 0.1 (\rightarrow p. 352), which is very crude compared to the frequency differential threshold. Hence, differences in loudness of sound are not perceived by the human ear until the intensity level changes by a factor of over 1.1, that is, until the sound pressure changes by a factor of over $\sqrt{1,1} = 1,05$.

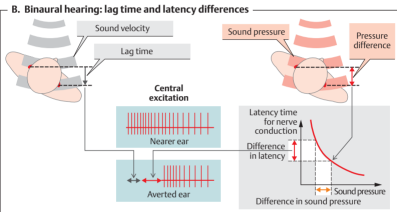
Direction. *Binaural hearing* is needed to identify the direction of sound waves and is based on the following two effects. (1) Sound waves that strike the ear obliquely reach the averted ear later than the other, resulting in a *lag time*. The change in direction that a normal human subject can just barely detect (*direction threshold*) is roughly 3 degrees. This angle delays the arrival of the sound waves in the averted ear by about $3 \cdot 10^{-5} \text{ s}$ (\rightarrow **B**, left). (2) Sound reaching the averted ear is also perceived as being *quieter*; differences as small as 1 dB can be distinguished. A lower sound pressure results in delayed firing of actions potentials, i.e., in increased *latency* (\rightarrow **B**, right). Thus,

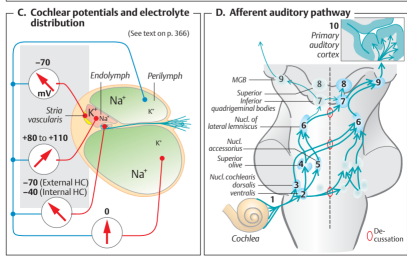
the impulses from the averted ear reach the CNS later (nucleus accessorius, \rightarrow **D5**). Effects (1) and (2) are additive effects (\rightarrow **B**). The external ear helps to decide whether the sound is coming from front or back, above or below. Binaural hearing also helps to distinguish a certain voice against high background noise, e.g., at a party. Visibility of the speaker's mouth also facilitates comprehension.

Distance to the sound source can be determined because high frequencies are attenuated more strongly than low frequencies during sound wave conduction. The longer the sound wave travels, the lower the proportion of high frequencies when it reaches the listener. This helps, for instance, to determine whether a thunderstorm is nearby or far away.

Auditory pathway $(\rightarrow D)$. The auditory nerve fibers with somata positioned in the spiral ganglion of the cochlea project from the cochlea $(\rightarrow D1)$ to the anterolateral $(\rightarrow D2)$. posteroventral and dorsal cochlear nuclei (→ D3). Afferents in these three nuclei exhibit tonotopicity, i.e., they are arranged according to tone frequency at different levels of complexity. In these areas, lateral inhibition $(\rightarrow p.313 D)$ enhances contrast, i.e., suppresses noise. Binaural comparison of intensity and transit time of sound waves (direction of sound) takes place at the next-higher station of the auditory pathway, i.e. in the superior olive (\rightarrow **D4**) and accessory nucleus (\rightarrow **D5**). The next stations are in the nucleus of lateral lemniscus (\rightarrow **D6**) and, after most fibers cross over to the opposite side, the inferior quadrigeminal bodies (\rightarrow D7). They synapse with numerous afferents and serve as a reflex station (e.g., muscles of the middle ear; \rightarrow p. 366). Here, sensory information from the cochlear nuclei is compared with spatial information from the superior olive. Via connections to the superior quadrigeminal bodies (\rightarrow **D8**), they also ensure coordination of the auditory and visual space. By way of the thalamus (medial geniculate body, MGB; \rightarrow **D9**), the afferents ultimately reach the *primary* auditory cortex (→ D10) and the surrounding secondary auditory areas $(\rightarrow p.311 E$, areas 41 and 22). Analysis of complex sounds, short-term memory for comparison of tones, and tasks required for "eavesdropping" are some of their functions.







Voice and Speech

The human voice primarily functions as a means of communication, the performance of which is based on the human capacity of hearing (\rightarrow p. 363 B). As in wind instruments, the body contains a wind space (trachea, bronchi. etc.). Air is driven through the space between the vocal cords (rima glottidis) into the air space (passages above the glottis), which sets the vocal cords into vibration. The air space consists of the throat and oronasal cavities $(\rightarrow A)$. The range of the human voice is so immense because of the large variety of muscles that function to modulate the intensity of the airstream (loudness), tension of the vocal cords, shape/width of the vocal cords (fundamental tone) and size/shape of the air space (timbre, formants) of each individual.

Joints and muscles of the **larynx** function to adjust the vocal cords and rima glottidis. A stream of air opens and closes the rima glottidis and sets off the rolling movement of the vocal cords (\rightarrow **B**). When a deep tone is produced, the fissure of the glottis remains closed longer than it opens (ratio of 5:1 at 100 Hz). This ratio drops to 1.4:1 in higher tones (400 Hz). The rima glottis remains open when whispering or singing falsetto (\rightarrow **C**, blue).

Motor signals originate in the motosensory cortex (\rightarrow p. 325 C/B, tongue/throat) and are conducted via the vagus nerve to the larynx. Sensory impulses responsible for voice production and the cough reflex are also conducted by the vagus nerve. Sensory fibers from the mucosa and muscle spindles of the larynx (\rightarrow p. 316) continuously transmit information on the position and tension of the vocal cords to the CNS. These reflexes and the close connection of the auditory pathway with bulbar and cortical motor speech centers are important for **fine adjustment of the voice**.

Vowels (\rightarrow **D**). Although their fundamental frequencies are similar (100–130 Hz), spoken vowels can be distinguished by their characteristic overtones (formants). Different formants are produced by modifying the shape of oral tract, i.e., mouth and lips (\rightarrow **D**). The three primary vowels [a:], [i:], [u:] make up the vowel triangle; [\overline{\text{ce}}:], [\overline{\text{ce}}:], [\overline{\text{ge}}:], [\overline{\text{ge}:}], [\overline{\text{ge}}:], [\overline{\text{ge}:}], [\overl

The phonetic notation used here is that of the *International Phonetic Society*. The symbols mentioned

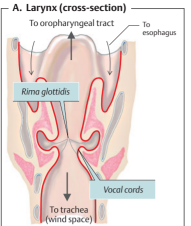
here are as follows: [a:] as in glass; [i:] as in beat; [u:] as in food; [œ:] as in French peur; [o:] as in bought; [ø:] as in French peu or in German hören; [y:] as in French menu or in German trüb; [æ:] as in bad; [ɛ:] as in head.

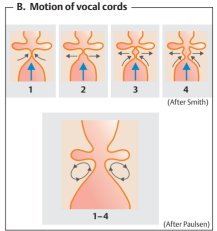
Consonants are described according to their site of articulation as *labial* (lips, teeth), e.g. P/B/W/F/M; *dental* (teeth, tongue), e.g. D/T/S/M; *lingual* (tongue, front of soft palate), e.g. L; *guttural* (back of tongue and soft palate), e.g. C/K. Consonants can be also defined according to their manner of articulation, e.g., *plosives* or *stop consonants* (P|B/T|D/K/G), *fricatives* (F)V/W/S/Ch) and *vibratives* (R).

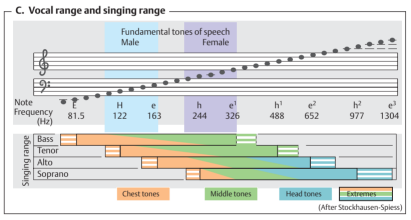
The **frequency range** of the voice, including formants, is roughly 40–2000 Hz. Sibilants like |s| and |z| have higher-frequency fractions. Individuals suffering from presbyacusis or other forms of sensorineural hearing loss are often unable to hear sibilants, making it impossible for them to distinguish between words like "bad" and "bass." The **tonal range** (fundamental tone, \rightarrow **C**) of the *spoken voice* is roughly one octave; that of the *singing voice* is roughly two octaves in untrained singers, and over three octaves in trained singers.

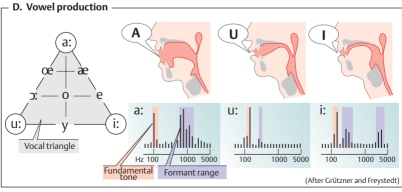
Language (see also p. 336). The main components of verbal communication are (a) auditory signal processing (\rightarrow p. 368), (b) central speech production and (c) execution of motor speech function. The centers for speech comprehension are mainly located in the posterior part of area 22, i.e., Wernicke's area $(\rightarrow p.311 E)$. Lesions of it result in a loss of language comprehension capacity (sensory aphasia). The patient will speak fluently yet often incomprehensibly, but does not notice it because of his/her disturbed comprehension capacity. The patient is also unable to understand complicated sentences or written words. The centers for speech production are mainly located in areas 44 and 45, i.e., Broca's area $(\rightarrow p.311 E)$. It controlls the primary speech centers of the sensorimotor cortex.

Lesions of this and other cortical centers (e.g., gyrus angularis) result in disorders of speech production (motor aphasia). The typical patient is either completely unable to speak or can only express himself in telegraphic style. Another form of aphasia is characterized by the forgetfulness of words (anomic or amnestic aphasia). Lesions of executive motor centers (corticobulbar tracts, cerebellum) cause various speech disorders. Auditory feedback is extremely important for speech. When a person goes deaf, speech deteriorates to an appreciable extent. Children born deaf do not learn to speak.









Dimensions and Units

Physiology is the science of life processes and bodily functions. Since they are largely based on physical and chemical laws, the investigation, understanding, assessment, and manipulation of these functions is inseparably linked to the measurement of physical, chemical, and other parameters, such as blood pressure, hearing capacity, blood pH, and cardiac output. The units for measurement of these parameters are listed in this section. We have given preference to the international system of SI units (Système International d'Unités) for uniformity and ease of calculation. Non-SI units will be marked with an asterisk. Conversion factors for older units are also listed. Complicated or less common physiological units (e.g., wall tension, flow resistance, compliance) are generally explained in the book as they appear. However, some especially important terms that are often (not always correctly) used in physiology will be explained in the Appendix, e.g., concentration, activity, osmolality, osmotic pressure, oncotic pressure, and pH.

The seven base units of the SI system.

Unit	Symbol	Dimension
Meter	m	length
Kilogram	kg	mass
Second	S	time
Mole	mol	amount of substance
Ampere	Α	electric current
Kelvin	K	temperature (absolute)
Candela	cd	luminous intensity

The base units are precisely defined autonomous units. All other units are derived by multiplying or dividing base units and are therefore referred to as **derived units**, e.g.:

- Area (length \cdot length): $m \cdot m = m^2$
- Velocity (length/time): $m/s = m \cdot s^{-1}$.

If the new unit becomes too complicated, it is given a new name and a corresponding symbol, e.g., force = $m \cdot kg \cdot s^{-2} = N (\rightarrow Table 1)$.

Fractions and Multiples of Units

Prefixes are used to denote decimal multiples and fractions of a unit since it is both tedious and confusing to write large numbers. We generally write 10 kg (kilograms) and $10\,\mu g$ (micrograms) instead of $10\,000\,g$ and $0.00001\,g$, for example. The prefixes, which are usually varied in 1000-unit increments, and the corresponding symbols and conversion factors are listed in Table 2. Prefixes are used with base units and the units derived from them (\rightarrow Table 1), e.g., $10^3\,Pa = 1\,kPa$. Decimal increments are used in some cases (e.g., da, h, d, and c; \rightarrow Table 2). Time is given in conventional nondecimal units, i.e., seconds (s), minutes (min), hours (h), and days (d).

Length, Area, Volume

The *meter* (m) is the SI unit of **length**. Other units of length have also been used.

Examples:

1 ångström (Å) = 10^{-10} m = 0.1 nm 1 migrap (μ) = 10^{-6} m = 1 μ m

1 micron (μ) = 10⁻⁶ m = 1 μ m 1 millimicron ($m\mu$) = 10⁻⁹ m = 1 nm

American and British units of length:

1 inch = 0.0254 m = 25.4 mm

1 foot = 0.3048 m

1 yard = 3 feet = 0.9144 m

1 (statute) mile = $1609.344 \, \text{m} \approx 1.61 \, \text{km}$

1 nautical mile = 1.853 km

The square meter (m²) is the derived SI unit of area, and the cubic meter (m²) is the corresponding unit of volume. When denoting the fractions or multiples of these units with prefixes (Table 2), please note that there are some peculiarities.

Examples:

 $1 \text{ m} = 10^3 \text{ mm, but}$

 $1 \text{ mm}^2 = 10^6 \text{ mm}^2$, and

 $1 \text{ m}^3 = 10^9 \text{ mm}^3$

The *liter* (L or l)* is often used as a unit of *volume* for liquids and gases:

 $1 L = 10^{-3} \text{ m}^3 = 1 \text{ dm}^3$

 $1 \text{ mL} = 10^{-6} \text{ m}^3 = 1 \text{ cm}^3$

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 $1 \mu L = 10^{-9} \,\mathrm{m}^3 = 1 \,\mathrm{mm}^3$.

Table 1 Derived units based on SI base units m, kg, s, cd, and A

Coulomb	С	electrical charge	s · A
Farad	F	electrical capacitance	$C \cdot V^{-1} = m^{-2} \cdot kg^{-1} \cdot s^4 \cdot A^2$
Hertz	Hz	frequency	s ⁻¹
Joule	J	heat, energy, work	$N \cdot m = m^2 \cdot kg \cdot s^{-2}$
Lumen	lm	light flux	cd · sr
Lux	lx	light intensity	$Im \cdot m^{-2} = cd \cdot sr \cdot m^{-2}$
Newton	N	force	m⋅kg⋅s ⁻²
Ohm	Ω	electrical resistance	$V \cdot A^{-1} = m^2 \cdot kg \cdot s^{-3} \cdot A^{-2}$
Pascal	Р	pressure	$N \cdot m^{-2} = m^{-1} \cdot kg \cdot s^{-2}$
Siemens	S	conductivity	$\Omega^{-1} = m^{-2} \cdot kg^{-1} \cdot s^3 \cdot A^2$
Steradian	SΓ	measure of solid angle ¹	1 (m ² · m ⁻²)
Tesla	T	magnetic flux density	$Wb \cdot m^{-2} = kg \cdot s^{-2} \cdot A^{-1}$
Volt	V	electric potential	$W \cdot A^{-1} = m^2 \cdot kg \cdot s^{-3} \cdot A^{-1}$
Watt	W	electric power	$J \cdot s^{-1} = m^2 \cdot kg \cdot s^{-3}$
Weber	Wb	magnetic flux	$V \cdot s = m^2 \cdot kg \cdot s^{-2} \cdot A^{-1}$

¹ The solid angle of a sphere is defined as the angle subtended at the center of a sphere by an area (A) on its surface times the square of its radius (r²). A steradian (sr) is the solid angle for which r = 1 m and A = 1 m², that is, 1 sr = 1 m²/m²².

Table 2 Prefixes for fractions and multiples of units of measure

Prefix	Symbol	Factor	Prefix	Symbol	Factor
deca-	da	10 ¹	deci-	d	10-1
hecto-	h	10 ²	centi-	С	10-2
kilo-	k	10 ³	milli-	m	10-3
mega-	M	10 ⁶	micro-	μ	10-6
giga-	G	10 ⁹	nano-	n	10-9
tera-	T	10 ¹²	pico-	р	10 ⁻¹²
peta-	Р	10 ¹⁵	femto-	f	10 ⁻¹⁵
exa-	E	10 ¹⁸	atto-	a	10 ⁻¹⁸

Conversion of American and British volume units into SI units:

1 fluid ounce (USA) = 29.57 mL 1 fluid ounce (UK) = 28.47 mL 1 liquid gallon (USA) = 3.785 L 1 liquid gallon (UK) = 4.546 L 1 pint (USA) = 473.12 mL 1 pint (UK) = 569.4 mL

Velocity, Frequency, Acceleration

Velocity is the distance traveled per unit time $(m \cdot s^{-1})$. This is an expression of **linear velocity**, whereas "**volume velocity**" is used to express the *volume flow per unit time*. The latter is expressed as $L \cdot s^{-1}$ or $m^3 \cdot s^{-1}$.

Frequency is used to describe how often a periodic event (pulse, breathing, etc.) occurs per unit time. The SI unit of frequency is s⁻¹ or *hertz* (Hz). min⁻¹ is also commonly used:

 $min^{-1} = 1/60 \text{ Hz} \approx 0.0167 \text{ Hz}.$

Acceleration, or *velocity change per unit time*, is expressed in $m \cdot s^{-1} \cdot s^{-1} = m \cdot s^{-2}$. Since *deceleration* is equivalent to negative acceleration, acceleration and deceleration can both be expressed in $m \cdot s^{-2}$.

Force and Pressure

Force equals mass times acceleration. Weight is a special case of force as weight equals mass times acceleration of gravity. Since the unit of mass is kg and that of acceleration $m \cdot s^{-2}$, force is expressed in $m \cdot kg \cdot s^{-2} = newton$ (N). The older units of force are converted into N as follows:

1 dyn = 10^{-5} N = $10 \,\mu\text{N}$ 1 pond = $9.8 \cdot 10^{-3}$ N = 9.8 mN.

Pressure equals *force per unit area*, so the SI unit of pressure is $N \cdot m^{-2} = pascal$ (Pa). However, the pressure of bodily fluids is usually measured in mmHg. This unit and other units are converted into SI units as follows:

 $\begin{array}{l} 1\ mm\ H_2O\approx 9.8\ Pa \\ 1\ cm\ H_2O\approx 98\ Pa \\ 1\ mm\ Hg = 133.3\ Pa = 0.1333\ kPa \\ 1\ torr = 133.3\ Pa = 0.1333\ kPa \\ 1\ technical\ atmosphere\ (at)\approx 98.067\ kPa \\ 1\ physical\ atmosphere\ (atm) = 101.324\ kPa \\ 1\ dyne\cdot cm^{-2} = 0.1\ Pa \\ 1\ bar = 100\ kPa. \end{array}$

Work, Energy, Heat, Power

Work equals *force times distance*, $N \cdot m = J$ (*joule*), or *pressure times volume*, $(N \cdot m^{-2}) \cdot m^3 = I$. **Energy** and **heat** are also expressed in I.

Other units of work, heat, and energy are converted into I as follows:

1 erg = 10^{-7} J = $0.1 \,\mu$ J 1 cal $\approx 4.185 \,$ J 1 kcal $\approx 4185 \,$ J = $4.185 \,$ kJ 1 Ws = 1 J 1 kWh = $3.6 \cdot 10^6 \,$ J = $3.6 \,$ MJ.

Power equals *work per unit time* and is expressed in *watts* (W), where W = $J \cdot s^{-1}$. Heat flow is also expressed in W. Other units of power are converted into W as follows:

 $1 \text{ erg} \cdot \text{s}^{-1} = 10^{-7} \text{ W} = 0.1 \,\mu\text{W}$ $1 \text{ cal} \cdot \text{h}^{-1} = 1.163 \cdot 10^{-3} \text{ W} = 1.163 \,\text{mW}$ $1 \text{ metric horse power (hp)} = 735.5 \,\text{W} = 1.163 \,\text{m}$

Mass, Amount of Substance

0.7355 kW.

The base unit of **mass** is the kilogram (kg), which is unusual insofar as the base unit bears the prefix "kilo". Moreover, 1000 kg is defined as a *metric ton** instead of as a megagram. **Weight** is the product of mass and gravity (see above), but weight scales are usually calibrated in units of mass (g, kg).

British and American units of mass are converted into SI units as follows.

Avoirdupois weight:
1 ounce (oz.) = 28.35 g
1 pound (lb.) = 453.6 g
Apothecary's and troy weight:
1 ounce = 31.1 g
1 pound = 373.2 g.

The mass of a molecule or an atom (**molecular** or **atomic mass**) is often expressed in *daltons* (Da)*. 1 Da = 1/12 the mass of a 12 C atom, equivalent to 1 kg/Avogadro's constant = $1 \text{ kg/}(6.022 \cdot 10^{23})$:

1 Da = $1.66 \cdot 10^{-27}$ kg 1000 Da = 1 kDa.

375

The **relative molecular mass (M_r)**, or *molecular "weight"*, is the molecular mass of a substance divided by 1/12 the mass of a ^{12}C atom. Since M_r is a ratio, it is a dimensionless unit.

The **amount of substance**, or *mole* (mol), is related to mass. One mole of substance contains as many elementary particles (atoms, molecules, ions) as 12 g of the nuclide of a ¹²C atom = 6.022 · 10²³ particles. The *conversion factor between moles and mass* is therefore: 1 mole equals the mass of substance (in grams) corresponding to the relative molecular, ionic, or atomic mass of the substance. In other words, it expresses how much higher the mass of the atom, molecule, or ion is than 1/12 that of a ¹²C atom.

Examples:

- Relative molecular mass of H2O: 18
 - \rightarrow 1 mol H₂O = 18 g H₂O.
- Relative atomic mass of Na⁺: 23
 - \rightarrow 1 mol Na+ = 23 g Na+.
- Relative molecular mass of CaCl₂:
 - $=40+(2\cdot35.5)=111$
 - \rightarrow 1 mol CaCl₂ = 111 g CaCl₂.

(CaCl₂ contains 2 mol Cl⁻ and 1 mol Ca²⁺.)

The **equivalent mass** is calculated as moles divided by the valency of the ion in question and expressed in equivalents (Eq)*. The mole and equivalent values of monovalent ions are identical:

 $1 \text{ Eq Na}^+ = 1/1 \text{ mol Na}^+$.

For bivalent ions, equivalent = 1/2 mole:

1 Eq Ca²⁺ = $\frac{1}{2}$ mole Ca²⁺ or 1 mole Ca²⁺ = 2 Eq $\frac{1}{2}$

The **osmole (Osm)** is also derived from the mole (see below).

Electrical Units

Electrical **current** is the flow of charged particles, e.g., of electrons through a wire or of ions through a cell membrane. The number of particles moving per unit time is measured in *amperes* (A). Electrical current cannot occur unless there is an **electrical potential difference**, in short also called potential, voltage, or tension. Batteries and generators are used to create such potentials. Most electrical potentials in the body are generated by ionic flow $(\rightarrow p. 32)$. The *volt* (V) is the SI unit of electrical potential $(\rightarrow Table \ 1)$.

How much electrical current flows at a given potential depends on the amount of **electrical resistance**, as is described in *Ohm's law*

(voltage = current · resistance). The unit of electrical resistance is ohm (Ω) (\rightarrow **Table 1**). Conductivity is the reciprocal of resistance ($1/\Omega$) and is expressed in siemens (S), where S = Ω^{-1} . In physiology, resistance is related to the membrane surface area ($\Omega \cdot m^2$). The reciprocal of this defines the **membrane conductance** to a given ion: $\Omega^{-1} \cdot m^{-2} = S \cdot m^{-2} (\rightarrow p. 32)$.

Electrical **work** or **energy** is expressed in *joules* (J) or *watt seconds* (Ws), whereas electrical **power** is expressed in *watts* (W).

The electrical **capacitance** of a capacitor, e.g., a cell membrane, is the ratio of *charge* (C) to *potential* (V); it is expressed in *farads* (F) (\rightarrow **Table 1**).

Direct current (DC) always flows in one direction, whereas the direction of flow of **alternating current** (AC) constantly changes. The **frequency** of one cycle of change per unit time is expressed in *hertz* (Hz). Mains current is generally 60 Hz in the USA and 50 Hz in Europe.

Temperature

Kelvin (K) is the SI unit of temperature. The lowest possible temperature is 0 K, or *absolute zero*. The **Celsius** or **centigrade** scale is derived from the Kelvin scale. The temperature in degrees Celsius (°C) can easily be converted into K:

 $^{\circ}$ C = K - 273.15.

In the USA, temperatures are normally given in degrees Fahrenheit (°F). Conversions between Fahrenheit and Celsius are made as follows:

$$^{\circ}F = (9/5 \cdot ^{\circ}C) + 32$$

 $^{\circ}C = (^{\circ}F - 32) \cdot 5/9.$

Some important Kelvin, Celsius, and Fahrenheit temperature equivalents:

K	°C	°F
+ 273	0°	+32°
293- 298	20°- 25°	68°-77°
310	37°	98.6°
311– 315	38°- 42°	100°- 108°
373	100°	212°
	+ 273 293- 298 310 311- 315	+ 273 0° 293- 20°- 298 25° 310 37° 311- 38°- 315 42°

Concentrations, Fractions, Activity

The word *concentration* is used to describe many different relationships in physiology and medicine. Concentration of a substance X is often abbreviated as [X]. Some concentrations are listed below:

- -Mass concentration, or the mass of a substance per unit volume (e.g., $g/L = kg/m^3$)
- -Molar concentration, or the amount of a substance per unit volume (e.g., mol/L)
- -Molal concentration, or the amount of substance per unit mass of solvent (e.g., mol/ $kg\,H_2O$).

The SI unit of mass concentration is g/L (kg/m^3 , mg/L, etc.). The conversion factors for older units are listed below:

 $\begin{array}{l} 1 \text{ g}/100 \text{ mL} = 10 \text{ g/L} \\ 1 \text{ g\%} = 10 \text{ g/L} \\ 1 \text{ \%} & (\text{w/v}) = 10 \text{ g/L} \\ 1 \text{ g\%} = 1 \text{ g/L} \\ 1 \text{ mg\%} = 10 \text{ mg/L} \\ 1 \text{ mg/} = 10 \text{ mg/L} \\ 1 \text{ mg/} 100 \text{ mL} = 10 \text{ mg/L} \\ 1 \text{ <math>\mu\text{g\%}} = 10 \text{ } \mu\text{g/L} \\ 1 \text{ <math>\gamma\%} = 10 \text{ } \mu\text{g/L}. \end{array}$

Molarity is the **molar concentration**, which is expressed in mol/L (or mol/m³, mmol/L, etc.). Conversion factors are listed below:

1 M (molar) = 1 mol/L 1 N (normal) = (1/valency) · mol/L 1 mM (mmolar) = 1 mmol/L

 $1 \text{ Eq/L} = (1/\text{valency}) \cdot \text{mol/L}.$

In highly diluted solutions, the only difference between the molar and molal concentrations is that the equation "1 L H_2O = 1 kg H_2O " holds at only one particular temperature (4°C). Biological fluids are not highly diluted solutions. The volume of solute particles often makes up a significant fraction of the overall volume of the solution. One liter of plasma, for example, contains 70 mL of proteins and salts and only 0.93 L of water. In this case, there is a 7% difference between molarity and molality. Differences higher than 30% can occur in intracellular fluid. Although molarity is more commonly measured (volumetric measurement), molality plays a more important role in biophysical and biological processes and chemical reactions.

The **activity** (a) of a solution is a thermodynamic measure of its physicochemical efficacy. In physiology, the activity of ions is measured by ion-sensi-

tive electrodes (e.g., for H * , Na * , K * , Cl * , or Ca 2*). The activity and molality of a solution are identical when the total **ionic strength** (μ) of the solution is very small, e.g., when the solution is an ideal solution. The ionic strength is dependent on the charge and concentration of all ions in the solution:

$$\mu \equiv 0.5 (z_1^2 \cdot c_1 + z_2^2 \cdot c_2 + \dots + z_i^2 \cdot c_i)$$
 [13.1]

where z_i is the valency and c_i the molal concentration of a given ion "i", and 1, 2, etc. represent the different types of ions in the solution. Owing to the high ionic strength of biological fluids, the solute particles influence each other. Consequently, the activity (a) of a solution is always significantly lower than its molar concentration (c). Activity is calculated as $a = f \cdot c$, where f is the **activity coefficient**.

Example: At an ionic strength of 0.1 (as it is the case for a solution containing 100 mmol NaCl/ kg $\rm H_2O$), f = 0.76 for Na $^+$. The activity important in biophysical processes is therefore roughly 25% lower than the molality of the solution.

In solutions that contain weak electrolytes which do not completely dissociate, the molality and activity of free ions also depend on the degree of electrolytic dissociation.

Fractions ("fractional concentrations") are relative units:

- Mass ratio, i.e., mass fraction relative to total mass
- Molar ratio
- Volume ratio, i.e. volume fraction relative to total volume. The volume fraction (F) is commonly used in respiratory physiology.

Fractions are expressed in units of g/g, mol/mol, and L/L respectively, i.e. in "units" of 1, 10^{-3} , 10^{-6} , etc. The unabbreviated unit (e.g., g/g) should be used whenever possible because it identifies the type of fraction in question. The fractions %, ‰, ppm (parts per million), and ppb (parts per billion) are used for all types of fractions.

Conversion:

1% = 0.01 $1\% = 1 \cdot 10^{-3}$ 1 vol% = 0.01 L/L $1 \text{ ppm} = 1 \cdot 10^{-6}$

1 ppb = $1 \cdot 10^{-9}$

Osmolality, Osmotic / Oncotic Pressure

Osmolarity (Osm/L), a unit derived from molarity, is the *concentration of all osmotically active particles* in a solution, regardless of which compounds or mixtures are involved.

376

However, measurements with osmometers as well as the biophysical application of osmotic concentration refer to the number of osmoles per unit volume of *solvent* as opposed to the total volume of the solution. This and the fact that volume is temperature-dependent are the reasons why **osmolality** (Osm/kg H₂O) is generally more suitable.

Ideal osmolality is derived from the molality of the substances in question. If, for example, 1 mmol (180 mg) of glucose is dissolved in 1 kg of water (1L at 4°C), the molality equals 1 mmol/kg H_2O and the ideal osmolality equals 1 mOsm/kg H_2O . This relationship changes when electrolytes that dissociate are used, e.g., $NaCl \rightleftharpoons Na^+ + Cl^-$. Both of these ions are osmotically active. When a substance that dissociates is dissolved in 1 kg of water, the ideal osmolality equals the molality times the number of dissociation products, e.g., 1 mmol $NaCl/kg H_2O = 2 mOsm/kg H_2O$.

Electrolytes weaker than NaCl do not dissociate completely. Therefore, their *degree of electrolytic dissociation* must be considered.

These rules apply only to ideal solutions, i.e., those that are extremely dilute. As mentioned above, bodily fluids are **nonideal (or real) solutions** because their real osmolality is lower than the ideal osmolality. The real osmolality is calculated by multiplying the ideal osmolality by the *osmotic coefficient* (g). The osmotic coefficient is concentration-dependent and amounts to, for example, approximately 0.926 for NaCl with an (ideal) osmolality of this NaCl solution thus amounts to $0.926 \cdot 300 = 278 \, \text{mOsm/kg} \, \text{H}_2\text{O}$.

Solutions with a real osmolality equal to that of plasma ($\approx 290 \, \text{mOsm/kg H2 O}$) are said to be isosmolal. Those whose osmolality is higher or lower than that of plasma are hyperosmolal or hyposmolal.

Osmolality and Tonicity

Each osmotically active particle in solution (cf. real osmolality) exerts an **osmotic pressure** (π) as described by *van't Hoff's equation*:

$$\pi = R \cdot T \cdot c_{osm}$$
 [13.2]

where R is the universal gas constant (8.314 J \cdot K⁻¹ \cdot Osm⁻¹), T is the absolute temperature in K, and c_{osm} is the real osmolality in Osm \cdot (m³

 $H_2O)^{-1} = mOsm \cdot (LH_2O)^{-1}$. If two solutions of different osmolality (Δc_{osm}) are separated by a water-permeable selective membrane, Δc_{osm} will exert an **osmotic pressure difference** ($\Delta \pi$) across the membrane in steady state if the membrane is less permeable to the solutes than to water. In this case, the selectivity of the membrane, or its relative impermeability to the solutes, is described by the reflection coefficient (a), which is assigned a value between 1 (impermeable) and 0 (as permeable as water). The reflection coefficient of a semipermeable membrane is $\sigma = 1$. By combining van't Hoff's and Staverman's equations, the osmotic pressure difference ($\Delta \pi$) can be calculated as follows:

$$\Delta \pi = \sigma \cdot \mathbf{R} \cdot \mathbf{T} \cdot \Delta \mathbf{c}_{osm}. \tag{13.3}$$

Equation 13.3 shows that a solution with the same osmolality as plasma will exert the same osmotic pressure on a membrane in steady state (i.e., that the solution and plasma will be isotonic) only if $\sigma = 1$. In other words, the membrane must be strictly semipermeable.

Isotonicity, or equality of osmotic pressure, exists between plasma and the cytosol of red blood cells (and other cells of the body) in steady state. When the red cells are mixed in a urea solution with an osmolality of 290 mOsm/kg $\rm H_2O$, isotonicity does not prevail after urea ($\sigma < 1$) starts to diffuse into the red cells. The interior of the red blood cells therefore becomes hypertonic, and water is drawn inside the cell due to osmosis (\rightarrow p. 24). As a result, the erythrocytes continuously swell until they burst.

An **osmotic gradient** resulting in the subsequent flow of water therefore occurs in all parts of the body in which dissolved particles pass through water-permeable cell membranes or cell layers. This occurs, for example, when Na¹ and Cl⁻ pass through the epithelium of the small intestine or proximal renal tubule. The extent of this water flow or *volume flow Jv* ($m^3 \cdot s^{-1}$) is dependent on the *hydraulic conductivity* k ($m \cdot s^{-1} \cdot Pa^{-1}$) of the membrane (i.e., its permeability to water), the *area* A of passage (m^2), and the pressure difference, which, in this case, is equivalent to the osmotic pressure difference $\Delta \pi$ (Pa):

$$[\mathbf{v} = \mathbf{k} \cdot \mathbf{A} \cdot \Delta \pi \, [\mathbf{m}^3 \cdot \mathbf{s}^{-1}]. \quad [13.4]$$

Since it is normally not possible to separately determine k and A of a biological membrane or cell layer, the product of the two $(k \cdot A)$ is often calculated as the *ultrafiltration coefficient* K_f $(m^3 s^{-1} Pa^{-1})$ (cf. p. 152).

The transport of osmotically active particles causes water flow. Inversely, flowing water drags dissolved particles along with it. This type of solvent $drag \, (\to p.24)$ is a form of convective transport.

Solvent drag does not occur if the cell wall is impermeable to the substance in question (σ = 1). Instead, the water will be retained on the side where the substance is located. In the case of the aforementioned epithelia, this means that the substances that cannot be reabsorbed from the tubule or intestinal lumen lead to osmotic diuresis (\rightarrow p. 172) and diarrhea respectively. The latter is the mechanism of action of saline laxatives (\rightarrow p. 262).

Oncotic Pressure / Colloid Osmotic Pressure

As all other particles dissolved in plasma, macromolecular proteins also exert an osmotic pressure referred to as oncotic pressure or colloid osmotic pressure. Considering its contribution of only 3.5 kPa (25 mm Hg) relative to the total osmotic pressure of the small molecular components of plasma, the oncotic pressure on a strictly semipermeable membrane could be defined as negligible. However, within the body, oncotic pressure is so important because the endothelium that lines the blood vessels allows small molecules to pass relatively easily ($\sigma \approx 0$). According to equation 13.3, their osmotic pressure difference $\Delta\pi$ at the endothelium is virtually zero. Consequently, only the oncotic pressure difference of proteins is effective, as the endothelium is either partly or completely impermeable to them, depending on the capillary segment in question. Because the protein reflection coefficient $\sigma \gg 0$ and the protein content of the plasma (ca. 75 g/L) are higher than that of the interstitium, these two factors counteract filtration, i.e., the blood pressure-driven outflow of plasma water from the endothelial lumen, making the endothelium an effective volume barrier between the plasma space and the interstitium.

If the blood pressure drives water out of the blood into the interstitium (filtration), the

plasma protein concentration and thus the oncotic pressure difference π will rise (\rightarrow pp. 152, 208). This rise is much higher than equation 13.3 leads one to expect (\rightarrow A). The difference is attributable to specific biophysical properties of plasma proteins. If there is a pressure-dependent efflux or influx of water out of or into the bloodstream, these relatively high changes in oncotic pressure difference automatically exert a counterpressure that limits the flow of water.

pH, pK, Buffers

The **pH** indicates the hydrogen ion $[H^+]$ concentration of a solution. According to *Sörensen*, the pH is the negative common logarithm of the molal H^+ concentration in mol/kg H_2O .

Examples:

 $1 \; mol/kg \, H_2O = 10^0 \, mol/kg \, H_2O = pH \, 0, \\ 0.1 \; mol/kg \, H_2O = 10^{-1} \, mol/kg \, H_2O = pH \, 1, \\ and \; so \; on \; up \; to \; 10^{-14} \, mol/kg \, H_2O = pH \, 14.$

Since glass electrodes are normally used to measure the pH, the H^+ activity of the solution is actually being determined. Thus, the following rule applies: $pH = -\log (f_H \cdot [H^+])$,

where f_H is the activity coefficient of H^* . Considering its ionic concentration (see above), the f_H of plasma is ≈ 0.8 .

The logarithmic nature of pH must be considered when observing pH changes. For example, a rise in pH from 7.4 (40 nmol/kg H_2O) to pH 7.7 decreases the H^+ activity by 20 nmol/kg H_2O , whereas an equivalent decrease (e.g., from pH 7.4 to pH 7.1) increases the H^+ activity by 40 nmol/kg H_2O .

The **pK** is fundamentally similar to the pH. It is the negative common logarithm of the *dissociation constant* of an acid (K_a) or of a base (K_b) :

$$pK_a = -\log K_a$$

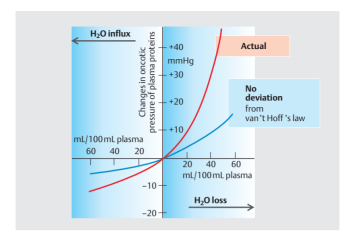
$$pK_b = -\log K_b$$
.

For an acid and its corresponding base, $pK_a + pK_b = 14$, so that the value of pK_a can be derived from that of pK_b and vice versa.

The *law of mass actions* applies when a weak acid (AH) dissociates:

$$AH \rightleftharpoons A^- + H^+$$
 [13.5]

It states that the product of the molal concentration (indicated by square brackets) of the dissociation products divided by the concentration of the nondissociated substance remains constant:



A. Physiological signification of deviations on oncotic pressure of plasma from van't Hoff's equation. A loss of water from plasma leads to a disproportionate rise in oncotic pressure, which counteracts the water loss. Conversely, the dilution of plasma due to the influx of water leads to a disproportionate drop in oncotic pressure, though less pronounced. Both of these are important mechanisms for maintaining a constant blood volume and preventing edema. (Adapted from Landis EM u. Pappenheimer JR. Handbook of Physiology. Section 2: Circulation, Vol. II. American Physiological Society: Washington D.C. 1963, S. 975.)

$$K_a = \frac{[A^-] \cdot [H^+] \cdot f_H}{[AH]}$$
 [13.6]

Converted into logarithmic form (and inserting H⁺ activity for [H⁺]), the equation is transformed into:

$$\log K_a = \log \frac{[A^-]}{[AH]} + \log ([H^+] \cdot f_H)$$
 [13.7]

or

$$-\log([H^+] \cdot f_H) = -\log K_a + \log \frac{[A^-]}{[AH]} \quad [13.8]$$

Based on the above definitions for pH and pK_a , it can also be converted into

$$pH = pK_a + log \frac{[A^-]}{[AH]}$$
 [13.9]

Because the concentration and not the activity of A^ and AH is used here, pK_a is concentration-dependent in nonideal solutions.

Equation 13.9 is the general form of the **Henderson–Hasselbalch equation** (\rightarrow p. 138ff.), which describes the relationship between the pH of a solution and the concentration ratio of a dissociated to an undissociated form of a solute. If [A⁻] = [AH], then the concentration ratio is 1/1 = 1, which corresponds to pH = pK_a since the log of 1 = 0.

A weak acid (AH) and its dissociated salt (A⁻) form a **buffer system** for H⁺ and OH⁻ ions: Addition of H⁺ yields A⁻ + H⁺ → AH

Addition of OH⁻ yields AH + OH⁻ \rightarrow A⁻ + H₂O. The *buffering power* of a buffer system is greatest when [AH] = [A⁻], i.e., when the pH of the solution equals the pK_a of the buffer.

Example: Both [A⁻] and [AH] = 10 mmol/L and pK_a = 7.0. After addition of 2 mmol/L of H⁺ ions, the [A⁻]/ [AH] ratio changes from 10/10 to 8/12 since 2 mmol/L of A⁻ are consequently converted into 2 mmol/L of AH. Since the log of 8/12 \approx -0.18, the pH decreases

by 0.18 units to pH 6.82. If the initial $[A^-]$ /[AH] ratio had been 3/17, the pH would have dropped from an initial pH 6.25 (7 plus the log of 3/17 = 6.25) to pH 5.7 (7 + log of 1/19 = 5.7), i.e., by 0.55 pH units after addition of the same quantity of H⁺ ions.

The titration of a buffer solution with H^+ (or OH^-) can be plotted to generate a **buffering curve** (\rightarrow **B**). The steep part of the curve represents the range of the best buffering power. The pK_a value lies at the *turning point* in the middle of the steep portion of the curve. Substances that gain (or lose) more than one H^+ per molecule have more than one pK value and can therefore exert an optimal buffering action in several regions. Phosphoric acid (H_3PO_4) donates three H^+ ions, thereby successively forming $H_2PO_4^-$, HPO_4^{2-} , and PO_4^{3-} . The buffer pair $HPO_4^{2-}/H_2PO_4^-$ with a pK_a of 6.8 is important in human physiology (\rightarrow p. 174ff.).

The absolute slope, $d[A^-]/d(pH)$, of a buffering curve (plot of pH vs. $[A^-]$) is a measure of **buffering capacity** $(mol \cdot L^{-1} \cdot [\Delta pH]^{-1}; \rightarrow p.$ 138).

Powers and Logarithms

Powers of ten are used to more easily and conveniently write numbers that are much larger or smaller than 1.

Examples:

 $100 = 10 \cdot 10 = 10^2$

 $1000 = 10 \cdot 10 \cdot 10 = 10^3$

 $10\,000 = 10 \cdot 10 \cdot 10 \cdot 10 = 10^4$, etc.

In this case, the exponent denotes the amount of times ten is multiplied by itself. If the number is not an exact power of ten (e.g., $34\,500$), divide it by the next lowest decimal power ($10\,000$) and use the quotient (3.45) as a multiplier to express the result as $3.45\cdot10^4$.

The number 10 can also be expressed exponentially (101). Numbers much smaller than 1 are annotated using negative exponents.

Examples:

 $1 = 10 \div 10 = 10^{0}$

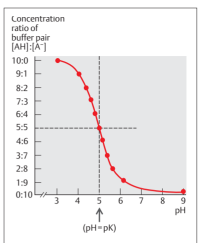
 $0.1 = 10 \div 10 \div 10 = 10^{-1}$

 $0.01 = 10 \div 10 \div 10 = 10^{-2}$, etc.

Similar to the large numbers above, numbers that are not exact powers of ten are expressed using multipliers, e.g.,

 $0.04 = 4 \cdot 0.01 = 4 \cdot 10^{-2}$.

Note: When writing numbers smaller than 1, the (negative) exponent corresponds to the



B. Buffering Curve. Graphic representation of the relationship between pH and the concentration ratio of buffer acid/buffer base [AH]/[A^-] as a function of pH. The numerical values are roughly equivalent to those of the buffer pair acetic acid/acetate (pKa = 4.7). The buffering power of a buffer system is greatest when [AH] = [A^-], i.e., when the pH of the solution equals the pKa of the buffer (broken lines).

position of the 1 after the decimal point; therefore, $0.001 = 10^{-3}$. When writing numbers greater than 10, the exponent corresponds to the number of decimal positions to the left of the decimal point minus 1; therefore, $1124.5 = 1.245 \cdot 10^{3}$.

Exponents can also be used to represent units of measure, e.g., m^3 . As in the case of 10^3 , the base element (meters) is multiplied by itself three times ($m \cdot m \cdot m$; \rightarrow p. 372). Negative exponents are also used to express units of measure. As with $1/10 = 10^{-1}$, 1/s can be written as s^{-1} , mol/L as $mol \cdot L^{-1}$, etc.

There are specific rules for performing calculations with powers of ten. Addition and subtraction are possible only if the exponents are identical, e.g.,

 $(2.5 \cdot 10^2) + (1.5 \cdot 10^2) = 4 \cdot 10^2$.

Unequal exponents, e.g., $(2 \cdot 10^3) + (3 \cdot 10^2)$, must first be equalized:

 $(2 \cdot 10^3) + (0.3 \cdot 10^3) = 2.3 \cdot 10^3$.

The exponents of the multiplicands are added together when multiplying powers of 10, and the denominator is subtracted from the numerator when dividing powers of ten.

Examples:

$$10^2 \cdot 10^3 = 10^{2+3} = 10^5$$

 $10^4 \div 10^2 = 10^{4-2} = 10^2$
 $10^2 \div 10^4 = 10^{2-4} = 10^{-2}$

The usual mathematical rules apply to the multipliers of powers of ten, e.g.,

$$(3 \cdot 10^2 \cdot (2 \cdot 10^3) = (2 \cdot 3) \cdot (10^{2+3}) = 6 \cdot 10^5.$$

Logarithms. There are two kinds of logarithms: common and natural. Logarithmic calculations are performed using exponents alone. The **common** (**decimal**) **logarithm** (log or lg) is the power or exponent to which 10 must be raised to equal the number in question. The common logarithm of $100 (\log 100)$ is 2, for example, because $10^2 = 100$. Decimal logarithms are commonly used in physiology, e.g., to define pH values (see above) and to plot the pressure of sound on a decibel scale $(\rightarrow p.363)$.

Natural logarithms (ln) have a natural base of 2.71828..., also called *base e*. The common logarithm ($\log x$) equals the natural logarithm of x ($\ln x$) divided by the natural logarithm of 10 ($\ln 10$), where $\ln 10 = 2.302585$. The following rules apply when converting between natural and common logarithms:

$$\log x = (\ln x)/2.3$$

$$\ln x = 2.3 \cdot \log x.$$

When performing mathematical operations with logarithms, the type of operation is reduced by one rank—multiplication becomes addition, potentiation becomes multiplication, and so on.

Examples:

Examples.
$$\log(a \cdot b) = \log a + \log b$$

$$\log(a/b) = \log a - \log b$$

$$\log a^n = n \cdot \log a$$

$$\log a^n = n \cdot \log a$$

$$\log \sqrt[n]{a} = (\log a)/n$$
Special cases:
$$\log 10 = \ln e = 1$$

$$\log 1 = \ln 1 = 0$$

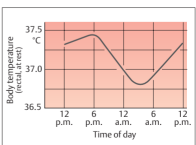
$$\log 0 = \ln 0 = \pm \infty$$

Graphic Representation of Data

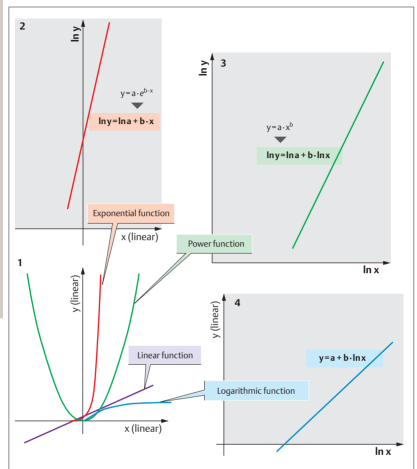
Graphic plots of data are used to provide a clear and concise representation of measurements, e.g., body temperature over the time of day $(\rightarrow C)$. The axes on which the measurements (e.g., temperature and time) are plotted are called coordinates. The vertical axis is referred to as the ordinate (temperature) and the horizontal axis is the abscissa (time). It is customary to plot the first variable x (time) on the abscissa and the other dependent variable v (temperature) on the ordinate. The abscissa is therefore called the x-axis and the ordinate the *y-axis*. This method of graphically plotting data can be used to illustrate the connection between any two related dimensions imaginable. e.g., to describe the relationship between height and age, lung capacity and intrapulmonary pressure, etc. (\rightarrow p. 117).

Plotting of data makes it easier to determine whether two variables correlate with each other. For example, the plot of height (ordinate) over age (abscissa) shows that the height increases during the growth years and reaches a plateau at the age of about 17 years. This means that height is related to age in the first phase of life, but is largely independent of age in the second phase. A correlation does not necessarily indicate a causal relationship. A decrease in the birth rate in Alsace-Lorraine, for example, correlated with a decrease in the number or nesting storks for a while.

When plotting variables of wide-ranging dimensions (e.g., 1 to 100000) on a coordinate



C. Illustration of how to plot data on a coordinate system. The plot in this example shows the relationship between body temperature (rectal, at rest) and time of day.



D. Types of functions. D1: Linear function (violet), exponential function (red), logarithmic function (blue), and power function (green) showing linear plotting of data on both axes. The three curves can be made into a straight line (linearized) by logarithmically plotting the data on the y-axis (**D2:** exponential function) or on the x-axis (**D4:** logarithmic function) or both (**D3:** power function).

system, it can be impossible to plot small values individually without having the axes become extremely long. This problem can be solved by plotting the data as powers of 10 or logarithms. For example, 1, 10, 100, and 1000 are written as 10⁰, 10¹, 10², and 10³ or as logarithms 0, 1, 2, and 3. This makes it possible to

obtain a relatively accurate graphic representation of very small numbers, and all the numbers fit on an axis of reasonable length (cf. sound curves on p. 363 B).

Correlations can be either linear or non-linear. *Linear correlations* (\rightarrow **D1**, violet line) obey the linear relationship

y = ax + b,

where a is the slope of the line and b is the point, or *intercept* (at x = 0), where it intersects the v-axis.

Many correlations are nonlinear. For simpler functions, graphic linearization can be achieved via a nonlinear (logarithmic) plot of the x and/or y values. This allows for the extrapolation of values beyond the range of measurement (see below) or for the generation of calibration curves from only two points. In addition, this method also permits the calculation of the "mean" correlation of scattered x-y pairs using regression lines.

An **exponential function** (\rightarrow **D1**, red curve), such as

 $y = a \cdot e^{b \cdot x}$

can be linearized by plotting $\ln y$ on the y-axis $(\rightarrow D2)$:

 $\ln y = \ln a + b \cdot x$,

where b is the slope and ln a is the intercept.

A **logarithmic function** (\rightarrow **D1**, blue curve), such as

 $y = a + b \cdot \ln x$,

can be linearized by plotting $\ln x$ on the x-axis (\rightarrow D4), where b is the slope and a is the intercept.

A power function (\rightarrow D1, green curve), such as

$$y = a \cdot x^b$$
,

can be graphically linearized by plotting $\ln y$ and $\ln x$ on the coordinate axes (\rightarrow D3) because

 $\ln y = \ln a + b \cdot \ln x$,

where b is the slope and ln a is the intercept.

Note: The condition x or y=0 does not exist on logarithmic coordinates because $\ln 0=\infty$. Nevertheless, $\ln a$ is still called the intercept in the equation when the logarithmic abscissa (\rightarrow **D3.4**) is intercepted by the ordinate at $\ln x=0$, i.e., x=1.

Instead of plotting In x and/or In y on the x- and/or y-axis, they can be plotted on **logarithmic paper** on which the ordinate or abscissa (semi-log paper) or both coordinates (log-log paper) are plotted in logarithmic units. In such cases, a is no longer treated as the intersect because the position of a depends on site of intersection of the x-axis by the y-axis. All values > 0 are possible.

Other nonlinear functions can also be graphically linearized using an appropriate plotting method. Take, for example, the **Michaelis-Menten equation** (\rightarrow **E1**), which applies to

many enzyme reactions and carrier-mediated transport processes:

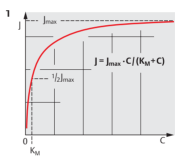
$$J = J_{\text{max}} \cdot \frac{C}{K_{\text{M}} + C}$$
 [13.10]

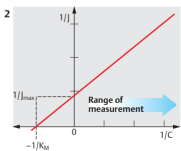
where J is the actual rate of transport (e.g., in $mol \cdot m^{-2} \cdot s^{-1}$), J_{max} is the maximal transport rate, $C (mol \cdot m^{-3})$ is the actual concentration of the substance to be transported, and K_M is the concentration (half-saturation concentration) at $^{1}/_{2} I_{max}$.

One of the three commonly used linear rearrangements of the Michaelis–Menten equation, the *Lineweaver–Burk plot*, states:

$$1/J = (K_M/J_{max}) \cdot (1/C) + 1/J_{max},$$
 [13.11]

Consequently, a plot of 1/J on the y-axis and 1/C on the x-axis results in a straight line $(\rightarrow E2)$. While a plot of J over J ove





E. Two methods of representing the Michaelis–Menten equation: The data can be plotted as a curve of J over C (E1), or as 1/J over 1/C in linearized form (E2). In the latter case, Jmax and K_M are determined by extrapolating the data outside the range of measurement.

permit accurate extrapolation of J_{max} (because an infinitely high concentration of C would be required), the linear rearrangement (\rightarrow E2) makes it possible to generate a regression line that can be extrapolated to $C = \infty$ from the measured data. Since 1/C is equal to $1/\infty = 0.1/J_{max}$ lies on the y-axis at x = 0 (\rightarrow E2). The reciprocal of this value is J_{max} . Insertion of 1/J = 0 into equation 13.11 yields

$$0 = (K_{M}/J_{max}) \cdot (1/C) + 1/J_{max}$$
 [13.12]

or $1/K_M = -1/C$, so that K_M can be derived from the negative reciprocal of the x-axis intersect, which corresponds to $1/J = 0 (\rightarrow E2)$.

The Greek Alphabet

α	A	alpha
β	В	beta
γ	Γ	gamma
δ	Δ	delta
ε	E	epsilon
ζ	Z	zeta
η	H	eta
ϑ, θ	Θ	theta
ι	I	iota
κ	K	kappa
λ	Λ	lamda
μ	M	mu
ν	N	nu
ξ	Ξ	xi
O	O	omicron
π	П	pi
ρ	P	rho
σ, ς	Σ	sigma
τ	T	tau
υ	Y	upsilon
φ	Φ	phi
χ	X	chi
ψ	Ψ	psi
ω	Ω	omega

Reference Values in Physiology

Total body and cells

Chemical composition of 1 kg fat-free body mass of an adult

Distribution of water in adult (child) as percentage of body weight (cf. p. 168)

Ion concentrations in ICF and ECF

720 g water, 210 g protein, 22.4 g Ca, 12 g P, 2.7 g K, 1.8 g Na, 1.8 g Cl, 0.47 g Mg

Intracellular: 40% (40%); interstitium: 15%

(25%); plasma: 5% (5%)

See p. 93 C

Cardiovascular system

Weight of heart

Cardiac output at rest (maximal)

Resting pulse = sinus rhythm

AV rhythm

Ventricular rhythm

Arterial blood pressure (Riva-Rocci)

Pulmonary artery pressure

Central venous pressure
Portal venous pressure

Ventricular volume at end of diastole/systole

Ejection fraction

384

Pressure pulse wave velocity

Mean velocity of blood flow

250-350 g

5-6 L/min (25 L/min); cf. p. 186

60-75 min-1 or bpm

40-55 min⁻¹

25-40 min-1

120/80 mm Hg (16/10.7 kPa) systolic/diastolic 20/7 mm Hg (2.7/0.9 kPa) systolic/diastolic

3–6 mm Hg (0.4–0.8 kPa)

3-6 mm Hg (0.4-0.8 kPa)

120 mL/40 mL

0.67

Aorta: 3-5 m/s; arteries: 5-10 m/s;

veins: 1-2 m/s

Aorta: 0.18 m/s; capillaries: 0.0002-0.001 m/s;

venae cavae: 0.06 m/s

Blood flow in organs at rest		
(See also pp. 187 A, 213 A)	% of cardiac output	per gram of tissue
Heart	4%	0.8 mL/min
Brain	13%	0.5 mL/min
Kidneys	20%	4 mL/min
GI tract (drained by portal venous system)	16%	0.7 mL/min
Liver (blood supplied by hepatic artery)	8%	0.3 mL/min
Skeletal muscle	21%	0.04 mL/min
Skin and miscellaneous organs	18%	_
Lungs and gas transport	Men	Women
Total lung capacity (TLC)	7 L	6.2 L
Vital capacity (VC); cf. p. 112	5.6 L	5 L
Tidal volume (V_T) at rest	0.6 L	0.5 L
Inspiratory reserve volume	3.2 L	2.9 L
Expiratory reserve volume	1.8 L	1.6 L
Residual volume	1.4 L	1,2 L
Max. breathing capacity in 30 breaths/min	110 L	100 L
Partial pressure of O ₂	Air: 21.17 kPa	(159 mm Hg)
•	Alveolar: 13.33 kPa	(100 mm Hg)
	Arterial: 12.66 kPa	(95 mm Hg)
	Venous 5.33 kPa	(40 mm Hg)
Partial pressure of CO ₂	Air: 0.03 kPa	(0.23 mm Hg)
	Alveolar: 5.2 kPa	(39 mm Hg)
	Arterial: 5.3 kPa	(40 mm Hg)
	Venous: 6.1 kPa	(46 mm Hg)
Respiratory rate (at rest)	16 breaths/min	
Dead space volume	150 mL	
Oxygen capacity of blood	180-200 mL O ₂ /L blo	od = $8-9 \text{ mmol } O_2/L \text{ blood}$
Respiratory quotient	0.84 (0.7–1.0)	
Kidney and excretion		
Renal plasma flow (RPF)	480-800 mL/min per	1.73 m ² body surface
	area	
Glomerular filtration rate (GFR)	80-140 mL/min per 1	1.73 m ² body surface area
Filtration fraction (GFR/RPF)	0.19	
Urinary output	0.7-1.8 L/day	
Osmolality of urine	250–1000 mOsm/kg	H_2O
Na ⁺ excretion	50–250 mmol/day	
K ⁺ excretion	25-115 mmol/day	
Glucose excretion	< 300 mg/day = 1.67	mmol/day
Nitrogen excretion	150-250 mg/kg/day	
Protein excretion	10-200 mg/day	
Urine pH	4.5-8.2	
Titratable acidity	10-30 mmol/day	
Urea excretion	10-20 g/day = 166-33	33 mmol/day
TT. Co. and Co	200 000/- 4	70 6 52 1/ 1

300-800 mg/day = 1.78-6.53 mmol/day

 $0.56-2.1 \text{ g/day} = 4.95-18.6 \mu\text{g/day}$

Uric acid excretion

Creatinine excretion

Energy expenditure during various activities • Bed rest • Light office work • Walking (4.9 km/h) • Sports (dancing, horseback riding, swimming) Functional protein minimum Vitamins, optimal daily intake (IU = international units) Electrolytes and trace elements, optimal daily intake For the state of the state	Men 6500 kJ/d (1550 kcal/d) 10 800 kJ/d (2600 kcal/d) 3.3 kW 4.5–6.8 kW 1 g/kg body weight A: 10 000–50 000 IU; I E: 200–800 IU; K: 65–25–300 mg of each; B₁ Ca: 1–1.5 g; Cr: 200–66 Fe: 15–30 mg; I: 50–30 Mg: 500–750 mg; Mn: Mo: 45–500 μg; Na ⁺ : 2 Se: 50–400 μg; Zn: 22-	80 μg; B ₁ , B ₂ , B ₅ , B ₆ : 1 ₂ : 25–300 μg; folate: 0 μg; C: 500–5000 mg 00 μg; Cu: 0.5–2 mg; 00 μg; K*: 0.8–1.5 g; 15–30 mg;
Bed rest Light office work Walking (4.9 km/h) Sports (dancing, horseback riding, swimming) Functional protein minimum Vitamins, optimal daily intake (IU = international units) Electrolytes and trace elements, optimal daily intake Nervous Systems, muscles Duration of an action potential	(1550 kcal/d) 10 800 kJ/d (2600 kcal/d) 3.3 kW 4.5–6.8 kW 1 g/kg body weight A: 10 000–50 000 IU; I E: 200–800 IU; K: 65–1 25–300 mg of each; B ₁ 0.4–1.2 mg; H: 25–30C Ca: 1–1.5 g; Cr: 200–61 Fe: 15–30 mg; I: 50–30 Mg: 500–750 mg; Mn: Mo: 45–500 µg; Na*: 2	(1300 kcal/d) 9600 kJ/d (2300 kcal/d) 2.7 kW 3.6–5.4 kW D: 400–600 IU 80 µg; B ₁ , B ₂ , B ₅ , B ₆ : 12: 25–300 µg; folate: 0 µg; C: 500–5000 mg 00 µg; Cu: 0.5–2 mg; 00 µg; K*: 0.8–1.5 g; 15–30 mg;
Light office work Walking (4.9 km/h) Sports (dancing, horseback riding, swimming) Functional protein minimum Vitamins, optimal daily intake (IU = international units) Electrolytes and trace elements, optimal daily intake M N N Nervous Systems, muscles Duration of an action potential	(1550 kcal/d) 10 800 kJ/d (2600 kcal/d) 3.3 kW 4.5–6.8 kW 1 g/kg body weight A: 10 000–50 000 IU; I E: 200–800 IU; K: 65–1 25–300 mg of each; B ₁ 0.4–1.2 mg; H: 25–30C Ca: 1–1.5 g; Cr: 200–61 Fe: 15–30 mg; I: 50–30 Mg: 500–750 mg; Mn: Mo: 45–500 µg; Na*: 2	(1300 kcal/d) 9600 kJ/d (2300 kcal/d) 2.7 kW 3.6–5.4 kW D: 400–600 IU 80 µg; B ₁ , B ₂ , B ₅ , B ₆ : 12: 25–300 µg; folate: 0 µg; C: 500–5000 mg 00 µg; Cu: 0.5–2 mg; 00 µg; K*: 0.8–1.5 g; 15–30 mg;
Light office work Walking (4.9 km/h) Sports (dancing, horseback riding, swimming) Functional protein minimum Vitamins, optimal daily intake (IU = international units) Electrolytes and trace elements, optimal daily intake Nervous Systems, muscles Duration of an action potential	10 800 kJ/d (2600 kcal/d) 3.3 kW 4.5–6.8 kW 1 g/kg body weight A: 10 000–50 000 IU; I E: 200–800 IU; K: 65–i 25–300 mg of each; B ₁ 0.4–1.2 mg; H: 25–30C Ca: 1–1.5 g; Cr: 200–66 Fe: 15–30 mg; I: 50–3C Mg: 500–750 mg; Mn: Mo: 45–500 μg; Na*: 2	9600 kJ/d (2300 kcal/d) 2.7 kW 3.6–5.4 kW D: 400–600 IU 80 μg; B ₁ , B ₂ , B ₅ , B ₆ : 12: 25–300 μg; folate: 0 μg; C: 500–5000 mg 00 μg; Cu: 0.5–2 mg; 00 μg; K*: 0.8–1.5 g; 15–30 mg;
Walking (4.9 km/h) Sports (dancing, horseback riding, swimming) Functional protein minimum Vitamins, optimal daily intake (IU = international units) Electrolytes and trace elements, optimal daily intake For the state of the stat	(2600 kcal/d) 3.3 kW 4.5–6.8 kW 1 g/kg body weight A: 10 000–50 000 IU; I E: 200–800 IU; K: 65–i 25–300 mg of each; B ₁ 0.4–1.2 mg; H: 25–300 Ca: 1–1.5 g; Cr: 200–66 Fe: 15–30 mg; I: 50–30 Mg: 500–750 mg; Mn: Mo: 45–500 µg; Na*: 2	(2300 kcal/d) 2.7 kW 3.6–5.4 kW D: 400–600 IU 80 µg; B ₁ , B ₂ , B ₅ , B ₆ : 12: 25–300 µg; folate: 0 µg; C: 500–5000 mg 00 µg; Cu: 0.5–2 mg; 00 µg; K ⁺ : 0.8–1.5 g; 15–30 mg;
Walking (4.9 km/h) Sports (dancing, horseback riding, swimming) Functional protein minimum Vitamins, optimal daily intake (IU = international units) Electrolytes and trace elements, optimal daily intake Nervous Systems, muscles Duration of an action potential	3.3 kW 4.5–6.8 kW 1 g/kg body weight A: 10 000–50 000 IU; I E: 200–800 IU; K: 65–: 25–300 mg of each; B ₁ 0.4–1.2 mg; H: 25–300 Ca: 1–1.5 g; Cr: 200–66 Fe: 15–30 mg; I: 50–30 Mg: 500–750 mg; Mn: Mo: 45–500 µg; Na ⁺ : 2	2.7 kW 3.6–5.4 kW D: 400–600 IU 80 μg; B ₁ , B ₂ , B ₅ , B ₆ : 12: 25–300 μg; folate: 0 μg; C: 500–5000 mg 00 μg; Cu: 0.5–2 mg; 15–30 mg;
Sports (dancing, horseback riding, swimming) Functional protein minimum Vitamins, optimal daily intake (IU = international units) Electrolytes and trace elements, optimal daily intake Nervous Systems, muscles Duration of an action potential	4.5–6.8 kW 1 g/kg body weight A: 10 000–50 000 IU; I E: 200–800 IU; K: 65–25–300 mg of each; B ₁ 0.4–1.2 mg; H: 25–300 Ca: 1–1.5 g; Cr: 200–66 Fe: 15–30 mg; I: 50–30 Mg: 500–750 mg; Mn: Mo: 45–500 μg; Na*: 2	3.6–5.4 kW D: 400–600 IU 80 μg; B ₁ , B ₂ , B ₅ , B ₆ : 12: 25–300 μg; folate: 0 μg; C: 500–5000 mg 00 μg; Cu: 0.5–2 mg; 00 μg; K*: 0.8–1.5 g; 15–30 mg;
Functional protein minimum Vitamins, optimal daily intake (IU = international units) Electrolytes and trace elements, optimal daily intake Nervous Systems, muscles Duration of an action potential	1 g/kg body weight A: 10 000–50 000 IU; I E: 200–800 IU; K: 65–25–300 mg of each; B ₁ 0.4–1.2 mg; H: 25–300 Ca: 1–1.5 g; Cr: 200–60 Fe: 15–30 mg; I: 50–30 Mg: 500–750 mg; Mn: Mo: 45–500 μg; Na ⁺ : 2	D: 400–600 IU 80 μg; B ₁ , B ₂ , B ₅ , B ₆ : 12: 25–300 μg; folate: 1 μg; C: 500–5000 mg 00 μg; Cu: 0.5–2 mg; 100 μg; K*: 0.8–1.5 g; 15–30 mg;
Vitamins, optimal daily intake (IU = international units) Electrolytes and trace elements, optimal daily intake Nervous Systems, muscles Duration of an action potential	A: 10 000–50 000 IU; I E: 200–800 IU; K: 65– 25–300 mg of each; B ₁ 0.4–1.2 mg; H: 25–300 Ca: 1–1.5 g; Cr: 200–60 Fe: 15–30 mg; I: 50–30 Mg: 500–750 mg; Mn: Mo: 45–500 µg; Na ⁺ : 2	80 μg; B ₁ , B ₂ , B ₅ , B ₆ : 1 ₂ : 25–300 μg; folate: 0 μg; C: 500–5000 mg 00 μg; Cu: 0.5–2 mg; 00 μg; K*: 0.8–1.5 g; 15–30 mg;
(IU = international units) Electrolytes and trace elements, optimal daily intake Nervous Systems, muscles Duration of an action potential	E: 200–800 IU; K: 65– 25–300 mg of each; B ₁ 0.4–1.2 mg; H: 25–300 Ca: 1–1.5 g; Cr: 200–60 Fe: 15–30 mg; I: 50–30 Mg: 500–750 mg; Mn: Mo: 45–500 μg; Na ⁺ : 2	80 μg; B ₁ , B ₂ , B ₅ , B ₆ : 1 ₂ : 25–300 μg; folate: 0 μg; C: 500–5000 mg 00 μg; Cu: 0.5–2 mg; 00 μg; K*: 0.8–1.5 g; 15–30 mg;
Electrolytes and trace elements, optimal daily intake Poptimal daily intake Nervous Systems, muscles Duration of an action potential	25–300 mg of each; B ₁ 0.4–1.2 mg; H: 25–300 Ca: 1–1.5 g; Cr: 200–60 Fe: 15–30 mg: I: 50–30 Mg: 500–750 mg; Mn: Mo: 45–500 μg; Na ⁺ : 2	12: 25–300 μg; folate: 1 μg; C: 500–5000 mg 00 μg; Cu: 0.5–2 mg; 00 μg; K+: 0.8–1.5 g; 15–30 mg;
Electrolytes and trace elements, optimal daily intake Poptimal daily intake Nervous Systems, muscles Duration of an action potential	0.4–1.2 mg; H; 25–300 Ca: 1–1.5 g; Cr: 200–60 Fe: 15–30 mg; I: 50–30 Mg: 500–750 mg; Mn: Mo: 45–500 µg; Na*: 2	0 μg; C: 500–5000 mg 00 μg; Cu: 0.5–2 mg; 00 μg; K ⁺ : 0.8–1.5 g; : 15–30 mg;
Electrolytes and trace elements, optimal daily intake Polymer of the second of the se	Ca: 1-1.5 g; Cr: 200-6 Fe: 15-30 mg: I: 50-30 Mg: 500-750 mg; Mn: Mo: 45-500 µg; Na†: 2	00 μg; Cu: 0.5–2 mg; 00 μg; K ⁺ : 0.8–1.5 g; : 15–30 mg;
optimal daily intake No. No. S. Nervous Systems, muscles Duration of an action potential	Fe: 15–30 mg: I: 50–30 Mg: 500–750 mg; Mn: Mo: 45–500 µg; Na+: 2	00 μg; K ⁺ : 0.8–1.5 g; : 15–30 mg;
Nervous Systems, muscles Duration of an action potential	Mg: 500-750 mg; Mn: Mo: 45-500 μg; Na ⁺ : 2	: 15–30 mg;
Nervous Systems, muscles Duration of an action potential	Mo: 45–500 μg; Na ⁺ : 2	
Nervous Systems, muscles Duration of an action potential		, g, i . 200 400 mg,
Nervous Systems, muscles Duration of an action potential		
Duration of an action potential		
•	Nerve: 1–2 ms; skeleta	al muscle: 10 ms:
	myocardium: 200 ms	ii iiiuscie. 10 iiis,
	See p. 49 C	
Blood and other bodily fluids ((see also Tables 9.1, and	l 9.2 on pp. 186, 187)
Blood (in adults)	Men:	Women:
((p)	4500 mL	3600 mL
	0.40-0.54	0.37-0.47
` ,	4.5-5.9 · 10 ¹² /L	$4.2-5.4\cdot 10^{12}/L$
	140-180 g/L	120-160 g/L
•	(2.2–2.8 mmol/L)	(1.9–2.5 mmol/L)
1 ,	80–100 fL	
• , ,	320–360 g/L	
9 , ,		
		180_400 . 10 ⁹ /I
	,	,
j ()		- 20 mm m mst nou
	66-85 g/L serum	
Total		55-64% of total
		2.5–4% of total
Albumin	1.3–4 g/L serum	
Albumin α_1 -globulins	1.3–4 g/L serum 4–9 g/L serum	7-10% of total
Albumin α_1 -globulins α_2 -globulins α_3 -globulins		7–10% of total 8–12% of total
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	4–9 g/L serum	
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	4–9 g/L serum 6–11 g/L serum 13–17 g/L serum	8–12% of total 12–20% of total
Albumin α_1 -globulins α_2 -globulins β	4–9 g/L serum 6–11 g/L serum 13–17 g/L serum (See p. 102 for coagula	8–12% of total 12–20% of total tion factors)
Albumin $\alpha_1\text{-globulins}$ $\alpha_2\text{-globulins}$ $\beta\text{-globulins}$ $\gamma\text{-globulins}$ α_3 $\gamma\text{-globulins}$ α_4 $\gamma\text{-globulins}$ α_5 α_7	4–9 g/L serum 6–11 g/L serum 13–17 g/L serum	8–12% of total 12–20% of total tion factors)
Mean Hb concentration in single RBC (MCH) Mean RBC diameter Reticulocytes Leukocytes (also refer to table on p. 88) Platelets Erythrocyte sedimentation rate (ESR) Proteins	27–32 pg 7.2–7.8 μm 0.4–2% (20–75 · 10 ⁹ /L) 3–11 · 10 ⁹ /L 170–360 · 10 ⁹ /L < 10 mm in first hour 66–85 g/L serum 35–50 g/L serum	180–400 · 10 ⁹ /L < 20 mm in first hou 55–64% of total

Parameters of glucose metabolism Glucose concentration in venous blood Glucose concentration in capillary blood Glucose concentration in plasma Limit for diabetes mellitus in plasma HBA _{1c} (glycosylated hemoglobin A) Parameters of lipid metabolism	3.9–5.5 mmol/L 4.4–6.1 mmol/L 4.2–6.4 mmol/L > 7.8 mmol/L 3.2–5.2%	(70–100 mg/dL) (80–110 mg/dL) (75–115 mg/dL) (> 140 mg/dL)
Triglycerides in serum Total cholesterol in serum HDL cholesterol in serum	< 1.71 mmol/L < 5.2 mmol/L > 1.04 mmol/L	(< 150 mg/dL) (< 200 mg/dL) (> 40 mg/dL)
Substances excreted in urine Urea concentration in serum Uric acid concentration in serum Creatinine concentration in serum Bilirubin	3.3–8.3 mmol/L 150–390 μmol/L 36–106 μmol/L	(20-50 mg/dL) (2.6-6.5 mg/dL) (0.4-1.2 mg/dL)
Total bilirubin in serum Direct bilirubin in serum Electrolytes and blood gases	3.4–17 μmol/L 0.8–5.1 μmol/L	(0.2–1 mg/dL) (0.05–0.3 mg/dL)
Osmolality Cations in serum:	280–300 mmol/kg H Na ⁺ K ⁺ Ionized Ca ²⁺ Ionized Mg ²⁺	₂ O 135–145 mmol/L 3.5–5.5 mmol/L 1.0–1.3 mmol/L 0.5–0.7 mmol/L
Anions in serum:	Cl ⁻ H ₂ PO ₄ ⁻ + HPO ₄ ²⁻	95–108 mmol/L 0.8–1.5 mmol/L
pH Standard bicarbonate TotaL buffer bases Oxygen saturation		7.35–7.45 22–26 mmol/L 48 mmol/L 96% arterial; 65–75% mixed venous
Partial pressure of O_2 at half saturation ($P_{0.5}$)		3.6 kPa (27 mmHg)
Cerebrospinal fluid Pressure in relaxed horizontal position Specific weight Osmolality Glucose concentration	1.4 kPa (10.5 mmHg) 1.006–1.008 g/L 290 mOsm/kg H ₂ O 45–70 mg/dL (2.5–3.5	9 mmol/L)

0.15-0.45 g/L

< 84 mg/dL

< 5 WBC/ μ L

Protein concentration

IgG concentration White cell count

Important Equations in Physiology

1. Fick's law of diffusion for membrane transport (see also p. 20ff.)

$$\mathbf{J}_{diff} = \mathbf{F} \cdot \mathbf{D} \cdot \frac{\Delta \mathbf{C}}{\Delta \mathbf{x}} [\text{mol} \cdot \text{s}^{-1}]$$

 J_{diff} = net diffusion rate [mol·s⁻¹];

 $A = area [m^2];$

D = diffusion coefficient $[m^2 \cdot s^{-1}]$;

 ΔC = concentration difference [mol·m⁻³];

 $\Delta x = membrane thickness [m]$

Alternative 1:

$$\frac{\mathbf{J}_{\text{diff}}}{\mathbf{F}} = \mathbf{P} \cdot \Delta \mathbf{C} \left[\text{mol} \cdot \mathbf{m}^{-2} \cdot \mathbf{S}^{-1} \right]$$

P = permeability coefficient [$m \cdot s^{-1}$]; I_{diff}, A an ΔC ; see above

Alternative 2 (for gas diffusion)

$$\frac{\dot{\mathbf{V}}_{\text{diff}}}{\mathbf{F}} = \mathbf{K} \cdot \frac{\Delta \mathbf{P}}{\Delta \mathbf{x}} [\mathbf{m} \cdot \mathbf{s}^{-1}]$$

 \dot{V}_{diff} = net diffusion rate [m³ · s⁻¹];

K = Krogh's diffusion coefficient $[m^2 \cdot s^{-1} \cdot Pa^{-1}]$ ΔP = partial pressure difference [Pa]

2. Van't Hoff–Stavermann equation

(see also p. 377)

$$\Delta \pi = \sigma \cdot \mathbf{R} \cdot \mathbf{T} \cdot \Delta \mathbf{c}_{\mathsf{osm}} [Pa]$$

 $\Delta\pi$ = osmotic pressure difference [Pa] σ = reflection coefficient [dimensionsless] R = universal gas constant [8.3144] · K⁻¹ ·

mol⁻¹]; T = absolute temperature [K];

 Δc_{osm} = concentration difference of osmotically active particles [mol·m⁻³].

3. Michaelis-Menten equation

(see also pp. 28, 383ff.)

$$\mathbf{J}_{\text{sat}} = \mathbf{J}_{\text{max}} \cdot \frac{\mathbf{C}}{\mathbf{K}_{\text{M}} + \mathbf{C}} [\text{mol} \cdot \text{m}^{-2} \cdot \text{S}^{-1}],$$

 J_{sat} = substrate transport (turnover)

[mol·m $^{-2}$ ·s $^{-1}$];

 J_{max} = maximum substrate transport (turnover) [mol·m⁻²·s⁻¹];

C = substrate concentration [mol \cdot m⁻³];

 K_M = Michaelis constant = substrate concentration at $^1/_2$ J_{max} [mol·m⁻³].

4. Nernst equation (see also p. 32)

$$E_x = -61 \cdot z_x^{-1} \cdot \log \frac{[X]_i}{[X]_a} [mV]$$

 E_x = equilibrium potential of ion X [mV];

 z_x = valency of ion X;

 $[X]_i$ = intracellular concentration of ion X

[mol·m⁻³]

 $[X_o]$ = extracellular concentration of ion X $[mol \cdot m^{-3}]$.

5. Ohm's Law (see also pp. 32, 188)

a. For ion transport at membrane

$$I_x = g_x \cdot (E_m - E_x) [A \cdot m^{-2}]$$

 I_x = ionic current of ion X per unit area of membrane [A · m⁻²];

 g_x = conductance of membrane to ion X [S · m⁻²]:

 E_m = membrane potential [V];

 E_x = equilibrium potential of ion X[V]

b. For blood flow:

$$\dot{\mathbf{Q}} = \frac{\Delta \mathbf{P}}{\mathbf{R}} \left[\mathbf{L} \cdot \mathbf{min}^{-1} \right]$$

Q = flow rate (total circulation:

cardiac output, CO) [L \cdot min $^{-1}$]

 ΔP = mean blood pressure difference (systemic circulation: $\overline{P}_{aorta} - \overline{P}_{vena cava}$;

lesser circulation: $P_{pulmonary\ artery} - P_{pulmonary\ vein}$

R = flow resistance (systemic circula-

total peripheral resistance = TPR) [mmHg \cdot min \cdot L⁻¹].

6. Respiration-related equations

(see also pp. 106, 120)

a. Tidal volume (V_T) : $V_T = V_D + V_A [L]$

b. Respiratory volume per minute $(\dot{V}_E \text{ oder } \dot{V}_T)$:

$$\dot{V}_E = f \cdot V_T = (f \cdot V_D) + (f \cdot V_A) =$$

 $\dot{\mathbf{V}}_{\mathbf{D}} + \dot{\mathbf{V}}_{\mathbf{A}} \left[\mathbf{L} \cdot \min^{-1} \right]$

c. O₂ consumption, CO₂ emission, and RO (total body:)

$$\dot{\mathbf{V}}_{\mathbf{O}_2} = \dot{\mathbf{V}}_{\mathsf{T}} \left(\mathbf{F}_{\mathbf{I}\mathbf{O}_2} - \mathbf{F}_{\mathbf{E}\mathbf{O}_2} \right) = \mathbf{CO} \cdot \mathbf{a} \mathbf{v} \mathbf{D}_{\mathbf{O}_2} \left[\mathbf{L} \cdot \mathbf{min}^{-1} \right]$$

 $\dot{\mathbf{V}}_{\mathbf{CO}_2} = \dot{\mathbf{V}}_{\mathsf{T}} \cdot \mathbf{F}_{\mathbf{E}\mathbf{CO}_2} \left[\mathbf{L} \cdot \mathbf{min}^{-1} \right]$

$$RQ = \frac{\dot{V}_{CO_2}}{\dot{V}_{O_2}}$$

 V_D = dead space [L];

 V_A = alveolar fraction of $V_T[L]$;

 $f = respiration rate [min^{-1}];$

 \dot{V}_D = dead space ventilation [L·min⁻¹];

 \dot{V}_A = alveolar ventilation [L · min⁻¹];

 $\dot{V}_{0_2} = O_2$ consumption [L·min⁻¹];

 \dot{V}_{CO_2} = CO_2 emission [L·min⁻¹];

 F_{1O_2} = inspiratory O_2 fraction [L/L];

 F_{EO_2} = exspiratory O_2 fraction [L/L];

 F_{ECO_2} = exspiratory CO_2 fraction [L/L];

RO = respiratory quotient (dimensionless)

d. O_2 consumption and CO_2 emission (organ):

 $\dot{\mathbf{V}}_{\mathbf{O}_2} = \dot{\mathbf{Q}} \cdot \mathbf{av} \mathbf{D}_{\mathbf{O}_2} \left[\mathbf{L} \cdot \min^{-1} \right]$

 $\dot{\mathbf{V}}_{CO_2} = \dot{\mathbf{O}} \cdot av \mathbf{D}_{CO_2} \left[\mathbf{L} \cdot min^{-1} \right]$

 \dot{Q} = blood flow in organ [L·min⁻¹]

 avD_{O_2} , avD_{CO_2} = arteriovenous

O2 and CO2 difference in total circulation and organ circulation [L/L blood]

e. Fick's principle:

$$\mathbf{CO} = \frac{\dot{\mathbf{V}}_{O_2}}{\text{av}\mathbf{D}_{O_2}} \left[\mathbf{L} \cdot \text{min}^{-1} \right]$$

CO = cardiac output $[L \cdot min^{-1}]$ >

f. Gas partial pressure \leftrightarrow gas concentration in liquids:

 $[X] = \alpha \cdot P_x [\text{mmol/L plasma}]$

[X] = concentration of gas X [mmol·L⁻¹]

 α = (Bunsen's) solubility coefficient $[mmol \cdot L^{-1} \cdot kPa^{-1}]$ P_X = partial pressure of gas X [kPa]

g. Bohr's formula (see also p. 115)

 $V_D = V_T \frac{(F_{ACO_2} - F_{ECO_2})}{F_{ACO_2}}$

 V_D = dead space [L]:

V_T = tidal volume [L];

FACO2 = alveolar CO2 fraction

FECO2 = exspiratory CO2 fraction [L/L]

h. Alveolar gas equation (see also p. 136)

$$P_{AO_2} = P_{IO_2} - \frac{P_{AO_2}}{RO} [kPa]$$

 P_{AO_2} and P_{IO_2} = alveolar and inspiratory partial pressure of O2 [kPa]

 P_{ACO_2} = alveolar partial pressure of CO_2 [kPa]

RQ = respiratory quotient [dimensionless].

Henderson-Hasselbalch equation

(see also pp. 138ff., 379)

a. General equation:

$$pH = pK_a + log \frac{[A^-]}{[AH]}$$

b. for bicarbonate/CO₂ buffer (37 °C):

pH = 6,1 +
$$\frac{[HCO_3^-]}{\alpha \cdot P_{CO_2}}$$

pH = negative common logarithm of H+ activity

 pK_a = negative common logarithm of dissociation constant of buffer acid in denominator (AH or CO2)

 $[A^-]$ and $[HCO_3^-]$ = buffer base concentration; $\alpha \cdot P_{CO_2} = [CO_2]$; see Equation 6f.

8. Equations for renal function

(see also p. 150ff.)

a. Clearance of a freely filtrable substance X

$$\mathbf{C}_{\mathbf{X}} = \dot{\mathbf{V}}_{\mathbf{U}} \cdot \frac{\mathbf{U}_{\mathbf{X}}}{\mathbf{P}_{\mathbf{X}}} \left[\mathbf{L} \cdot \min^{-1} \right]$$

b. Renal plasma flow

RPF =
$$\dot{\mathbf{V}}_{U} \cdot \frac{\mathbf{U}_{PAH}}{\mathbf{0.9 \cdot P}_{PAH}} [L \cdot min^{-1}]$$

c. Renal blood flow (RBF):

$$RBF = \frac{RPF}{1 - HCT} [L \cdot min^{-1}]$$

d. Glomerular filtration rate (GFR):

$$GFR = \dot{V}_{U} \cdot \frac{U_{ln}}{P_{ln}} [L \cdot min^{-1}]$$

e. Free water clearance (CH2O)

$$C_{H_2O} = \dot{V}_U \cdot (1 - \frac{U_{osm}}{P_{osm}}) [L \cdot min^{-1}]$$

f. Filtration fraction

$$FF = \frac{GFR}{RPF}$$
 [dimensionless]

g. Fractional excretion of substance X (FE_X): $FE_X = \frac{C_X}{GFR}$ [dimensionless];

$$FE_X = \frac{C_X}{GFR}$$
 [dimensionless];

h. Fractional reabsorption of substance X

 $FR_X = 1 - FE_X$ [dimensionslos];

 \dot{V}_U = urinary excretion rate [L·min⁻¹] U_X , U_{PAH} , U_{In} = urinary concentration of substance X, para-aminohippuric acid, and indicator (e.g., inulin, endogenous creatinine) $[\text{mol} \cdot L^{-1}]$ or $[\text{g} \cdot L^{-1}]$

 $\begin{array}{l} U_{osm} = osmolality \ of urine \ [Osm \cdot L^{-1}] \\ P_{Xx} \ P_{PAH}, \ P_{In} = plasma \ concentration \ of substance \ X, para-aminohippuric acid, and indicator (e.g., inulin, endogenous creatinine) \ [mol \cdot L^{-1}] \ or \ [g \cdot L^{-1}] \\ P_{osm} = osmolality \ of \ plasma \ [Osm \cdot L^{-1}] \\ HCT = hematocrit \ [L \ of \ blood \ cells/L \ of \ blood] \end{array}$

9. Equations for filtration

(see also pp. 152, 208)

a. Effective filtration pressure at capillaries (P_{eff})

$$P_{eff} = P_{cap} - P_{int} - \pi_{cap} + \pi_{int} [mmHg]$$

b. Effective filtration pressure at capillaries of renal glomerulus:

$$\textbf{P}_{\text{eff}} = \textbf{P}_{\text{cap}} - \textbf{P}_{\text{Bow}} - \pi_{\text{cap}} \left[mmHg \right]$$

c. Filtration rate (\dot{Q} at glomerulus = GFR) $\dot{Q} = P_{eff} \cdot F \cdot k [m^3 \cdot s^{-1}]$

 $P_{cap}(P_{int})$ = hydrostatic pressure in capillaries (interstitium) [mm Hg]

 $\pi_{cap}(\pi_{int})$ = oncotic pressure in capillaries (interstitium) [mm Hg]

P_{eff} = mean effective filtration pressure [mm Hg] A = area of filtration (m²)

k = permeability to water (hydraulic conductance) $[m^3 \cdot s^{-1} \cdot mm Hg^{-1}]$

10. Law of Laplace

(see also pp. 118, 188, and 210)

a. Elliptical hollow body (with radii r_1 and r_2)

$$P_{tm} = T \left(\frac{1}{r_1} + \frac{1}{r_2} \right) [Pa];$$

b. Elliptical hollow body, considering wall thickness:

$$P_{tm} = S \cdot w \left(\frac{1}{r_1} + \frac{1}{r_2} \right) [Pa];$$

c. Spherical hollow body $(r_1 = r_2 = r)$:

$$P_{tm} = 2 \frac{T}{r} [Pa] \text{ or } P_{tm} = 2 \frac{S \cdot w}{r} [Pa];$$

d. Cylindrical hollow body

$$(r_2 \rightarrow \infty$$
, therefore $1/r_2 = 0$):

$$P_{tm} = \frac{T}{r} [Pa] \text{ or } P_{tm} = \frac{S \cdot w}{r} [Pa]$$

P_{tm} = transmural pressure [Pa]

T = wall tension $[N \cdot m^{-1}]$

S = wall tension $[N \cdot m^{-2}]$

w = wall thickness [m]

11. Equations for cardiovascular function

(see also items 2, 5b, 6c, and 9 as well as p. 186ff.)

a. Cardiac output (CO):

$$CO = f \cdot SV [1 \cdot min^{-1}]$$

b. Hagen-Poiseuille equation

$$R = \frac{8 \cdot 1 \cdot \eta}{\pi \cdot r^4};$$

f = heart rate [min-1]

SV = stroke volume [L]

R = flow resistance in a tube $[Pa \cdot s \cdot m^{-3}]$ of known length [m] and inner radius [m]

 $\eta = \text{viscosity} [Pa \cdot s]$

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Index, Abbreviations

A	gastrointestinal tract 234	myosin interaction 68
- (-++	heart action 194	smooth muscle 70
a (atto-, submultiple of a unit) 393	motor end plate 56 NO-synthase activation 82	skeletal muscle 60 Action potential 42, 62
A (ampere), unit 375	pankreas 246	all-or-none depolarization
band, muscle 60	parietal cells 242	46
AA (amino acids) 240, 258, 182,	release 82	depolarization phase 46
184, 296	receptors → cholinoceptors	heart 59 A, 192, 194
Abdominal muscles 108	second messenger 55 F, 274,	overshoot 46
pressure 108, 238, 264	276	pacemaker, heart 192
reflex 320	synthesis 82	postsynaptic neurons 82
Aberration, spherical 346	Acetylcholinesterase 56, 82	propagation 48
AB0 system 100	synaptic cleft 82	purkinje fibers 200
Absolut threshold, eye 352	Acetyl coenzyme A 284	repolarization phase 46
smell 340	A chain, insulin 282	retina 354
ear 362	Achalasia 238	skeletal muscle 56, 59 A
Absorption, intestinal, of amino	Acetylsalicylic acid 104	smooth muscle 59 A
acids 258 carbohydrates 258	cyclooxygenase inhibition 269	Activating energy 40 Activation system, subcortical
electrolytes 262	Achromat 356	336
fat 252	Acid, fixed 174	Active immunization 94
iron 90	production 174	transport 26, 44
lipids 252	titratable 174, 178	Activin, FSH secretion 306
peptides 258	Acid-base balance 138, 142	Activity coefficient 376
vitamins 260	compensatory mechanisms	Acuity, visual 348
Abstract thinking 336	142	Acyl-CoA-cholesterol acyl trans-
ABP (androgen-binding protein)	disturbances 142, 146, 176	ferase (ACAT) 256
306	kidney 174	Adam-Stokes attack 200
ACAT (acyl-CoA-cholesterol acyl	liver 176	Adaptation, eye 350, 352, 354,
transferase) 256	metabolic disturbances 142	sensors 312
Acceleration (unit) 374	normal parameters 142	smell 340
angular (rotational) 342	regulation 142	taste 338 thermosensation 314
detection 342 linear (translational) 342	respiratory disturbances 144 status, assessment 146	ADCC (antigen-dependent cell-
sensor 314, 342	Acidosis 142	mediated cytotoxic-
Accessory nucleus 368	hyperkalemic 180	ity) 96
Accident, electrical 200	influence on protein-bound	Addison's disease 182
Acclimatization 224	Ca ²⁺ 290	Adenin 8
Accommodation, eye 342, 346	K+ concentration, plasma 180	Adenohypophysis 268, 280
range 342, 346	lact- 76	Adenosine, coronary vasodilata-
reflex, intestine 244	metabolic 142, 262, 284	tion 210
rectum 264	nonrespiratory (metabolic)	second messenger of 274
stomach 240	142, 262, 284	Adenosine diphosphate → ADP
ACE (angiotensin-converting	in diarrhea 262	monophosphat → AMP
enzyme) 184 inhibitor 184	phosphaturia 178	triphosphat → ATP
A cells, islets of Langerhans 282,	renal tubular 142, 176 respiratory 126, 142, 144	Adenylate cyclase 37 C1, 274, 276 inhibition with acetylcholine
284	Acini, salivary gland 236	82
Acenocumarol 104	Acne 306	ADH → adiuretin
Acetate, conjugates 160	Aconitase, iron absorption 90	Adhesion molecule (VLA-4) 98
Acetazolamide 172	Acoustic information, central	Adhesion, platelets 102
Aceton 284	processing 368	Adiadochokinesis 328
Acetylcholine 34, 52, 78 ff., 236,	pathways 368	Adiuretin 24, 162, 170, 269, 280
242	thresholds 362, 368	activated Cl ⁻ -channels 162
antagonists 82	Acrosomal reaction 308	deficiency 166
cerebral cortex 332	ACTH → corticotropin	effects 212, 280
control of circulation 212 ff.	ACTH reserve 294	receptor types 24, 55 F, 166,
coronary vasodilatation 210	Actin 14, 30, 56, 58, 62, 70, 82	280
esophagus 238	filament 14, 60	salt/water homeostasis 173 B
esterase 56, 82	molecular structure 60	second messenger 24, 55 F

secretion 170, 218	sound conduction 364	cell types 118
ADP 72	Air space, voice 370	contact time 120
role in autoregulation 212	Airway resistance 134	diffusion distance 120
ADP-ribosylation factor (ARF) 30	A-kinase (proteinkinase A, PKA)	surface 118
Adrenal cortex 254, 272, 294,	84, 274	tension 118
303, 306	Akinesia 326	Amacrine cells, retina 344, 354
androgens 306	Alanin, glucagon secretion 284	Ambient temperature pressure
fetal 304	Alarm reaction 86, 330	H ₂ O-saturated (ATPS)
glucocorticoids 296	Albumin 92, 154, 158, 268, 306	112
HDL receptors 254	bilirubin binding 250	Amenorrhea 230
hormone synthesis 294 ff.,	calcium binding 290	Amiloride 172
304	pressure, oncotic 208	L-Amino acid decarboxylase 84
progesterone 303	renal filtration 154	Aminoaciduria 258
zones 294, 296	reabsorption 158	Amino acids (AA) 240, 258, 182,
medulla 78, 86 , 268, 272, 296	testosterone binding 306	184, 296
hormones 86	T ₃ /T ₄ transport 288	co-transport with Na ⁺ 158,
Adrenalin → epinephrine	Alcohol, energy source 226	258
Adrenergic transmission 84	Aldosteron 162, 170, 182 ff. , 184,	essential 226
	216	
Adrenoceptors 84, 212, 214		glucagon secretion 284
α- 84, 212 f.	antagonists 172	gluconeogenesis from 282
α ₁ - 80 ff., 214, 276, 304	deficiency 182	influence on insulin release
renin secretion 184	degradation 183 D	282 ff.
α ₂ - 52, 80 ff., 86, 146, 230, 248	effects 182	intestinal absorption 258,
insulin secretion 282	induced protein (AIP) 182	262
second messenger 55, 274	K+ metabolism 180	metabolism, cortisol effect
agonists 87 B	Na+-transport, intestinal 262	296
antagonists 87 B	tubular 180 ff.	pylorus, effect on 240
β- 86, 212 f.	receptor 172	renal transport 156, 158
β ₁ -, renin secretion 184	glucocorticoids 296	storage 284
second messenger 85 ff., 274	secretion 182, 183 D	transmitter 55 F
β ₂ -, 85 ff., 212, 284	synthesis 294	γ-aminobutyric acid (GABA) 52,
coronary vasodilatation 210	Alkaline phosphatase 250	320
second messenger 55, 85 ff.,	Alkalosis 142	p-aminohippurate 150, 160
274	Ca ²⁺ concentration, serum 290	Aminopeptidases, small intestine
salivary glands 236	hypokalemic 172, 180	258
β_3 -, fatty tissue 84, 222	K ⁺ plasma concentration 180	Ammonia \rightarrow NH ₃
second messenger 55, 274	metabolic 142	Ammonium \rightarrow NH ₄ ⁺
stomach 242	nonrespiratory 142	Amnesia 336
distribution 81, 87 B	phosphate reabsorption, renal	Amount of substance, unit 375
G- proteins 55, 87 B	178	5'-AMP 276
kidney 214	respiratory 142, 144	AMP, cyclic 84, 236, 274, 286
localisation 81 ff.	high altitude 136	adrenergic transmission 84
receptor types 84 ff.	vomiting 238	TSH 286
skin 214	Allergens 98, 100	AMPA receptors → glutamate
types 84, 87 B	Allergy 100	Ampere (A), unit 375
uterus 304	anaphylactic shock 218	Amplification, cochlear 366
Adrenocortical hormones 294	delayed type hypersensitivity	Ampulla, labyrinth 342
insufficiency 182	100	Amygdala 310, 320, 330, 336,
tumor 216	immediate reaction 100	340
Adrenocorticotropic hormone →	Allodynia 318	α-amylase 246, 258
corticotropin	All-or-none-response 46, 66, 192	Amylopectin 258
Aerobic glucose metabolism 72	all-trans-retinal 348, 350	Amylose 258
Affinity, transport system 28	Alternating current 50, 375	Anabolic, action 280
Afterload, heart 204	Altitude gain 136	insulin 284
maxima, curve 203	Alveolar gas 114	STH 280
Afterloaded contraction 66	partial pressure 120	testosterone 306
heart 202	equation 120, 136, 389	Anaerobic glycolysis 72
Agglutination, red blood cells 100	exchange 120, 124	threshold 72
Aggregation, platelets 102	pressure (P _A) 108, 116	Analgesic action 318
AIP (aldosterone-induced pro-	Alveolar ventilation 106, 114,	Anal sphincters 264
teins) 182	120 ff.	Anaphylaxis 100
Air. composition 107 A. 385	Alveoli 106, 118 ff.	Anastomoses, arteriovenous 224

Androgen-binding protein 306	Antihemophilic factors 102	interlobular 150
Androgens 300, 306	Antiovulatory effect 303	pressure 206
adrenal cortex 294	α_2 -Antiplasmin 104	umbilical 220
synthesis 294, 306	Antiport, definition 26	Arteriole 186 ff.
follicle 300	Antipyrine 168	afferent 184
Androstenedione 300	Antithrombin 104	efferent 184
Anemia 226, 260	Antrum, stomach 240	Arteriosclerosis 208, 210, 302
hyperchromic 90	α ₁ -Antitrypsin 104	coronary artery 210
hypochromic, iron deficiency	Anulospiral endings 316	hypertension 216
90	Anuria 164	Artery (→ also arteries), carotid,
pernicious 260 sickle cell 92	Anus 264 Aorta 188	pressure sensors 214 interlobular 150
Anergy 98	blood flow velocity 190	pulmonary 122
ANF → Atriopeptin	flow rate 191 A	pressure 122
Angina pectoris 318	pressure 190, 204	fetus 221 B
ECG 198	influence on heart functions	Ascending reticular activating
Angiotensin I 184	204	system (ARAS) 322,
lung 106	sensors 214	332
Angiotensin II (AT II) 158, 170,	valves 190	Ascites 208
182, 184, 212	APC (antigen-presenting cells)	Ascorbate → vitamin C
aldosterone secretion 182	96, 97 B, 98	Aspartate 174, 258
effects 184, 212, 216	Aphasia 370	intestinal absorption 258
CSF 168	Apnea 108, 132	receptor types 55 F
degradation, renal 158	Apolipoproteins 92, 254	second messenger 55 F
receptors 184	Apomorphine 238	Aspirin® 104, 268, 318
second messenger 274	Apoptosis 98, 272, 300	Associative cortex 324, 329, 336
Angiotensin-converting enzyme	Apotransferrin 90, 92	Asthma 100, 118
(ACE) 184	Apparatus, juxtaglomerular 172,	Astigmatism 346
Angiotensinogen 184	184	Astrocytes 338
synthesis, cortisol effect 296	Aprotinin 105 C	Astrup method 146
Ångström (unit) 372	AQP → Aquaporins	at, conversion into SI unit 374
Anion exchange, red blood cells	Aquaporins 24, 166, 174	AT I (A prints print II) 150, 170
124 Anode 50	Aqueous humor 344 Arachidonic acid 268, 276	AT II (Angiotensin II) 158, 170, 182, 184, 212
Anomaloscope 356	Arachnoid villi 310	receptor 184
Anovulatory cycle 298	ARAS (ascending reticular acti-	Ataxia 328
Anoxia (→ also hypoxia) 130, 134	vating system) 322,	Atelectasis 118
ANP → Atriopeptin	332	Atenonol 87 B
ANS (Autonomic nervous system)	Archeocerebellum 326	Athosclerosis → Arteriosclerosis
78 ff., 194, 212 ff., 234,	Area prostrema, chemosensors	atm, conversion into SI unit 374
266, 330	238	Atmosphere, physical, 374
Anterior pituitary 268, 280	Areflexia 320	technical, 374
Anterolateral funiculus, tracts	ARF (ADP-ribosylation factor) 30	Atomic mass 374
318, 322	Arginine 174, 226, 258, 282, 284	ATP 41, 64, 72, 86, 228
Antibodies (→ also immuno-	glucagon secretion 284	control of K+ channels 282
globulins) 94, 98, 100	intestinal absorption 258	coronary vasodilatation 210
Anticoagulants 104	insulin release 282	cotransmitter 84
Anticodon 8, 10	Aromatase 300, 306	creatine phosphate 228
Antidiuresis 164, 166	Arousal activity 332	free enthalpy 41
Antidiuretic hormone → adi-	Arrhenius 40	gain, glucose 73 B
uretin	Arrhythmia 218 ff	neurotransmitter 86
Antigen-antibody complex 98 allergy 100	absolute 200 Areflexia 328	production 12 aerobic 39 C
presentation 96, 98	Arrestin 350	supply of energy, muscle 72
Antigen-dependent cell-medi-	Arsenic (AS) 226	synthesis 12
ated cytotoxicity	Arteries (\rightarrow also artery) 186 ff.,	transport, active 26
(ADCC) 96	188	tubular epithelium 154
Antigens 94, 98	arcuate 150	ATPase (\rightarrow also Na ⁺ -K ⁺ -ATPase,
incomplete 100	bronchial 186	Ca ²⁺ -ATPase) 26, 58,
presentation 96, 98	compliance 188	72, 84
thymus-dependent (TD) 98	coronary myocardial perfusion	,

thymus-independent (TI) 98

ATPS (Ambient temperature	hillock 42	Bilayer of membranes 14
pressure H ₂ O-satu-	membrane capacity 48	Bile 232, 248 ff., 256, 260
rated) 112	myelin sheath 42	acid → bile salts
ATP-synthetase 17	nodes of Ranvier 42	bilirubin excretion 250
Atrial fibrillation 200	reflexes 214	canaliculi 248
flutter 200	saltatory conduction 48	components 248
septal defect 220	Axonal transport 42, 280	ductules 248
tachycardia 200		excretory function 248
Atriopeptin (ANF, ANP) 152, 162,	_	formation 248
170, 182, 218	В	hepatic 248
aldosterone secretion 182	P7	salts 160, 246, 248, 252, 256
receptor 278	B7 protein 98	absorption, terminal ileum
Atrioventricular (AV) block 162,	B cells, islets of Langerhans 282	252
200	lymphocytes 94, 98	body pool 248
node 192, 194 valves 190	B chain, insulin 282	carrier 248 conjugated 248, 252
	B lymphocytes 94, 98 Bacteria, defense 94 ff., 242	enterohepatic circulation
Atrophy, compensatory 272 Atropine 82	intestinal 232, 240, 250, 264	252
transport, tubular 160	lysis 94	function 252
Attention 326, 336	Bainbridge reflex 216	primary 248
automated 336	Balance, body 328, 342	secondary 248
EEG 332	glomerulo-tubular 166	synthesis 248
selective 336	Bar (unit) 374	steroid hormone excretion 294
Atto- (submultiple of a unit) 373	Barbiturates, tubular secretion	Bilirubin 248, 250
Audiometer 366	160	direct 250
Auditory canal 364	Barium contrast medium 264	diglucuronide 250
pathway 368	Barotrauma 134, 110	excretion 250
threshold 339	Barrier, blood-brain 18, 230, 238,	indirect 250
Auerbach plexus (myenteric	310, 330	Biliverdin 250
plexus) 244	blood-CSF 18, 310	Binary digit 312
Autoantibody 100	Basal body temperature 298	Biologic clock 224, 334
TSH receptor 288	Basal ganglia 310, 324	Biotin, intestinal absorption 260
Autogenous inhibition 316 Autoimmune disease 94	labyrinth of tubule cells 154 metabolic rate 226	Biotransformation 232, 250 Bipolar cells, retina 344
Automaticity, heart 192	Base, DNA 8	Birth 94, 304, 220
Autonomic nervous system (ANS)	excess 142	circulation during 220
78 ff., 194, 212 ff., 234,	measurement 146	2,3-Bisphosphoglycerate (2,3
266, 330	triplet, mRNA 8	BPG) 128, 138
Autophagia 12	units, SI system	Bit 312
Autophagic vacuole 12	Basic frequency of vowels 364,	Bitemporal hemianopia 358
Autophagosomes 12	366	Bitter sensors 338
Autophosphorylation, receptor	Basilar membrane 364, 366	Bladder, urinary 70, 79 ff., 148
tyrosine kinase 278	Basket cells 328	control 79 ff., 308
Autoreceptor 86	Basolateral membrane, tubule	Bleeding 218
presynaptic 52, 82, 146	cell 162	time 102
Autoregulation 4, 212	Basophilic granulocytes 100	Blind spot, eye 348, 358
coronary 210	Bathorhodopsin 348	Blindness 344
gastrointestinal 232	Bayliss effect 212	Blinking 344
renal 150	BDGF (brain-derived growth fac-	Blood 88 ff.
Auxotonic contraction 64, 66 AV → atrioventricular	tor) 338 Beat tone 362	brain barrier 18, 230, 238, 280, 310, 330
avD_{O_2} (arterial venous O_2 differ-	Behavior, defensive 330	cells 88
ence) 106, 107 A	nutritive 330	clotting 102
aVF (ECG leads) 196	programmed, hypothalamus	activation 102, 104
aVL (ECG leads) 196	330	endogenous 102
Avogadro's constant (N _A) 20, 374	reproductive 330	exogenous 104
aVR (ECG leads) 196	thermoregulation 224, 330	coagulation disorders 226
Axolemma 42	BEI → iodine	estrogen effects 302
Axon 42	Bel 339	factors 102
capacitive flow of charge 48	Beriberi 226	composition 88
conduction velocity 48, 49 C	Bicarbonate → HCO ₃ ⁻	CSF barrier 18, 126, 310
diameter 48	Bicycle ergometry 76	fibrinolysis 102

Blood flow 122, 212, 232 viscosity 92, 188 second messenger 276 brain, control 212 high altitudes 136 Bradypnea 108 volume 88, 182, 186, 188, 206 central regulation 214 Brain 310ff. coronaries 210, 212 central 204 anatomy 310 determination 168 blood flow 186 gastrointestinal 232 lungs 122, 186 distribution 187 A regulation 212 muscle 214 endurance athletes 77 C cortex properties 92 measurement 168 afferents 324 rate 188 sensors 214 area 1 311 F 324 325 C total 186 area 2 311 E. 324, 325 C regulation, neuronal 214 renal 150 B-lymphocytes 94, 98 area 3 311 E. 324, 325 C skin 212, 214, 224 BMAI.1 334 area 4 311 E. 324, 325 C velocity 92, 188, 190 BMI (body mass index) 230 area 5 311 E. 324, 325 C vessel segments 189 A BMR (basal metabolic rate) 226 area 6 331 E. 324, 325 C function 88 ff. Body fluids 92, 168 area 7 311 E, 324, 325 C glucose 282, 284 compartments 168 area 22 311 E. 370 measurement 168 area 44 311 E. 370 groups 100 area 45 311 E. 370 HCO₃⁻ concentration 142, 146 homeostasis 168 hormone transport 268 osmolality 170 area MI 324, 325 C lipids 256 mass index (BMI) 230 area PM 324, 325 C loss 90, 173, 218 plethysmography 114 area S1 318, 322, 323 A pH 138 temperature 132, 222, 224, area S2 318, 323 A puffer 138, 144 298, 330, 375, 381 C area SMA 324, 325 C normal range 142 circadian variation 224. asssociation areas 328, 336 measurement 146 381 C auditory 368 pressure 74, 86, 184, 188, 202, measurement 224 Brodman's areas 311 E 206, 212, 214 menstrual cycle 298 columns 332 pressure saturated (BTPS) arterial 206 corticocortical signal loops amplitude 206 112 324 angiotensin II effect 184 respiration 132 cortical afferents 324 set point 224 efferents 324 mean 188, 189 A, 206 water → body fluids ultrastructure 332 measurement 206 weight 230 motor 324, 328 normal range 206 regulatory mechanism 230, motosensory 370 physical work 74 280 neuronal circuitry 332 Bohr effect 128 regulation 216 orbitofrontal area 340 renin-angiotensin sysequation, dead space 114, 115 A organisation 332 tem 184 Boiling point, water 375 potentials 332 salt deficit 170 Bomb calorimeter 228 prefrontal association 324 sensors 214 Bone 290, 292 premotor area (PMA) 324 shock 218 break down 290 prepiriform 340 control 214, 216 calcitonin effect 292 primary motor area (MI) 324 diastolic 206 calcitriol effect 292 pyramidal cells 332 pulmonary 122, 206 conduction, sound 364 scarring 338 sensors 214 estrogen effects 302 sensory association 324 systolic 190, 206 growth 302 input 322 venous 204 marrow 88, 90, 94 supplementory motor area sugar 282 ff. Fe homeostasis 90 (SMA) 324 testis barrier 306 megakarvocvtes 102 death, diagnosis 332 transfusion 100 preprocessed precursor cells derived growth factor (BDGF) vessels 188, 212 ff. 338 autoregulation 212 metabolism 290 glucose deficit 242 chemosensors 212 mineralization 292 glutamate, transmitter 336 cross-sectional area 189 A PTH effect 290 hemisphere, dominant 337 diameter 189 A Botulinum toxin 56 metabolism 282 nerve cells 338, 340, 359 function 188 ff. Bowman's capsule 148 holding capacity 189 A Bradycardia, sinus 200 stem (→ also medulla oblongnervous control 214 Bradykinesia 326 ata, pons, and midprecapillary sphincter Bradykinin 184, 208, 214, 269, brain) 310, 318 188 318 motor centers 324, 328 skin 224 coronary vasodilatation 210 survival time, anoxia 130 wall tension 188 salivary glands 236 swelling 172

visual areas 328, 354, 356, 358	Butanol-extractable iodine (BEI)	daily requirement 226, 290
Breathing capacity, maximum (MBC) 118	288 Byte 312	darkness-induced 350 deficiency 290
flat 120	byte 312	equilibrium potential 44
gas (\rightarrow also O ₂ and CO ₂) 106		excretion 178, 290
mechanics 108	C	exocytosis 30
Breast 302, 303	-	fat absorption 252
development 304	c (submultiple of a unit) 373	fetal 290
estrogen effect 302	C17/C20-lyase 300	free 178, 290
progesterone effect 303	C cells, thyroid gland 292	hormone release 280
prolactin effect 303	°C, conversion to °F 375	intake 290
enlargement 304	Ca^{2+} (\rightarrow also Calcium) 36, 226,	intestinal absorption 262, 290,
feeding 303	262	292
iodine 288	absorption, intestine 262, 278, 290, 292	intracellular stores 36 ionized 178, 290
Brightness channel, visual path- ways 356	antagonists 194	light sensors 352
Bristle cell, olfactory bulb 340	ATPase 17 A, 17 B2, 26, 36, 178,	metabolism 178, 290
Broca's area 370	292	milk 290
Brodman's map 310	heart 194	muscle contraction 60 ff
Bronchi 370	muscle 64	/3 Na+ antiport 36
epithelium 110	renal 178	plasma 178, 290
innervation 79 ff.	balance 290	pregnancy 290
Bronchial mucus 110	calmodulin-dependent protein	protein bound 178, 288, 290,
tree 186	kinase II 36, 50	292
Bronchitis 118	cell regulation 36	reabsorption, renal 156, 178
Brown fat 224, 288 Brown-Sequard syndrom 322	channels 36, 62, 178, 348 activation 276	paracellular 172 renal excretion 178
Brunner's glands 244	β-adrenoceptors 84	saliva 236
Brush border 9 f., 154, 158	hair cells 342	sensors 36, 292
intestinal mucosa 244	heart muscle 63 B3, 65 D,	kidney 178
renal tubule 154	194	serum 290
stomach, parietal cells 242	inhibition 84, 276	smooth muscle 70
BSC (bumetanide-sensitive co-	muscle 62	solubility 290
transporter) 162	photosensors 348	store, intracellular 10,36
NH ₄ ⁺ reabsorption 176	regulation 36	IP ₃ 276
BSEP (bile salt export pump) 248,	renal 178	myocard 194
250	ryanodine-sensitive 194	skeletal muscle 60
BTPS (body temperature pressure saturated) 112	voltage-gated 194 clotting process 102	third messenger 276 trigger effect 62
α -BTX (α -bungarotoxin) 56	complexed 290	urine 290, 178
Buffer 124, 138, 140, 142, 144,	concentration 50	CaBP (calcium-binding protein)
378 f.	cytosolic 36, 66, 236	262, 278, 292
base 140	adrenergic transmission	Cabrera circle (ECG) 198
blood 138	84	Caffeine 276
concentration 138, 142	electromechanical coupling 62,	Caisson disease 134
total 146	194	Cajal cells 240, 244
capacity 138, 140	epinephrine 87 B	Cajal's horizontal cells, retina
closed system 140	exocytosis 30	344, 354
curve 380 B hemoglobin 124, 128	extracellular 36, 46 heart 194	cal (calorie), unit 374
nonbicarbonate 126, 140, 144	intracellular 36, 276	Calcidiol (25-OH-cholecalciferol) 158, 292
open system 140	muscle 63 B	Calciferol 226, 292
titration curve 380	smooth 71 B	$1,25-(OH)_2$ -calciferol \rightarrow calcitriol
Bulb, ocular 344	norepinephrine 87 B	Calciol (= cholecalciferol = Vi-
Bumetanide 172	oscillations 36	tamin D) 260, 292
Bundle of His 192	photosensors 350, 352	Calcitonin 36, 37 C2, 274, 286,
Bundle-branch 200	smooth muscle 70	290, 292
block 200	muscle fibers 62, 64, 194	gene-related peptide (CGRP)
α-bungarotoxin (α-BTX) 56	neurons 336, 338	214, 234, 306, 318
Bunsen solubility coefficient 126	serum 290	
Bunte glass burette 114	conductance, GABA _B receptors	

Burning 218

metabolism 282

Calcitriol (1,25-(OH) ₂ -cholecal-	cortisol effect 296	antigen-presenting (APC) 98
ciferol, D-hormone)	nutrition 226	body 42
158, 262, 268, 278,	Carbon dioxide \rightarrow CO ₂	chief 240
290 f. , 292	monoxide (CO) 128	dendritic 96
deficiency 290	Carbonic anhydrase 124, 174, 236	dedifferentiation 272
nuclear receptor 278	gastric 242	differentiation 272
PTH effect on formation 290	inhibitors 142, 172, 344	division 8
Calcium → also Ca ²⁺	red blood cells 124	excitable 42
Calcium-binding protein (CaBP)	renal 174	fat 254
262, 278, 292	salivary glands 236	ganglion, retina 344, 358
Calcium oxalate, urinary stones	γ-carboxylation 102	granular 326
178	Carboxylesterase 246, 252, 256	granulosa 300
Calcium phosphate, solubility	Carboxypeptidase 258	killer 96, 98
product 290	Cardia 240	Leydig, interstitial 306
urinary stones 178	Cardiac (→ also heart)	mast 100, 256
Caldesmon 70	arrest 200	membrane 2, 8, 14
Calmodulin 36, 276, 278, 336	arrhythmias 180, 200	apical 18
smooth muscle 70	cycle 190	basolateral 18
photo sensors 350	failure 204	function 2. 14
Calmodulin-dependent protein	muscle 68	ion conductance 32
kinase II 36, 50		
	output (CO) 70, 74, 122, 136,	proteins 14
neurons 336	186, 188, 204, 206,	structure 14
Caloric equivalent 228	218, 220	permeability 2
value, physical 228	fetus 220	migration 30, 58
Calorie 228	paralyis 198	neuroendocrine 266
conversion into SI unit 374	valves 190	nuclear pores 10
Calorimetry 228	work 202	nucleus 8, 288
$cAMP \rightarrow AMP$, cyclic	Carnosine 158	structure and function 8 ff.
Canaliculi 248	Carotinoids 350	organelles 8 f.
bile 248	Carrier 22	pacemaker 244
lachrymal 344	affinity to 28	parietal 240, 242
parietal cells 242	free fatty acids 252	principal (chief) 162, 180
Canals, semicircular 342	liver cells 250	potential 366
Can sugar → saccharose 258	passive 28	replacement 244
Capacitation of sperm 302, 308	transport 158	retina 358
Capacitive current, membrane 48	CART (cocaine and amphet-	Sertoli 306
Capacity, buffering 140	amine-regulated tran-	structure and function 8 ff.
electric, nerve 48	script) 230	T-helper 98
Capillaries 188	Cartilage 280	Cellulose 226, 264
blood pressure in 150, 209 B	Catabolism, cortisol 296	Celsius, conversion to °F 375
lung 208	Catalase 14, 96	Centi- (submultiple of a unit) 373
characteristics 188	Cataract 346, 358	Central blood volume 204
diffusion processes 208	Catecholamines 84, 268, 274	canal 310
exchange processes 208, 377	receptor types 55 F	chemosensors 126
filtration 208	Catechol-O-methyltransferase	nervous system (CNS) 303,
fluid exchanges 208	(COMT) 86	310 ff.
glomerular 150	Cathode 50	anatomy 310
	cBAT (canalicular bile acid trans-	
peritubular 150, 151 A, 166		autonomic centers 78
permeability 208, 377	porter) 248	venous pressure 186, 190 , 204 shock 218
pulmonary 106, 122	CBG (cortisol binding-globulin =	
reabsorption 208	transcortin) 296, 303	Centralization, circulation 218
Capsaicin sensors 314, 318	C cells, parafollicular 36, 286	Centrosome 14
Carbamate, CO ₂ transport in	CCK → Cholecystokinin	Cerebellum 310, 322, 324, 326ff. ,
blood 124	CD4 protein 98	328, 342, 360, 370
Carbohydrate 226, 228, 246, 258,	CD8 protein 98	eye movement 360
282, 296	CD28 protein 98	lesions 328
absorption, intestinal 258	CD40 ligand 98	nystagmus 360
caloric equivalent 228	CD40 protein 98	speech 370
digestion 258	CD45 receptor 278	Cerebrosides 14
pancreatic enzymes 246	CD95 protein (= Fas) 98	Cerebrospinal fluid (CSF) 126,
energy substrate 228	Cecum 232, 264	144, 168, 310

Cell(s) 8 ff.

CO₂ 126, 144

pH 126, 144	types 55 F	Circulation, blood 186 ff., 212 ff.
Cerebral cortex 310	second messenger 55 F, 276	autoregulation 212
function, glucocorticoid effect	stomach 242	Bayliss-effect 212
296	Cholera toxin 262, 276	hormonal 212
Ceruloplasmin, iron oxidation 90	Choleresis 248	myogenic effect 212
Cervix uteri 298, 302	Cholesterol (CHO) 14, 114, 254,	neuronal 214
menstrual cycle 298	268, 294, 303	local metabolic effect 212
os 298, 303	bile 248	sympathetic nervous sys-
progesterone effects 303	crystals 248	tem 214
cGMP → GMP, cyclic	esters 252, 256	birth 220
cGMP phosphodiesterase, retina 348	hydrolase 252	centralisation, flow 214, 218
	LDL 254	control 212
CGRP (calcitonin gene-related peptide) 214, 234,	lipoproteins 254 feces 256	coronary 210 fetal 220
306, 318	LDL 254	portal 232
gallbladder 248	lipoproteins 256	pulmonary 186, 188
second messenger 274	membranes 14	pulse wave velocity 190
stomach 242	micelles 248	regulation 212, 214 ff.
Chamber, eye 344	steroid hormone synthesis 294	resistances 187 A, 188, 206
Channels, ion 34	storage 255 B	total peripheral resistance
Chaperone protein 10	synthesis 256	206
Chemokines 94, 266, 274	Cholesterol-7α-hydroxylase 248	venous return 204
receptors 94	Cholic acids (cholates) 248	cholehepatic 248
Chemosensors 142	Choline 82, 160	enterohepatic 248, 252
area postrema 238	acetyl transferase 82	bile salts 252
central 126, 132, 144	nerve terminals 82	bilirubin 250
for glucose 282	$receptor \rightarrow cholinoceptors$	Circulatory "center" 214
respiratory control 132	tubular transport 160	failure 216
small intestine 240	Cholinesterase 56	reflexes, homeostatic 216
Chemotaxis 94, 308	inhibitors 56	sensors 214
Chenodeoxycholic acid 248	Cholinoceptors 82	shock 218
Chest 108	antagonist 82	Circumventricular organs 280,
Chest leads (ECG) 196	heart 194	310, 330
Chiasma, optic 358	ionotropic 56	11-cis retinal 348
Chief (principal) cells, kidney 162 stomach 240, 242	M-types, second messenger 274, 276	Cisterns, endoplasmic reticulum 10
Chinin, taste 338	nicotinergic 56	Citrate 12. 102
Chloramphenicol 250	salivary glands 236	complex former 178
Chloride → Cl ⁻	stomach 242	cycle 12, 73 B3
CHO (cholesterol) 14, 114, 254,	types 55 F, 56	inhibition of blood clotting 102
268, 294, 303	Chorda tympani 339 C	renal excretion 174
Cholagogue 248	Chorionic gonadotropin, human	Citric acid → citrate
Cholates (→ also bile salts) 248	(HCG) 304	Cl- 378
Cholecalciferol (calciol = Vitamin	Choroid plexus 310	absorption in intestine 262
D) 260, 292	Christmas factor 102	balance, body 170
formation in skin 292	Chromium (Cr) 226, 228	channels, lysosomes 14
25-OH-cholecaliferol (calcidiol)	Chromatin 8	tubular 162
158	Chromosomes 8	concentration, intracellular 44
1,25 (OH) ₂ -cholecalciferol (cal-	single set (haploid) 306	conductance membrane 34, 44
citriol, D-hormone)	Chronotropism, heart 194	distribution, active/passive 34
Chalagratakinin (CCV) 220, 224	Chylomicron 254, 260	equilibrium potential 44 excretion 162
Cholecystokinin (CCK) 230, 234, 240, 248	remnants 254, 256	
cerebral cortex 332	Chyme 232, 258 pH 240	renal handling 156, 162 secretion in intestine 262
esophagus 238	Chymotrypsin 246, 258	in pancreas 246
gallbladder 248	Chymotrypsinogen 246	salivary glands 236
pancreas secretion 246	Cilia, bronchial 58, 110	stomach, parietal cells 242
receptors	hair cells 342, 364	Clarifying factor → Heparin
gallbladder 248	Ciliary body 70	Clathrin 14, 28
hypothalamus 330	muscle 344	Clearance factor (heparin) 256
pancreas 246	process 344	esophagus 238
stomach 242	zonules 344	renal 152

response curve 132

Cleft, synaptic 42, 56, 82 solubility coefficient 126 factors 94, 96 Climacteric 298 total 126 Complementary colors 356 Compliance Climax 308 transport, blood 124 Climbing fibres, cerebellum 328 α - γ -Coactivation 316, 320 blood vessels 188, 206 Clitoris 308 Coagulation, disseminated intralung and thorax 116, 118 CLOCK 334 vascular 104 measurement 112 Clock, biologic 224, 240, 334, 354 Coagulopathy consumptive 104 COMT (catechol-O-methyl-Clonal selection 98 Coated pits 28 transferase) 86 Clonidine 84 87 B Coatomer 30 Concentration, units 372 f. Closed system, thermodynamics Coatomer-coated vesicles 30 Conception 308 Cobalamines (vitamin B 12) 90, Conditioned reflexes 236, 242 41 buffer 140 92, 226, 260 Conductance 46, 375 Clothing 224 absorption 260 electrical 22 for Cl- 44 Clotting, blood (→ also coaguladeficiencies 90, 260 tion) 102 ff. receptor 28 for K+ 44 disorders 226 storage 260 hydraulic 24 cmH2O, conversion into SI unit ionic 32 transport proteins 260 unit 375 374 Cobalt (Co) 226 cMOAT (canalicular multispecific Cocaine and amphetamine-regu-Conduction, atrioventricular organic anion translated transcript (ECG) 198 deafness 364 porter) 250 (CART) 230 CNS → central nervous system Cochlea 364, 368 defects, heart 200 CO (carbon monoxide) 128 Cochlear duct → scala media heart 192, 194, 200 binding to hemoglobin 129 C amplifier 366 heat 222 CO → also cardiac output nucleus 368 nerve fibres 42, 50 CO2 (carbon dioxide) 106, 120, Code, genetic 8 saltatory, nerve 48 264 Coding, neuronal signals 312 velocity, measurement 50 binding, blood 124ff. Codogen 8 Conductivity → conductance buffer system 138 Codon 10 Cones 344, 348, 350, 356 concentration Coital capacity, testosterone 306 adaptation 352 blood 124ff. Cold sensors 224, 314 classes 356 influence of O2 saturation Colipase 246, 252 density 348 Collaps, orthostatic 204 light absorption maximum 356 cerebrospinal fluid 126, 144 Collagen 102, 103 A Conjugate eye movement 360 influence O2 saturation 126 Collaterals, axon 42 Conjugation processes 160 chemosensors 134, 144 with glucuronic acid in liver Collecting duct 148, 162, 166, 174 chronic retention 132 150, 250 Colliculus superior 326 diffusion, tissue 130 Colloid, thyroid 286 with glutathione in liver 160, dissociation curve 126 Colloidal osmotic pressure 92, dissolved in plasma 124ff. 378 Connectin (= titin) 60, 66 Connexin 16, 19 C distribution in blood 124 Colon 262, 264 H₂O absorption 262 elimination 106 Connexons 16 /HCO3- buffer 138, 140 K+ secretion 262 Conn's syndrome 182 Consciousness 322, 336 Krogh's diffusion coefficient Color blindness 352, 356 120 constancy 356 Consensual light response 359 large intestine 264 opponency 354 Consolidation, memory 336 partial pressure 124 perception 356 Consonants 370 alveolar 120 triangle 356 Constancy, of color 356 diving 134 vision 356 of shape 356 blood 126 contrast 354 of size 356 increased 144 Colors, additive mixture 356 Constipation 264 chemical respiratory complementary 356 Contact phase, blood clotting 103 stimulant 132 subtractive mixture 356 chronical increase 132 Comfort zone (ambient tempera-Contraceptives 300, 303 hyperventilation 120 ture) 224, 226 Contractility, heart muscle 194 Contraction, afterloaded 66 normal range 142 Compensation, renal 144 respiratory 142 auxotonic 64, 66 regulation by respiration Compensatory atrophy 272 isometric 64, 66 venous blood 120 hypertrophy 272 isotonic 64, 66 plasma 124 ff., 138, 144 pause 200 velocity, skeletal muscle 68 Competitive inhibition 56 myocard 204 production 142, 228

Complement cascade 96

Contracture 66

Contrast, auditory pathway 368	receptors, nypotnaiamus 330	Cyclooxygenase (COX) 242, 269
enhancement 368	second messenger 274	inhibitors 104
of stimuli 312, 328	releasing factor → cortico-	Cyclosporin A 98
retina 354	liberin	Cystic fibrosis 110, 246
simultanous 354	$hormone \rightarrow corticoliberin$	Cystine 158, 174
successive 352, 354	Corti's tunnel 364	Cystinuria 158, 258
Control circuit 4	Cortisol 92, 182, 256, 272, 280,	Cytochrome oxidase 130
humoral 272	296	Cytochrome P450-epoxygenase
system 4	binding-globulin (CBG) 296,	269
Controller 4	303	Cytokines 266, 268, 274
Convection 24, 222	effect, permissive 296	cortisol 296
Converting enzyme, angiotensin-	fight or flight 330	receptors 278
(ACE) 184	hypothalamus 330	Cytokinesis 58
Convergence of signals 358	precursors 296	Cytolysis 96
response, eyes 360	stress 296	Cytoplasm 8
Cooperativity, positive, hemo-	synthesis 296	Cytosin 8
globin 128	transport 296	Cytosis 28, 58
Copper (Cu) 226	Cortisone 296	Cytoskeleton, migration 30
iron oxidation 90	Cotransmitter 52, 84, 86, 326	Cytosol 8
Coprosterol 256	Cotransport, definition 26	ions 45 B, 93 C
		1011S 43 B, 93 C
Core sleep 334	Costimulatory signal 98	
temperature 222, 224	Coughing 132, 320, 370	
Cornea 344, 346	Coumarin 104	D
Corneal reflex 320, 358	Countercurrent exchange, heat 224	1/ 1 1/ 1 6 1/ 000
Corona radiata 308	kidney 164	d (submultiple of a unit) 373
Coronary blood flow 210	system 164	da (multiple of a unit) 373
insufficiency 210	Countertransport 26	DAG (Diacylglycerol) 36, 82, 84,
reserve 210	Coupling, electromechanical 62	274, 276
Corpus amygdaloideum 310, 330,	COX (cyclooxygenase) 104, 262,	Dalton (Da), unit 374
336	269	Dalton's law 106
callosum 310	CPPV (continuous positive pres-	Dark adaptation, eye 348, 352,
geniculatum laterale 356, 358	sure ventilation) 110	354
mediale 368	Creatine 72	receptive field 354
luteum 298, 300	Creatine phosphate 72, 76, 228	Darkness-induced Ca ²⁺ 350
pregnancy 304	muscle reserve 73 B	Dead space, artificial respiration
progesterone production 303	standard free enthalpy 41	110
stomach 240	clearance, endogenous 152	functional, respiration 114,
Correlations 383	Cretinism 288	122, 134
Corresponding areas, retina 360	CRF → cortiboliberin	increase 120
Cortex, auditory 368	CRH → corticoliberin	functions 114
associative 326	Crigler-Najjar syndrome 250	snorkel breathing 134
		0
cerebellum 326	Cristae, mitochondria 12	ventilation 106
cerebral 332, 310, 326	Crista, ampulla 342	volume (V _D) 114
frontal 330	Cross test, blood types 100	Deafness 364f., 370
Corticoids, albumin binding 268	Crypts, large intestine 264	Debré-Toni-Fanconi syndrome
globulin binding 268, 303	small intestine 244, 262	158
Corticoliberin (= CRH) 230, 269,	Crypts of Lieberkühn 244, 262	Deca- (multiple of a unit) 373
272, 274, 280, 296,	CSF → cerebrospinal fluid	D cells, islets of Langerhans 282,
304	Cupula 342	284
birth 304	Curare 56,110	stomach 242
cortisol secretion 296	Current, alternating 50, 375	DCT (distal convoluted tubule)
placenta 304	direct 50, 375	148
receptors, hypothalamus 330	endplate 56	Deceleration, unit 374
second messenger 274	miniature endplate 56	Deci- (submultiple of a unit) 373
secretion, interleukin effect	unit 375	Decibel (dB), unit 362
296	Current-voltage curve 33 B3, 34	Decompression sickness 134
Corticosterone 182	CVP (central venous pressure)	Decurarinization 56
synthesis 294	190, 204	Dedifferentiation, cells 272
Corticotropin (ACTH) 269, 272,	Cyanide 130	Defecation 264
280, 294, 296	Cyanosis 110, 130	Defense mechanism 96 ff., 232,
aldosterone synthesis 182	Cyclic AMP → AMP, cyclic	330
cortical synthesis 206	CMD CMD cyclic	intectine 222

Defensins 96	Diencephalon 310	vision 360
Defibrillation 200	Diet 226	DIT (diet-induced thermogene-
Deflation reflex 132	induced thermogenesis (DIT)	sis) 228
Deglutition 238	228	DIT (diiodotyrosine residues) 286
Degrees Celcius 375	vegetarian 142	Diuresis 164, 172
Fahrenheit 375	Differential threshold, sound,	osmotic 172, 176
7-dehydrocholesterol 292	frequency 368	influence on K+ excretion 182
Dehydroepiandrosterone (DHEA)	intensity 368	salt and water homeostasis
295, 304, 306	optical 352	173 B
adrenal cortex 296	smell 340	Diuretics 172, 182, 218
sulfate (DHEA-S) 304	taste 338	influence on Ca2+ reabsorption
Deiodinase 286, 288	Differentiation, cells 272	178
Deiter's nucleus 326, 328	Diffusion 20, 208	K+ excretion 172, 182
Delayed immune reaction 100	capacity 22	osmotic 172
Deletion, clonal 94	capillaries 208	tubular secretion of 156, 160,
Dendrites 42	coefficient (D) 20	172
Deoxycholate 248	Krogh's (K) 22	Diurnal rhythm 334
Depth perception, visual 358	conductance 22	Diving 134
Depolarization 46, 66	distance 20	oxygen toxicity 136
action potential 46	driving force 20, 22	DNA (desoxyribonucleic acid) 8
permanent, muscle 56, 60	equilibrium 20	double helix 8
smooth muscle 70	faciliated 23, 158, 258	DNAses, pancreas 246
Derepression 8	Fick's first law 20, 22, 120	Döderlein bacilli 302
Dermatitis, allergy 100	gas 22	Dominance columns, ocular 358
Dermographism 214	ions 22	L-Dopa 84
Desensitization 52, 276, 318	net 20	Dopamine 84, 268, 269, 274, 280,
smell sensors 340	non-ionic 22, 156, 176	303, 326
Desmin filaments 14	potential 22, 32, 44	cerebral cortex 332
Desoxycorticosterone 295	rate 20	menstrual cycle 298
Desoxyribonucleic acid → DNA	"simple" 20	neurons 330
Desoxyribose 8	unidirectional 20	receptor, striatum 326
Detoxication processes 160	Digestion 232, 236, 252, 258	second messenger 274
Deuteranomaly 356	carbohydrates 236, 258	synthesis 84
Deuteranopia 356	impaired 246	transmitter 326
Dextrin, α-limit 258	lipids 252	Dopamine-β-hydroxylase 84
DHEA → dehydroepian-	organs 232	Dorsal root, spinal cord 322
drosterone	proteins 258	Double helix, DNA 8
DHPR (dihydropyridine receptor)	Digitalis 194	vision 360
62, 63 B, 65 D	Dihydropyridine receptor (DHPR)	Down regulation, hormon recep-
DHT (5α-dihydrotestosterone)	62, 63 B, 65 D	tors 30
306	heart 194	Na+-phosphate symport carrier
Diabetes insipidus 166, 218	5α-Dihydrotestosteron (DHT)	178
mellitus 142, 158, 218, 230, 284	306	Dreams 334
coma 218	synthesis 294	Driving "force" 20, 38
osmotic diuresis 172	testes 306	pressure difference 108
type II, obesity 230	1,25-dihydroxycholecaliciferol →	Dromotropism, heart 194
Diacylglycerol (DAG) 36, 82, 84,	calcitriol	D-Sensor 312, 314, 316
274, 276	Diiodotryrosin residues (DIT) 286	Dubin-Johnson syndrome 250
Diadochokinesia 328	Dilator muscle, pupil 344	Duct, collecting 174
Diapedesis 94	Dim-light vision 348	Ductus arteriosus 220
Diaphragm 108	Diopters (dpt) 346	patent 220
Diarrhea 138, 142, 173, 258, 262,	Dipeptidases 258	venosus 220
264	Dipeptides 158, 258	Duodenum (→ also intestine)
acid-base balance 138	Diplopia 360	234, 244, 258
lactase deficiency 258	Direct bilirubin 250	gastrin production 234
salt and water homeostasis	Disaccharide, digestion 258	GIP production 234
173 B	Disinhibition 340	secretin production 234
Diastole, heart 190	thalamus 326	Dwarfism, T ₃ /T ₄ deficiency 288
Diastolic depolarization 192	Dissociation constant 378 f.	Dynamic work, negative 74
Diathermy 50	Distal stomach 240	positive 74
Dicarboxylates, tubular transport	tubule 148, 162, 166, 178	Dyne, conversion into SI unit 374
158, 160	Distant hearing 368	Dynein 58

Dynorphin 86, 318	Electrochemical gradient 26	Endothermic reaction 38
Dysmetria 328	potential 32	Endplate current 56
Dyspnea 108	Electrodiffusion 22, 34	motor 56
Dystrophin 60	Electroencephalogram → EEG	blocking substances 56
	Electrolyte homeostasis 168 ff.	reversal potential 56
-	Electromotility, outer hair cells	potential (EPP) 56
E	366	Endurance limit 72
E (multiple of a unit) 272	Electrophoresis 93 B	Enema 264
E (multiple of a unit) 373	Electrotonic transmission 48, 54 D	Energy, activating 39
E1 → estrone E2 → estradiol	Embolisms 104	basic daily 226 chemical 228
E3 → estriol	diving 134	expenditure, total (TEE) 226
Ear 362, 364, 366, 368	Eminence, median 280	homeostasis 230
drum 364	Emission, sperms 308	metabolism 38, 222, 228, 282
in diving 134	Emissions, evoked otoacoustic	cortisol effect 296
Eavesdropping 368	366	need 226
ECF → extracellular fluid	Emotions 336	production 38, 228
ECL (enterochromaffin-like cells)	expression 330	reserves 282
242	limbic system 330	sources 226
ECG (electrocardiogram) 190,	Emphysema 114, 118	storage 230
196 ff.	Emptying rates, gastrointestinal	substrates 72, 226 ff., 254, 282
atrial depolarization 196	tract 233, 240	254
cardiac cycle 190	Emulsification of fats 252	turnover 38, 226, 228
electrical axis 198	Encoding, information 312	units 374
electrolyte disturbances 198	End diastolic pressure 202	Enkephalin 52, 86, 234, 258, 326
integral vector 196	volume (EDV) 190, 202	ENS (enteric nervous system)
leads 196	systolic volume (ESV) 190, 202,	234
myocardial infarction 198	220	Enteroglucagon (= GLP-1) 282,
ventricular depolarization 196	Endergonic reaction 38	284
repolarisation 196	Endings, annulospiral 316	Enterohepatic circulation 248,
Ectopic pacemaker, heart 200	Endocochlear potential 366	250
Edema 172, 208	Endocrine cells 268	Enteropeptidase 246
causes 208	gland 268	Enthalpy 38
extracellular 172, 208	growth 272	Entrainment, brainwaves 332
intracellular 172	system 266 ff.	Entropy 38
local 302	Endocytosis 12, 28, 90, 286	Environment, internal 2, 78, 266
pulmonary 118, 120, 122, 132,	receptor-mediated 12	Enzyme, function 40
144, 172, 208	kidney 158	Ependymal cells 338
Edinger-Westphal nucleus 360 EDHF (endothelium-derived	thyroid gland 286 transferrin 90	EpETrE (= Epoxyeicosatrienoates EE) 269
	Endolymph 342, 364	Epidermal growth factor (EGF)
hyperpolarizing factor) 214	Endonymph 342, 364 Endometrium 303	278
EDTA (ethylendinitrilo-	Endopeptidases 242, 246, 258	Epilepsy, EEG 332
tetraacetat), blood	gastric juice 242	Epinephrine (= adrenalin) 84ff. ,
clotting 102	renal tubule 158	160, 194, 212, 256,
EDV (end diastolic volume) 190,	Endoplasmic reticulum 10, 12, 26	268, 274, 282, 288
202	Endorphin 280, 318	adrenal medulla 86
EEG (electroencephalogram) 332,	Endosomes 12	circulatory shock 218
334	receptors 28	coronary vasodilatation 210
11,12-EET (Epoxyeicosatrienoate)	transcytosis 28	defensive behavior 330
214, 269	Endothel(ium) 102	heart 194
EGF (epidermal growth factor)	exchange processes 208	influence on insulin release
278	function 18	282
Eicosanoids 214, 269, 272, 276	heparin source 254	on K+ uptake 180
Einthoven leads (ECG) 196, 198	lipoprotein lipase (LPL) 254,	lipolysis 218
Ejaculate 308	256	metabolic effect 283 A, 285 C
Ejaculation 308	NO synthase 82	pheochromocytoma 216
Ejaculatory center 308	Endothelium-derived hyper-	production, cortisol influence
Ejection fraction 190	polarizing factor	296
phase, heart 190, 202	(EDHF) 214	receptor types → adrenocep-
Elastase 258	Endothelin 212, 280	tors

 $Electrocardiogram \rightarrow ECG$

second messenger 276

Epinephrine second messengers	ECG leads 196	volume, forced 118
→ adrenoceptors	pleural pressure, measurement	first second (FEV ₁) 118
synthesis 84	108	relative 118
tubular transport 160	sphincter 238	Expired air 107 A
vasoactivity 212	Essential amino acids 226	Exponent, calculation with 380 f
Epiphysis (= pineal gland) 334	fatty acids 226	Export protein, synthesis 12
Epiphysis (of bones), estrogens	Estradiol (E ₂) 294, 298, 300, 302 ,	External auditory canal 364
302	306	intercostal muscle 108
testosterone 306	synthesis 294	Extracellular fluid (ECF) (\rightarrow also
Epithel, function 18	testis 306	H ₂ O) 34, 93 C, 148,
Epoxyeicosatrienoates (EpETrE =	Estriol (E ₃) 294, 300, 302, 304	152, 168
EE) 269	Estrogens 268, 294, 298, 300,	indicators 168
11,12-Epoxyeicosatrienoic acid	302, 304	ions 93 C
(11,12 EET) 214	actions 302	Extrafusal fibers, muscle 316
EPP (endplate potential) 56	comparison 302	Extrasystole 200
EPSP (excitatory postynaptic	degratation 302	Eye (→ also visual, retina and
potential) 52, 56, 82, 320	menstrual cycle 298	opt), accommoda-
cerebral cortex 332	oral therapy 302	tion 344, 346
early 82	placenta 304 synthesis 294, 302	adaptation to light 352 blind spot 348
late 82	Estrone (E ₁) 300, 302	color vision 356
peptidergic 82	synthesis 294	dark adaptation 356
Eq (equivalent), unit 375	ESV (end systolic volume) 190,	far point 344
Equilibrium organ 328, 342	202	focal lenght 346
nystagmus 360	Euphoria, diving 134	point 346
concentration 32	Evans blue, indicator 168	intraocular pressure 346
constant (K) 40	Evaporation of water, heat loss	light rays, physics 346
potential 32, 44	222	movements 360
ions 45 B	Exa- (multiple of a unit) 373	muscles, external 360
Equivalent, caloric 228	Exchange carrier, Na+/H+ 26 f.,	near point 344
mass 375	162, 174 f., 276	nodal point 344
ER → endoplasmic reticulum	Excitable cells 42, 44, 46	optical axis 346
Erection, genital organs 214, 308	Excitatory postsynaptic potential	photochemistry 348
Erg, conversion into SI unit 374	(EPSP) 52, 56, 82, 320	postural motor control 328
Ergocalciferol 292	Excretion (→ also kidney) 160,	reflex movements 342
Ergometry 76	174	refraction 346
Erogenous areas 308	fractional 152	structure 344
Erythroblasts 90 Erythrocytes 88, 90	Exercise 72, 76, 107, 120, 282 cardiac output 76, 74	Eyelids 359
crenation 92	O_2 uptake 74	
effect of hypertonicity 88, 100,	respiration 74	F
124, 128	Exergonic reaction 38	•
fluidity 92	Exocrine glands 246	F (Faraday constant) 22, 32 F
life span 88	Exocytosis 28 f., 30, 50, 86, 286	(fluorine) 226
maturation 88	epinephrin 86	f (submultiple of a unit) 373
mean corpuscular hemoglobin	constitutive 30	°F, conversion to °C 375
(MCH) 88	glucagon 284	Facilitated diffusion 158
concentration (MCHC)	salivary enzymes 236	Fåhraeus-Lindqvist effect 92
88	thyroid glands 286	Fahrenheit, conversion 375
volume (MCV) 88	Exopeptidases 246	Fallopian tube 308
metabolism 282	Exothermic Reaction 38	Falsetto 346
pH 126	Expansion, clonal 94, 98	Fanconi-Debré-Toni syndrome
proliferation 88	Expectancy potential, cortical	158
viscosity 92	324	Faraday constant (F) 22, 32
Erythropoiesis 90	Expiration 108	Far point, vision 344, 346
cobalamin 90 folic acid 90	maximum flow rate 118 muscle 108, 132	Farsightedness 346 Fas (= CD95) 98
high altitude 136	pressure difference, driving	ligand 98
inefficient 90	force 108	Fascicular zone, adrenal cortex
Erythropoietin 88, 148, 218	work 116	294
Escalator, cilial 110	Expiratory curves, maximal 117	Fasting 282
Fsonhagus 232 238 240	flow maximum 118	T3 synthesis 288

Force-velocity curve, muscle 68

Fat 228, 246, 252, 256, 282 absorption, intestine 252	Femto- (submultiple of a unit) 373	Flexor muscles 328 withdrawal reflex 320
brown 224, 288	Ferrireductase 90	Floccunodular lobe 326
caloric equivalent 228	Ferritin 90	Flow rate, blood 188
cells 256	Fertility 306	unit 374
chemical structure 227 B	Fertilization 308	resistance, blood circulation
daily intake 252	Fetal circulation 220	188
depots 230, 282	zone, adrenal cortex 304	velocity 188
digestion 236, 252	Fetoplacental unit 304	unit 374
pancreas 246	Fetus 220	Fluid compartments 168
phases 253 B	O ₂ supply 220	measurement 168
emulsification 240, 252	FEV ₁ (forced expiratory volume,	Fluid exchange, capillaries 208
energy substrate 228	first second) 118	extracellular 168
metabolism, insulin 284	Fever 100, 224, 288	intracellular 92, 168
Fatigue 72, 76	FF (filtration fraction) 24, 152,	ounce, conversion into SI unit
Fat-soluble vitamins 252, 260	389	374
Fattened animals 284	FGF (fibroblast growth factor)	Fluidity of erythrocytes 92
Fatty acids 210, 252, 254, 262,	278	Fluorine (F) 226
282, 284	Fiber → nerve, neuron, motor	Flutter, atrial 200
cyclooxygenase 269	neuron and muscle	Folic acid 90, 226, 260
essential 226	Fibers, diet 264	absorption, intestine 260
free (FFA) 282	Fibrillation, atrial 200	daily requirement 260
carrier 252	ventricular 200	deficiencies 90
glucagon action 284	Fibrin 102, 104	storage 260
lipoprotein lipase 254	Fibrinogen 102, 104	Follicle, dominant 298, 300,
myocard metabolism 210	Fibrinolysis 102, 104	303
sources 257 D	Fibrinopeptides 104	graafian 298
storage 257 D	Fibrin-stabilizing factor 102, 104	primordial 298
target sites 256	Fibroblast 94	progesteron synthesis 303
transport in blood 254	interferons, release 96	stimulating hormone → fol-
uptake in cells 254	migration 30	litropin
supply of energy 72	Fibroblast growth factor (FGF)	thyroid gland 286
liver 254, 284	278	Follicle-stimulating hormone-re
Fe (iron) 88, 90, 92, 128, 226, 262	Fibronectin 102	leasing hormone →
absorption in intestine 90	Fick's first law of diffusion 20,	gonadoliberin
carrier, intestinal mucosa 90	120, 388	Follicular phase 298, 300
deficiency 90	Fick's principle 106, 130, 389	Follitropin (FSH) 269, 298, 300,
functional 90	kidney 150	306
hemoglobin 128	Field, receptive 314, 354	man 306
degradation 250	visual 358 Fila olfactoria 340	menstrual cycle 298
intake 90, 226		peak 300
metabolism 90 overload 90	Filament sliding, smooth muscle 71 B	receptor density 300 second messenger 274
poisoning 90	striated muscle 62	secretion, activin 306
pool 90	Filling phase, heart 190	DHT 306
recycling 90	pressure, receptors for, heart	estrogens 300, 306
storage 90	214	inhibin 300, 306
transferrin 90	Filtration 24	neuropeptid Y 300
transport, plasma 90	capillaries 208	norepinephrine 300
FE (fractional excretion) 152	coefficient 208	progesterone 300
Feces 250, 262, 264	equilibrium 150	pulsatile 298
Fechner's law 354	fraction, glomerular (FF) 24,	testosterone 306
Feedback, control, hormones 272	152, 389	Food deprivation, energy reserve
negative 4, 272, 286, 300, 306,	glomerular 148	282
318	pressure, effective 208	Foot, conversion into SI unit 372
TSH secretion 288	kidney 152	Foramen ovale 220
neuroendocrine 272	shock 218	patent 220
positive 272, 300, 304	First messenger → hormones	Force, unit 374
tubuloglomerular (TGF) 172,	Fitzgerald factor 102	Forced expiratory volume (FEV)
184	Fixed acids 174	118
Feet, conversion into SI unit 372	Fletcher factor 102	vital capacity 118

Flexion reflex 320

Forgetfulness of words 370	insulin secretion 282	transducin 348
Formant 370	Gall bladder 248	Gene expression 8
Formatia reticularis 238, 322,	stones 248	regulation 12
328, 340	posthepatic jaundice 250	Genetic code 8
vomiting center 238	Gallon, conversion into SI unit	Genital tract 214, 298, 306
Formic acid, non-ionic diffusion	374	female 298
22	Ganglia, vegetative 78, 83 A	male 306
Forskolin 276	neurotransmission 83 A	innervation 79 ff.
Fovea centralis, retina 344, 348,	Ganglion cells, retina 344, 354,	Germ cell 306
358	358	Gestagens 303
Fraction, respiration gas 106	vestibular 342	$GFR \rightarrow glomerular filtration$
units 376	GAP (GTPase-activating protein)	GH (growth hormone) → soma-
Fractional concentration → frac-	350	totropin
tion	Gap junction 18, 58	GH-IH → somastotatin
excretion (FE) 152, 154	astrocytes 338	GH-RH → somatoliberin
Frank-Starling mechanism, heart	heart muscle 192	GI → gastrointestinal
68, 202, 204, 216, 218	regulation 37 A	Gibbs-Donnan distribution 44
FRC (functional residual capacity)	smooth muscle 70	Gibbs-Helmholtz equation 38
112, 114, 116	uterus 304	Giga- (multiple of a unit) 373
Free enthalpy 38	Gas constant 20, 24, 32	GIP (glucose-dependent in-
fatty acids 210, 252, 254	equation, alveolar 120, 136, 389	sulinotropic peptide)
water, urine 172	ideal 112	234, 240
Freezing point, H ₂ O 375	exchange 106, 120	esophagus 238 insulin seretion 282
Frequency, fundamental 362, 370 inotropism 194, 204	impairment 120 Gases 106, 264	stomach 242
unit 374	Gastric (\rightarrow also stomach), acid	Glands, Brunner 244
Fructose, intestinal, absorption	242	bulbo-urethral 309
258	function, glucocorticoid effect	endocrine 268
renal reabsoption 158	296	exocrine 246
FSF (fibrin-stabilizing factor) 102	inhibitory peptide (obsolete	gastrointestinal 232 ff., 244,
FSH → follitropin	name for GIP) 240	262
/LH-RH → gonadoliberin	juice 238, 242, 258	parathyroid 236
Fuel value, physiological 228	pH 242	pineal 334
Functional minimum, protein in-	reflux 238	salivary 214
take 226	mucosa, protection 242	sublingual 204, 236
residual capacity (FRC) 112,	secretion 242	submandibular 236
114, 116	ulcers 242	sweat 222
Fundamental frequency, vowels	Gastrin 234, 240, 242	Glasses 346
370	esophagus 238	Glaucoma 344
Fundus, stomach 240	insulin secretion 282	Glia 338
Fungi, defense against 94	second messenger 276 stomach 234, 240	Glial cell line-derived neu-
Funicular myelosis 260 Furosemide 172	Gastrin-releasing peptide → GRP	rotropic factor (GDNF) 338
Fusimotor set 316	Gastrii-releasing peptide → GKi Gastrocolic reflex 264	Glioma 338
Tushilotoi set 510	Gastrocone renex 204 Gastrointestinal (GI) tract 78,	Globulin 92
	232 ff.	cortisol-binding 296
G	bacteria 240, 264	testosterone-binding 307
_	blood flow 186, 232	thyroxin-binding 288
ΔG (free enthalpy) 38	hormones 234	Globus pallidus 310
G (multiple of a unit) 373	interdigestive phase 240	Glomera aortica 132
g (membrane conductance) 32	neuronal and hormonal inte-	carotica 132
GABA (γ-aminobutyric acid) 34,	gration 234	Glomerular filter 148
52, 284, 320, 326 f.	neurotransmitters 234	zone, adrenal cortex 294,
cerebral cortex 332	passage time 232, 233 A	296
Gn-RH secretion 300	Gauer-Henry reflex 170	Glomeruli olfactorii 340
receptors 52, 55 F, 320	GCAP (guanylyl cyclase-activat-	Glomerulotubular balance
second messenger 55 F, 274	ing protein) 350	166
Galactorrhea 303	G cells, antrum 242	Glomerulus, renal 148
Galactose, intestinal absorption	GDNF (glial cell line-derived neu-	filtration 154, 156

258

renal reabsorption 158 Galanin 52, 86

rotropic factor) 338

GDP (guanosine diphosphate)

274, 276

pressure 150

rate (GFR) 152

GLP-1 (glucagon-like peptide =	Glutamate 34, 52, 174, 258, 274	Golgi apparatus 12, 286
enteroglucagon) 282,	AMPA-receptors 55 F, 336	cells, cerebellum 328
284	cochlear transmitter 366	tendon organs 316
Glucagon 230, 256, 272, 274, 282,	astrocytes 338	Gonadoliberin (GnRH) 230, 269,
284, 288	dehydrogenase, renal 176	280, 298, 300, 306
actions 283 A, 284, 285 C	genetic code 8	menstrual cycle 298, 300
gluconeogenesis 284	intestinal absorption 258	Gonadotropic hormone-releasing
influence on insulin release	long-term potentiation 336	hormone →
282	NH ₄ + excretion 176	gonadoliberin
lipolysis 256	NMDA receptor 52, 55 F, 336	Gonatropin-releasing hormone (GnRH) →
second messenger 274 secretion 284	receptor types 55 F second messenger 55 F, 274,	(GIRH) → gonadoliberin
	_	
Glucagon-like peptide (GLP-1) 230, 282, 284	276 taste quality 338	Gonads 294, 306 G proteins 37 C1, 55 F, 84, 274,
Glucocorticoids (→ also cortisol)	transmitter function 326, 336	276, 338
296	hair cells 342	adrenoceptors 84, 87 B
receptors 10, 296	photosensors 350, 354	subunits 274
synthesis 294, 296	Glutaminase, kidney 176	types, G _i 36, 82, 84, 274
Gluconeogenesis 72, 282, 284	Glutamine 176	G _o 36, 276
renal 148	astrocytes 338	G _{olf} 340
Glucose 72, 154ff., 172, 210, 228,	formation, hepatic 176	G _q 82, 276
258, 284	gluconeogenesis 282	G _s 84, 274, 340, 348
aerobic oxidation 72, 282	renal metabolism 176	G_t (transducin) 348
anaerobic degradation 142, 282	Glutathione 158, 160, 286	Graafian follicle 298
blood 282	conjugates 250	Gradient, electrochemical 26, 32,
caloric equivalent 228	secretion 160	44
value 228	conjugation 160, 250	Granular cells, cerebellum 328,
carrier 22, 26, 156, 158, 258,	S-transferase 250	340
262, 284	Glycerol 282, 284	olfactory bulb 340
chemosensors 282	Glycine 34, 52, 248, 255	Granular mesangial cells 184
co-transport with Na ⁺ 262	conjugation 250	Granules, secretory 268
deficiency 242	receptor 52, 55 F	Granulocytes 88, 104
energy substrate 72, 210	second messenger 55 F	basophilic 100, 254
glucagon secretion 284	transmitter function 320	eosinophile 94
intestinal absorption 258, 262	Glycocalyces 12, 14	neutrophilic 30, 94
metabolism 72, 282, 296	Glycogen 58, 72, 246, 282	allergy 100
influence of cyclic AMP 274	metabolism 282	immune defense 94
muscle 72	phosphorylase 274	migration 30
myocard 210	skeletal muscle 58, 73 B	production 94
osmotic diuresis 172	synthase 274	Granulosa cells 300
plasma concentration 282	synthesis 274	Granzyme B 98
pregnancy 304	Glycogenesis 282, 284	Graves' diseases 288
regulation 272	Glycogenolysis 72, 85, 274, 282,	Grey matter, spinal cord 310
production 72, 148, 282, 284	284	GRH → somatoliberin
renal reabsorption 156, 158	STH 280	Growth 282, 284, 288, 306, 330
storage 282	Glycolipids 14	factors 88, 232, 268, 278, 280,
uptake 86	Glycolysis 142, 282	338
Glucose-dependent in-	anaerobic 72	nerval 338
sulinotropic peptide	muscle 72	receptor types 268, 278
(GIP) 234	Glycoprotein 274, 286	hormone → somatotropin
Glucose-6-phosphate 41, 72	hormones 274	release-inhibiting hormone
Glucosuria 158, 172, 284	Glycosylation 12	(or factor) → somato-
Glucuronic acid 160, 250, 294	GMP, cyclic 268, 274, 278, 348,	statin
conjugates, carrier 250	350	releasing factor (or hor-
steroid hormones 294	photosensors 350	mone) → somato-
Glucuronides 156, 160	GnRH → gonadoliberin	liberin
Glucuronyl transferase 250	GNRP (guanine nucleotide-re-	influence of thyroid hormones
GLUT (Glucose transporter) 23,	leasing protein) 30	288
158, 258, 284	Goal-directed movement 328	insulin 284
GLUT2 158, 258	Goblet cells 244, 264	T_3/T_4 288
GLUT4 284	Goiter 272, 286, 288	GRP (gastrin-releasing peptide)
GLUT5 158, 258	Goldberger leads, ECG 196, 198	86, 230, 234, 240, 242

GRP second messenger 276	concentration, blood 138	buffer system 138
stomach 242	normal range 142	/Cl ⁻ antiport 124
GSC (glomerular sieving coeffi-	excretion 174	/CO ₂ buffer 140, 144
cient) 154	renal 142	CO ₂ transport, blood 124
GTP (guanosine triphosphate)	gradient 158	erythrocytes 124
274, 276, 278	Na+ antiport 26 ff., 161 C, 162,	excretion 138, 142, 144, 172,
transducin 348	174 ff.	176
GTPase 276	secretion, renal 174	high altitude 136
transducin 350	symport 158, 258	gastric mucosa 242
GTPase-activating protein (GAP)	uniporter (UCP) 222, 230	intestinal absorption 262
350	H+-K+-ATPase 26 f., 28, 174, 182,	loss, diarrhea 262
Guanine 8	242	plasma 142, 146
Guanine nucleotide-releasing	collecting duct 174, 182	production, amino acid metab-
protein (GNRP) 30	colon 262	olism 176
Guanosine	production 174	renal 145 B2
$diphosphate \rightarrow GDP$	renal excretion 174	renal reabsorption 156, 174
monophosphate, cyclic → GMP,	secretion, gastric 242	saliva 236
cyclic	renal tubular 174	secretion 242, 246
triphosphate → GTP	H+-ATPase 14, 26, 174	bile ducts 248
Guanosylmonophosphate →	lysosomes 14	pancreas 246
GMP	H ⁺ -peptide cotransport 26 ff.,	salivary glands 236
Guanylyl cyclase-activating pro-	158, 258	stomach 242
tein (GCAP) 350	H+-phosphate cotransport 17 B2	titration curve 147
Guanylyl cyclase, coronary arter-	H ⁺ -pyruvate cotransport 17 B2	hCS (human chorionic soma-
ies 210	H ₂ , large intestine 264	totropin) 304
cytoplasmatic 278	Habituation 336	HDL (high density lipoproteins)
retina 350	Hageman factor 102	254. 256
α-gustducin 338	Hagen-Poiseuille's equation 188,	estrogen effect 302
Gut 234ff.	390	Head's reflex 132
absorption, amino acids 262	Hair cells 58, 342	Head's zones 318
Ca ²⁺ 262	inner 364	Hearing (\rightarrow also sound) 362 ff.
Cl ⁻ 262	outer 364, 368	audibility limit 362
mechanisms 263 D	electromotility 366	auditory cortex 368
glucose 262	potential 366	pathway 368, 370
Mg ²⁺ 262	sensor potential 366	binaural 368
Na+ 262	vestibular ganglion 342	cochlear amplification 366,
mechanisms 263 D	follicle receptors 314	368
phosphate 262	Haldane effect 124, 126	direction 368
H ₂ O 262	Hamburg shift 124	threshold 368
mechanism 263 B		loss 364, 366
	Haptenes 100	sensors 364 ff.
bacteria 232, 250, 264	Haptoglobin 90 Haustration 264	
Brunner's glands 244		thresholds 362, 366, 368
calcitriol effect 292	Hay fever 100	Heart (→ also cardiac and myo-
CCK production 234	hBSEP (human bile salt export	card) 86, 186, 210
defecation 264	pump) 248	action potential 192, 193 A
gases 264	H (histamine) cells, stomach 242	afterload 202, 204
motility 244	hCG (human chorionic	all-or-none contraction 68, 192
structure 244	gonadotropin) 304	arrhythmias 180
Gynecomastia 294	hCL (human placental lactogen)	atrial contraction 190
Gyrus, angularis 370	304	flutter 200
cingulate 310, 330	HCl (hydrochloric acid) 142, 238	fibrillation 200
parahippocampal 330	gastric 238, 242, 258	septum defect 220
postcentral 322, 338	production in metabolism 174	tachycardia 200
	HCO ₃ ⁻ (bicarbonate) 124, 126,	atrioventricular block 200
	140, 146, 236, 262	node 192
Н	actual 146	valves 190
1 / 1 / 1 / 6 / 1 / 1	blood 142	automaticity 192
h (multiple of a unit) 373	concentration, actual 142	autonomic nervous system
H zone, muscle 62	measurement 146	194
H+ 138, 140, 142, 144, 242	normal range 142, 146	beats → heart rate
buffering 138, 140, 142, 144	standard 142	bundle of His 192
chemosensors 132	measurement 146	cholinergic transmission 83 B

conduction 192, 194, 200	blood flow 186	semilunar valves 190
disturbances 200	Ca ²⁺ -ATPase 194	sinoatrial node 192
system 192	Ca ²⁺ channels 194	rhythmicity 192
velocity 195 C	Ca ²⁺ concentration, cytosolic	threshold potential 192
contractility 194, 204	194	sounds 190, 191 A
influences 194, 296	contraction 59 A	stroke volume 186, 190, 202,
coronary blood flow 186, 190,	isovolumetric 190	204
212, 210	velocity 204	maximum 77 C
vessels 210, 212	dihydro-pyridine receptors	endurance, athlet 77 C
cycle 190	194	measurement 106
diastolic time 190	ECG 196	regulation 204
ejection fraction 190	electromechanical coupling	training 76
phase 202	194	ventricles, stretch sensors 214
electric activity, action poten-	frequency inotropism 204	ventricular diastolic volume
tial 58, 192	impulse generation 192	190
arrhythmias 180, 200	infarction 218	pressure 190
AV block 200	ECG 198	curve 191 A
AV rhythm 192	influence of Ca ²⁺ 194	work diagram 202
bundle branch block 200	contractility 194	volume load 204
cholinoceptors 82	inotropism 194, 204	weight, endurance athlet 77 C
chronotropism 194	ischemia, ECG 198	work 202
dromotropism 194	isotonic maxima 203	Heat 222, 228, 288
ECG 196	isovolumetric maximum 203	countercurrent exchange 164
ectopic 200	metabolism 210	flow, external 222
electrolyte disturbances	Na+/Ca2+ exchange carrier	internal 222
198	194	loss by evaporation of water 222
extrasystoles 200	Na+/K+-ATPase 194	of maintenance 74
impulse generation 192	O ₂ supply 210	production 222, 228, 288
disturbances 200	tension-time index 210	energy metabolism 228
inotropism 194, 204	mechanics 202	influence of thyroid hor-
pacemaker 192	metabolic activity 210	mones 288
diastolic depolarization	O ₂ supply 210	sensors (→ also warm sensors)
192	output, cardiac (CO) 74, 106,	314
ectopic 192	122, 186, 188, 218	unit 375
potential 192	fetus 220	Heat shock proteins (HSP) 278
reentry 194, 200	high altitude 136	Hecto- (multiple of a unit) 373
spreading time 195 C	maximum 77 C	Helicine arteries 308
vulnerable phase 193 A,	endurance, athlets 77 C	Helicotrema 364, 366
200	physical work 74	Helium 114
electrical axis 198	shock 218	Helium dilution, measurement of
end diastolic volume (EDV)	power 202	residual volume 114
190, 202	preload 202, 204	Helper cells, T- 98
systolic volume (ESV) 190, 202	pressure-volume diagram 68	Hematocrit 88, 150, 168
influence of aortic pres-	relation 202	blood viscosity 92
sure 204	work 202	high altitude 136
energy sources 210	Purkinje fibers 192	Hematopoiesis 88
exercise 74	rate 74, 136, 186, 194, 288	testosterone 306
failure 208, 216, 218	AV block 200	Heme 88, 128
cause of edema 208	blood pressure regulation 4,	Fe(II) 90
salt and water homeostasis	218	oxygenase 90
173 B	fetus 220	Hemeralopia → night blindness
filling phase 202	high altitude 136	Hemochromatosis 90
Frank-Starling mechnism 68	influence of Ca ²⁺ influx 194	Hemoglobin 88, 90, 92, 124, 126,
frequency → heart rate	maximum 77 C	128 , 138, 140, 146, 250
glycosides 194	endurance, athlet 77 C	blood puffer 124, 128, 138, 140,
influence of Ca ²⁺ 194	normal 186	146
inotropism 194, 204	physical work 74	carbamate 124
isovolumetric contraction 190,	shock 218	concentration, blood 128
202	T ₃ /T ₄ effect 288	degradation 250
muscle 46, 58, 68, 192 ff.	reentry 194	fetal 128
action potential 46, 59 A	reflexes 216	synthesis 10, 90
adrenoceptors 214	resting pressure-volume curve	types 128
	202	.J

hNBC (human Na+-bicarbonate Hemolysis 88, 100, 250 male sex 306 fetus 100 cotransporter) 162, mammotropic 269 prehepatic jaundice 250 174 medication 272 Hemopexin 90 H₂O (→ also extracellular fluid α-melanocyte-stimulating hor-Hemophilia 104 and body fluid). mone (α -MSH) 269 Hemoproteins 250 balance 148, 166, 168 menstrual cycle 300 Hemorrhage 218 disturbances 172, 173 B natriuretic 170 Hemorrhagic diatheses 104 clearance 164 nomenclature 269 Hemosiderin 90 concentration 24 pancreatic 282 Hemostasis 102 daily intake 168, 262 paracrine 266, 268 Henderson-Hasselbalch equation deficit 170, 180 peptide 268, 274 138, 139 A, 140, 146, diffusion 24 placenta 304 379, 389 diuresis 164, 166, 172 pregnancy 304 maximum 166 Henry-Gauer reflex 170, 218 principle functions 272 Heparin 104, 254 urea excretion 166 receptors 266, 268, 274 ff. Hepatic → liver excess 170 second messengers 274 ff. Hepatocyte growth factor (HGF) excretion, feces 262 steroid 248, 268 278 renal → kidney, excretion placenta 304 Hering-Breuer reflex 132 flow, osmotic 24 therapeutic administration 272 Hering's opponent color theory free, urine 164, 172 thyroid 286 homeostasis 168 stimulating (TSH) 269 354 Hertz, unit 362, 374 intake 168 tissue 268 HGF (hepatocyte growth factor) intestinal absorption 262 transport, blood 268 278 intoxication 170 types 268 hGH → chorionic growth horlosses 170 Horopter 360 Horse power (hp), metric, unit mone partial pressure 106 HLA protein (human leukocyte permeability 208 374 antigen) 96, 98 reabsorption 154, 164 5-HPETE (= 5-hydroperoxyeico-High altitude 130, 132 renal handling 166, 170 satetraenoate) 269 respiration 136 regulation 170 hPL (human placental lactogen) erythropoietin secretion 88 transport 24 hypoxic hypoxia 130 turnover 168 H₂SO₄ (sulfuric acid) production density lipoproteins (HDL) 256 Hoffmann's reflex 316 142, 174 pressure system 186 Homeostasis 2, 4, 212, 266 HSP (heat shock protein) 278 Hippocampus 330, 336 Homeothermy 222 H₁(istamine)-receptor, second Hippurate, transcellular secre-Horizontal cells, retina 344, 354 messenger 55 F, 212, Hormone(s) 234, 266, 268 tion, renal 156 Histamine 70, 100, 160, 208, 212, abbreviations 268, 269 H2-receptor, second messenger 214, 218, 234, 242, adrenocorticotropic (= ACTH) 55 F. 274 269, 274, 276 269 Hüfner number 128 allergy 100 aglandotropic 266, 280 Human chorionic gonadotropin anabolic 306 (hCG, HCG) 304 cerebral cortex 332 coronary vasodilatation 210 autocrine 266 somatotropin (hCS) 304 binding proteins 268 gastrointestinal tract 234 leukocyte-associated antigen gastric juice secretion 242 catecholamines 274 (HLA) 96 influence on vessel permeabilchemical structure 268 Na+-bicarbonate cotransporter down regulation 30 (hNBC) 162, 174 ity 214 endocrine 266, 268 receptors, types 55 F placental lactogen (hPL, HPL) H₁- 212, 276 feedback mechanism 272 H2-242, 274 femal sex 294, 298, 302 Humidity, air 222, 224 second messenger 274 follicle-stimulating (= FSH) 269 Humor, aqueous 344 shock 218 gastrointestinal 234 Hunger 142 tubular transport 160 gland, atrophy 272 edema 208 vasodilator 214 hypertrophy 272 energy reserves 282 Histidine 174, 226 influence of hormonal medimetabolic effect 283 A 3-HMG-CoA-reductase 256 Hydraulic filter, arteries 188 cation 272 glandotropic 266, 269, 280 Hydrocarbon continuum 252 HMK (high-molecular-weight kininogen) 102 glycoprotein 274, 268 Hydrocephalus 310 hNaDC-1 (human Na+-dicarboxyhierarchy 268 Hydrochloric acid → HCl late transporter) 160 hypothalamic 268, 280 $Hydrocortisone \rightarrow cortisol$ Hydrogen ion → H⁺ and pH lactogenic 269

412

lipophilic 268

peroxide 94

β-Hydroxybutyrate 142	salt and water homeostasis	1
1α-Hydroxylase, calcitriol syn-	173 B	
thesis 292	Hypervitaminosis 226	I band, muscle 62
11-Hydroxylase	Hypervolemia 178	ICAM (intercellular adhesion
adrenal cortex 294	Hypocalcemia 178, 198, 290, 292	molecule) 96
17-Hydroxylase	ECG 198	ICF → intracellular fluid
adrenal cortex 294	renal phosphate reabsorption	ICSH (= LH) \rightarrow lutropin
21-Hydroxylase	178	Icterus 250
adrenal cortex 294	Hypocapnia 144	IDDM (insulin-dependent dia-
24-Hydroxylase, calcidiol 292	Hypoglycemia 86, 284	betes mellitus) 284
17β-Hydroxysteroid dehydro-	glucagon secretion 284	IDL (intermediate-density lipo-
genase 300	Hypokalemia 172, 198, 262	proteins) 254, 256
11β-Hydroxysteroid oxidoreduc-	diarrhea 262	IEL (intraepithelial lymphocytes)
tase 182, 296	ECG 198	232
Hyperalgesia 318	vomiting 238	IgA (immunoglobulin A) 98, 232,
Hyperaldosteronism 182, 238	Hyponatremia 170	236, 344
Hyperaminoaciduria 158	Hypoosmolality, extracellular	IgE (immunoglobulin E) 93, 98,
Hyperbicarbonaturia, osmotic di-	fluid 172	100
uresis 172	Hypoparathyroidism 290, 292	IGF-1 (insulinlike growth factor)
Hypercalcemia 178, 198, 290, 292	Hypophosphatemia 290, 292	278, 280
ECG 198	Hypopnea 108	IgG (immunoglobulin G) 93, 98
renal phosphate reabsorption	Hypotension 206	IgM (immunoglobulin M) 93, 98,
178	orthostatic 182	100
Hypercapnia 144	Hypothalamus 78, 134, 170, 224,	Ileum (→ also intestine) 232, 244
Hypercholesterolemia 256	230, 266, 268, 280 ,	bile salt reabsorption 248
Hypercolumns, cerebral cortex	300, 310, 330, 358,	cobalamin absorption 260
358	359	Immune defense 94 ff., 232, 244
Hyperemesis gravidarum 238	afferents 330	antigen presentation 96
Hyperemia, reactive 212	angiotensin II 184	cellular 94, 232
Hyperglycemia 284	body weight 230	gastrointestinal 232, 244
osmotic diuresis 172	function 266	specific 98
Hyperkalemia 142, 180 ff., 198	hormones, regulation 268, 280	cellular 96
ECG 198	limbic system 340	humoral 98
Hyperlipoproteinemia 256	nuclei, arcuate 230	unspecific 94
Hypermagnesiemia 178	dorsomedial 230	response, delayed 96
Hyperopia 346	lateral 230	secondary antigen contact 100
Hyperosmolality, extracellular	paraventricular 224	Immunity 94
fluid 172	magnocellular 280	nonspecific 94
Hyperoxia 136	ventromedial 224	specific 98
	osmosensors 170	Immunization 100
Hyperparathyroidism 292 Hyperpnea 108	smell 340	active 94
	somatostatin secretion 286	
Hyperpolarization, action poten- tial 46	testosterone effect 306	passive 94
		Immunoglobulins (Ig) 93 f., 94, 98, 232
afterpotential 46	thermoregulation 222, 224 thirst center 168	
photo sensors 354	TRH secretion 286	class switching 98
Hyperprolactinemia 303		concentration, serum 93 D
Hyperreflexia 320	Hypothyreosis 288, 303	IgA 98, 232, 344
Hypersensitivity, delayed type	Hypoventilation 108, 144	saliva 236
100	Hypovitaminosis 226	IgE 93, 98, 100
immediate type 100	Hypovolemia 218	IgG 93, 98
Hypertension 184, 206, 216	chronic vomiting 238	IgM 93, 98
renal 184, 216	shock 218	AB0 system 100
resistance 216	Hypoxia 130 , 136, 142	function 94
Hyperthyreoidism 288	autoregulation 210	placental barrier 92
Hypertrophy, compensatory 272	brain 186	receptors 96
Hyperventilation 108, 120, 136,	fetus 220	infections 94
144, 290	vasoconstriction 122, 212, 220	Immunologic tolerance, central
diving 134	Hypoxic vasoconstriction, lung	94
high altitude 136	122	peripheral 98
H ₂ O losses 168	Hz, unit 374	Immunological memory 94
nonrespiratory acidosis 142		Immunosuppression 98

Impedance matching, middle-ear receptor 268, 278 Intestine 232 regulation of secretion 282 364 Cl- secretion 263 Implantation of fertilized ovum secretion 282 defense mechanism 232 amino acids 282 lymphatics 232 Impotence 303 β₂-adrenoceptors 84 mucosal surface 244 Inch. conversion into SI unit 372 GIP 234 small 240 Incus 364 inhibition 330 structure 244 Indicator dillution technique 168 synthesis 282 passage time 232 Indifference point, hydrostatic Insulin-like growth factors (IGFs) Intracellular fluid 168 204 278, 280 composition 92 Indirect bilirubin 250 Insulin receptor substrate-1 ions 93 C Indoor climate 224 (IRS-1) 278, 284 Intraesophageal leads (ECG) 196 Induction, hormon-dependent Integration of body functions Intrafusal fibers, muscle 316 266, 310 Intrapleural pressure 108 278 Infarction, myocardial 198 Intention tremor 328 Intrapulmonic pressure 108 Inflammation 96, 100, 269, 318 Intercalated cells, tubule 174, 182 Intrathoracic pressure 108 neurogenic 318 Intercellular adhesion molecule Intrinsic factor 260 (ICAM) 96 Information 312 cobalamin deficiency 90 encoding 312 Intercostal muscles 108 gastric juice 242 storage 336 Intercourse 308 Intron 8 unit 312 Interdigestive phase 240 Inulin, indicator for extracellular Inhibin 300, 306 motility 234 space 168 kidney, clearance 152, 154 FSH secretion 306 Interferon (IFN) 95 f. menstrual cycle 298 IFNy 95, 98 5-iodinase 286 Iodine/Iodide, butanol-ex-Inhibition, antagonistic 320 Interleukin (IL) 94, 224, 232 tractable (BEI) 286 ff. autogenic 316 IL1 296 competitive 28, 56 IL2 98, 296 daily requirement 226, 288 descending 318 11498 Ion channels 32, 34 feed-forward 320 II.5 98 control 34, 276 lateral 312, 368 IL6 98 with ligands 34, 55 F acoustic pathway 368 II 8 94 diffusion 22 postsynaptic 320 Intermediate filaments 14 open-probability 34, 46, 50 presynaptic 320 type, heart axis 198 pump (ATPases) 26 Ionic conductance 32, 375 recurrent 316, 320 Internal clock 240, 334 current 32, 375 Inhibitory postsynaptic potential heat flow 222, 224 intercostal muscles 108 Ions (→ also individual ions) 375 (IPSP) 52 milieu 168, 266 body fluid 93 C Inner ear 364ff. potentials 366 tissue respiration 130 concentrations 45 B IP₃ (inositol-1,4,5-trisphophate) Inositol-1,4,5-trisphosphate (IP₃) Interneuron 316, 320, 324 82, 84, 268, 274, 276 inhibitory 320 82, 274, 276 Inotropism, heart muscle 194, 204 stimulatory 320 IPPV (intermittant positive pres-Inspiration 108, 110, 204 Interstice 45 B, 92, 93 C, 168 sure ventilation) 110 pressure differences, driving Interstitial cell-stimulating hor-IPSP (inhibitory postsynaptic force 108 potential) 52 mone (ICSH) \rightarrow work 116 lutropin cerebral cortex 332 Inspiratory pressure, maximum fluid 92, 93 C, 168 peptidergic 82 116 Internode region 48 Iris 344 Inspired air, composition 107 A Iron (→ also Fe) Intestinal absorption (→ also ab-Instinctive behavior, limbic syssorption) 244 ff. Iron lung 110 IRS-1 (insulin receptor substratetem 330 amino acids 258 Insulin 180, 230, 254 ff., 256, 268, Ca2+ 262, 292 1) 278 272 ff., 274, 282 ff, electrolytes 262 Ischemia 130, 198 284, 288 glucose 258 Islets of Langerhans 282 anabolic effects 284 vitamins 260 Isoleucin 226 deficiency, diabetes 284 water 262 Isomaltase 258 bacteria 264 Isophones 362 degradation 282 effects 180, 283 A, 284, 285 C blood flow 232 Isoprenaline 84, 87 B half-life 282 glands 244, 262 Isotonic peak cuve 202 Isovolumetric contraction, heart K+ homeostasis 180 lymph 232, 254 lipoproteinlipase 254 movements 244 190 lipolysis 256 neurons 244 relaxation, heart 190 overdose 218 work diagram 203 phase in gastric secretion 242

homeostasis 180

peak curve 202	renal handling 156, 180 ff.	ammonia/ammonium 76,
IP ₃ (inositol-1,4,5-trisphosphate)	role in autoregulation 212	174
274, 276	secretion, intestine 262	Ca ²⁺ 178
	saliva 236	electrolytes 157 D
	tubule 172	fractional (FE) 152 f.
J	stomach, parietal cells 242 K _M (Michaelis-Menten constant)	H+ 142, 174 titratable acid 178
(Joule), unit 298, 374	28	
(Joule), difit 298, 374 [aundice 250	Kallidin 214	HCO ₃ 138, 144 Mg ²⁺ 178
ejunum (→ also intestine) 234,	Kallikrein 104. 214	organic substances 157 D,
244	salivary gland 236	158, 160
let lag 334	Karyocytes 88	phosphate 176 f.
[GA (juxtaglomerular apparatus)	Karyolymph 8	steroid hormones 294
172, 184	kcal, conversion into SI unit 374	urobilinogen 250
oint position, information on	Kelvin (K), unit 375	water 154, 157 D, 164
316	Keratin filaments 14	glucocorticoid effect 296
oule (J), unit 298, 374	Kerckring's fold 244	extraction fraction 152
sensors 132	α -Ketoglutarat, renal production	filtration 148, 152
unctions, tight 154	176	amount of a substances 158
uxtaglomerular apparatus (JGA)	renal transport 160	dissolved substances 154
172, 184	Ketosis 284	equilibrium 152
	17-ketosteroids 294, 302, 306	fraction (FF) 152
1/	synthesis 294	pressure, effective 152, 390
K	Kidney (→ also tubule and renal) 148 ff., 292	function 148 ff.
k (multiple of a unit) 373	acid base balance 138, 174	glomerular filtration rate (GFR) 150
K (dissociation constant) 378 f.	α_1 -adrenoceptors 214	glucocorticoid effect 296
K (Kelvin), unit 375	ADH receptors 166	glomerulotubular balance
K ⁺ , absorption 180	aldosterone effect 182	glomerulus 148 ff.
balance 180	anatomy 148, 149 A, B	glucocorticoid effect 296
in intestine 262	aquaporin (AQP) 166	glutaminase 176
adaptation 182	ATP 154	glutamine 176
balance 182	atriopeptin effect 152	H+/K+-ATPase 174, 181 B4
effect of aldosterone 182	autoregulation 150, 212	H+-ATPase 174
cellular uptake, influences on	range of 151 C	H+-excretion 142, 174
180	balance glomerulotubular 166	H+-secretion 174
channels, activation 32 ff.	blood flow 150, 186	handling of amino acids 158
adrenoceptors 84	measurement 150	angiotension 158
G-proteins 276	medullary 172	Ca ²⁺ 178, 292
ATP-controlled 282	Ca ²⁺ excretion 178, 292	diuretics 156
collecting duct 180	sensors 178	drugs 156
conductance, action poten- tial 44, 46	calcitriol synthesis 292 capillaries 148 f., 151 A, 164,	glucose 158 glucuronides 156
GABA _B receptors 320	184	glutathione 158
hair cells 342	pressure 151 B	H+ 144, 174
pacemaker cells, heart 192	hypertension 216	HCO ₃ - 142, 174
resting potential 44	circulation 150	K+ 156, 180 ff.
voltage-gated 46	clearance 152, 389	lactate 158
renal 162, 180	ratio of 152	Na+ 156, 162, 170
concentration, intracellular 26,	water 164	oxalate 158
44	concentration and dilution	peptides 158
contracture, skeletal muscle 66	164, 389	phosphate 176, 178, 290,
deficiency 172, 180	mechanism 164, 166	292
diffusion 32	conjugation process 160	proteins 158
distribution 284	connecting tubule 166	urea 166
equilibrium potential 44	cortex 164, 166	uric acid 156, 158
excretion 172, 180, 182	countercurrent exchange 164	inulin clearance 152, 154
diuretics 172	diuresis 172, 176	juxtaglomerular apparatus
influence 182	endopeptidase 158	184
intake, high 180 loss 262	energy metabolism 150 excretion	juxtamedullary nephrons 150 K+ channels 162
1033 202	CACICUUII	ix challifeld 102

outflow, motor end plate 56

Kidney loss 172	high-molecular-weight (HMK)	LCAT (lecithin cholesterol acyl
loop of Henle 164	102	transferase) 254
Ca ²⁺ -reabsorption 178	salivary gland 236	LDL (low density lipoproteins)
Mg ²⁺ reabsorption 178	Kinocilium 342	254, 256, 302
Na ⁺ transport 162	Knee jerk reflex 316	estrogen effect 302
water permeability 166	Kohlrausch's fold 264	receptors 254, 256
medulla 150	Kohlrausch break 352	L-dopa 84
medullary blood flow 166, 172	Korotkow sounds 206	Learning 336
metabolism 150, 282	Korsakoff syndrome 337	Lecithin 14, 248, 252
Mg ²⁺ sensors 178 Na ⁺ channels 172	Krebs cycle → citrate cycle Krogh's diffusion coefficient 22,	bile 248 cholesterol acyl transferase
Na*-K*-ATPase 154, 156, 162	120, 388	(LCAT) 254
function 154	Krogh's cylinder model 130	Left axis (heart) 198
nephron types 148, 166	Kupffer cells 96, 232	Lemniscus, medial 322
O ₂ consumption 150	kWh (kilowatt hour), unit 374	trigeminalis 322
PAH clearance 150	KVII (Kilowatt iloui), uliit 3/4	Length, units 372
pH homeostasis 142, 174		Lens 344, 346
plasma flow 150	L	Leptin 230
potential, transepithelial 156,	-	receptors 230, 330
162	Labia minora, glands 308	Leucine 226
PTH effect 178	Labyrinth 364	insulin release 282
reabsorption	basal, tubulus epithelium 154	Leu-enkephalin 234
Ca ²⁺ 172, 290	Labyrinth reflexes, tonic 328	Leukocytes 88
Cl- 162	Lacrimal canaliculi 344	interferon secretion 96
electrolytes 157 D	glands 344	Leukotrien 268, 269
fractional 154	sac 344	allergy 100
D-glucose 158	Lactacidosis 72, 76	second messenger 276
HCO ₃ - 174	Lactase 258	Lewis's response 214
K+ 180	Lactate 72, 73 B2, 74, 142, 174,	Leydig cells 302, 306
Mg ²⁺ 172	282	LFA 1 (lymphocyte function-as-
Na+ 162	concentration in plasma 76	sociated antigen 1) 98
collecting duct 182	glugoneogenesis from 282	LH → lutropin
driving force 156	muscle metabolism 72	LHRH → gonadoliberin
organic substances 157 D,	myocard metabolism 210	Libido 306
158	physical work 76	Lids 344
phosphate 178 PTH effect 292	renal reabsorption 156, 158 vagina 302	Light, adaptation 350, 354 sensors 344
water 154, 164	Lactation 292	stimuli 354
renal failure 142, 176	reflex 303	wavelength 356
vitamin D 292	Lactic acid → lactate	Light chain protein, myosin II 60
renin 184	Lactogenesis 303, 304	Lignin 226, 264
secretion 156	Lactose 258	Limb leads (ECG) 196
K+ 172, 180	Lamellipodia 30	Limbic system 310, 330
organic substances 160	L-amino acids → amino acids	α-limit dextrin 246, 258
solvent drag 162	Landolt rings 349 A	Limited capacity control system
structure 148	Langerhans cells 96	(LCCS) 336
tight junctions 154	Language 337, 370	Linear velocity, unit 374
transport processes 154ff.,	Lanosterol 294	Linoleic acid 226
155 B,C	Laplace's law 118, 188, 202, 210,	Lipase 246, 252, 256
tubuloglomerular feedback	390	acid 256
(TGF) 172	Large intestine 232, 264	gastric fundus 252
ultrafiltrate 154	Larynx 370	hepatic 254
urea 166	testosterone 306	in human milk 252
Killer cells, natural 94, 96 T-killer cells 98	Lateral geniculate body 356, 358	lingual 252
Kilo- (multiple of a unit) 373	lemniscus 368 signal flow, retina 354	non-specific 252 pancreatic 252
Kilowatt hour (kWh), conversion	Lateralization, sound 364	saliva 236
into SI unit 374	Latency, hearing 368	Lipid membrane 14
Kinase cascades 274	Law of mass actions 379	Lipids 252, 256
Kinesia 238	Laxatives 262	absorption 252
Kinesia 256 Kinesia 42, 58, 62	LCCS (limited capacity control	digestion 252
Kininogen 102, 214, 236	system) 336	distribution 254

storage 254	LTP (long-term potentiation) 336
synthesis 12, 13 F, 282	Luminous intensity, unit 372
Lipogenesis 282	Lumirhodopsin 348
Lipolysis 86, 222, 246, 256, 282,	Lung (→ also pulmonary) 106,
284	108, 110, 112
influence 256	acid base balance 138 ff.
insulin effect 284	alveolar contact time 141 B
STH effect280	blood flow 106, 122, 186
stimulation, thermoregulation	fetus 220, 221 B
222	bronchial, obstruction 120
Lipoproteinlipase (LPL) 254, 256	capacity, total (TCL) 112, 113 A
insulin effect284	capillaries 106
thermoregulation 222	blood pressure 122
Lipoproteins 254ff.	disease, obstructive 118
receptors 28, 254	restrictive 118
Lipostasis 230	edema 118, 120, 122, 132, 144,
•	172, 208
Lipoxygenase 269	
Liter, definition 372 f.	fetal 220
Lithocholate 248	function test, dynamic 112, 118
Liver 90, 160, 176, 232, 248, 250,	gas exchange 120 ff.
282	hypoxic vasoconstriction 122
acid-base homeostasis 144, 176	inflation 118
bypass 232	iron 110, 111 A2
calcidiol synthesis 292	O2 diffusion capacity 22
coagulation factors, synthesis	opening pressure 118
102	perfusion → lung, blood
conjugation process 160	flow122
damage, blood clotting 104	stretch receptors 132
excretory function 160, 248 ff.	surface tension 118
fatty 254	total capacity (TCL) 112
Fe homeostasis 90	ventilation/perfusion ratio
fetal blood cell formation 88	122
formation of 7-dehydro-	volumes 112 ff.
cholesterol 292	measurement 112 ff.
gluconeogenesis 282	Lung and thorax
glutamine formation 176	compliance 116, 118
glycogenesis 282	pressure-volume relationship
icterus 250	116
jaundice 250	Luteal phase 298, 300
metabolism 282	Luteinization 300
steroid hormones, degradation	Luteinizing hormone (LH) →
294	lutropin
urea production 176	releasing hormone (LTH) →
Load, filtered 158	
	gonadoliberin
Locus coeruleus 328	Luteotropic hormone (LTH) →
Logarithms, calculations 380 f.	prolactin
Longitudinal tubules, muscle 60	Luteotropin (LTH) → prolactin
Long-term potentiation (LTP) 336	Lutropin 269, 280, 294, 306
Loop, cortico-thalamo-cortical	menstrual cycle 298
326	peak 300
diuretics 172, 178	receptor, cells of Leydig 306
oculomotor 324, 326	secretion, pulsatile 298
skeletomotor 326	Lymph flow 208, 209 B
Loop of Henle 148, 164, 166, 178	intestinal 232, 254
Loudness 362, 370	nodes 88, 96
level 362	vessels 244
voice 370	Lymphocyte function-associated
Low density lipoproteins (LDL)	antigen 1 (LFA 1) 98
256	Lymphocytes 88, 94
Low pressure system 186	B- 94, 98
LRH → gonadoliberin	activation 98
$LTH \rightarrow prolactin$	clonal selection 98

differentiation 98 clonal deletion 94 expansion 94 selection 94 intraepithelial (IEL) 232 naive 94 T- 94 "armed" 94, 96 CD4 98 CD8 98 cytotoxic cells 98 clonal expansion 98 selection 98 differenziation 98 naive 96 T helper cells 98 T_{H1} cells 98 T_{H2} cells 98 T killer cells 94, 98, 232 receptor 98 Lymphocytopoiesis 88 Lymphokins, cortisol 296 Lysine 174, 226, 258 intestinal absorption 258 Lysis, bacterial 94 Lysosomes 12, 14, 26 f., 28, 286 Lysozyme 94, 96, 232 renal rabsorption 158 saliva 236 tears 344

M

Malleus 364

μ (submultiple of a unit) 373 M (multiple of a unit) 373 m (submultiple of a unit) 373 M line, muscle 60 α₂-Macroglobulins 104 Macrophages 30, 94, 96, 98, 224, 250 activation 96 break down of red blood cells hemoglobin degradation 250 immune defense 94, 96 iron metabolism 90 migration 30 respiratory tract 110 Macula densa 148, 172, 184 Maculae 342 Magnesium → Mg²⁺ Magnetoencephalography (MEG) 332 Maintenance heat 74 Major histocompatibility complex (MHC) 96, 98 Malabsorption, folic acid 90 Maldigestion, enzyme deficit 246 Malpighian bodies 148

Maltase 258 Megakarvocytes 88, 102 immunological 94 Meiosis 306 Maltose 246, 258 knowledge 336 Meiotic division, spermatocyte loss 336 Maltotriose 246, 258 Mammary glands 292 306 motoric 326 Mammatropic hormone → profirst, ovum 300 short-term 336, 368 lactin second, ovum 308 Menaguinone (Vitamin K2) 260 Manganese (Mn) 226 Meissner's corpuscles 314 Menarche 298 Mannitol, osmotic diuresis 172 Meissner's plexus (plexus sub-Menopause 298, 302 Mannose-6-phosphate 12, 14 mucous) 244 Menses 280, 298 f. Mannose-binding protein (MBP) α -Melanocortin (= α -MSH = α -Menstrual cycle 224, 298, 300, melanotropin) 230, 96 303 MAO (monoamine oxidase) 86 269, 280 body temperature 224, 298 MAP kinase (mitogen-activated Melanocortin receptor (MC4 hormonal control 300 protein kinase) 276 receptor) 230 interactions 300 Margination 94 Melanocytes 280 Menstruation 280, 298 f. Masking, sound 362 Melanocyte-stimulating Mercapturic acid 250 Masklike facial expression 326 Merkel's cells 314 $hormone \rightarrow$ Mass, units 374 Mesencephalon 310 melanotropin Mass actions, law 379 α -Melanocyte-stimulating Mesentery 244 Mass concentration 376 hormone (α-MSH) Messenger, first (→ also hormones) 268, 274 Mass movement, large intestine 230, 269, 280 264 Melanopsin 334 ribonucleic acid → mRNA Mast cells 104, 256 Melanotropin (α-MSH) 230, 269, second 268, 274 allergy 100 280 substances 266 Maturation, influence of thyroid release-inhibiting factor → third 268 hormones 288 melanostatin Metabolic alkalosis, 142 sexual 298, 306 releasing hormone → rate 226, 228 Matrix, extracellular 14 melanoliberin basal (BMR) 226 total 226 Maximal breathing capacity Melatonin 334 (MBC) 118 second messenger 274, 276 Metabolism, amino acids 158 Maximum diastolic potential Membrane bone 174, 292 (MDP) 192 apical 18 carbohydrate 282, 296 MBC (maximal breathing capacbasolateral 18, 162, 180 energy 72, 284 ity) 118 capacity, nerve 48 glucose 282, 296 MBP (mannose-binding protein) conductance 32 heart 210 iron 90 96 fractional 32 M cells, mucosal epithelium 232 electric properties 42, 52 lipids 254, 282 MCH (mean corpuscular function 2 muscle 254 permeability for K+ 46, 180 Metarhodopsin I 348 hemoglobin) 88 f., Metarhodopsin II 348, 352 89 C influence of cyclic AMP MCHC (mean corpuscular 274 phosphorylation 350 hemoglobin concenfor Na+ 46, 56, 162 Metastasization 30 tration) 88, 89 C postsynaptic 42, 50 Met enkephalin 234 MCV (mean corpuscular volume) potential 32, 44, 48 Methan, intestine 264 88.89 C hair cells 366 Methemoglobin (MetHb) 128 MDR1 (multidrug resistance prophotosensors 354 reductase 128 tein 1) 250 renal tubule 156 Methionine 174, 226 MDR3 250 smooth muscle 70 Methotrexate 260 Measles 94 presynaptic 42, 50 5-Methyltetrahydrofolic acid 260 Mechanosensors, skin 314, 316 proteins, glycosylation 12 Methopyrapone 294 Medial geniculate body 368 synthesis 12 Metopyrone 294 Mediators 268 structure 14 Mg²⁺ 178, 262, 274 Medications, bile excretion 248 transport 16 ff. absorption, intestine 262 Medulla, adrenal 272 active 26 f inhibition of Ca2+ channels, 50 oblongata 132, 310, 322 carrier-mediated 22 f. CNS 336 intracellular 16 circulatory "center" 214 excretion, renal 178 nonionic 22 plasma concentration 178 rhythm generator, respiration 132 paracellular 18 renal reabsorption 156, 172, vomiting center 238 passive 20 178 MEG (magnetoencephalography) potential, driving force 22, sensors, kidney 178 32 mGLU-receptors, second mes-Mega- (multiple of a unit) 373 Memory 336 senger 274, 276

multi-unit type 70

MgSO ₄ 248	Mitral cells, olfactory bulb 340	tupos 50
MHC (major histocompatibility	valves 190	types 58 Mouth-to-mouth resuscitation
complexes) 96	MLCK (myosin light chain kinase)	110
MHC proteins 96, 98	36	MPS (mononuclear phagocytotic
MI (cortex area) 324	MMC (migrating motor complex)	system) 96
Micelle, bile 248	240	MRF → melanoliberin
intestine 252, 260	mmHg, conversion into SI unit	MRH → melanoliberin
Michaelis-Menten constant (K _M)	374	mRNA (messenger ribo nucleic
28, 158, 383 f., 388	Modification, posttranscriptional	acid) 8
kinetics 28, 158, 383 f., 388	10	MRP2 (multi-drug-resistance
renal glucose tranport 158	posttranslational 12, 268	protein type 2) 160,
Micro- (submultiple of a unit) 373	Mol, unit 374 f. Molality 374 ff.	250
Microfilaments 14, 16	Molarity 374 ff.	MSH \rightarrow also melanotropin α -MSH (α -melanocyte-stimulat-
Microglia 96, 338	Mole, unit 374f.	ing hormone) 230,
Micrography 326	Molecular layer 326, 328	269. 280
α ₁ -Microglobulin, renal reab-	"weight", unit 374 f.	Mucoviscidosis 246
sorption 158	Molybdenum (Mo) 226	Mucus 110, 236, 244
β ₂ -Microglobulin, renal reabsorp-	2-Monoacylglycerol 246, 252	bronchial 110
tion 158	Monoamine oxidase (MAO) 86	cervical os 298
Micron, conversion into SI unit	Monoaminergic pathways sys-	intestine 242, 244, 264
372	tem 330	neck cells (MNC), stomach 240
Microtubules 14	Monocytes 88, 94, 96	saliva 236
Micturition 79 ff., 148	Monoiodotyrosin residue (MIT)	stomach 242
Middle ear 364	286	Müller's maneuver 116
MIF → melanostatin	Mononuclear phagocytotic sys-	Multidrug resistance protein 1
Migrating motor complex (MMC)	tem (MPS) 96	(MDR1) 160, 250
240	Monosaccharides (→ also glu-	3 (MDR3) 250
Migrating wave 366 Migration 30	cose) 258 Monooxigenases 250	Multi organ failure 218 Muramidase → lysozyme
phagocytes 94	Monosynaptic stretch reflex 316	Muscarine 82
MIH → melanostatin	Morning sickness 238	Muscle 42, 44, 46, 56 ff., 58, 59 A,
Mile, conversion into SI unit 372	Morphine 318	254
Milk 252, 260, 290	tubular transport 160	abdominal 108
Ca ²⁺ 290	Moss fibre, cerebellum 328	activity, heat production 224
ejection 280, 303	Motilin 240	ATP 72
fat 252	esophagus 238	afterloaded contraction 66
human, lipase 252	interdigestive motility 234	cardiac 46, 192 ff.
lactation reflex 303	secretion 234	ciliary 344
oxytocin 303	Motility, molecular basis 58	contractile machinery 60 ff.
production 303	Motion sickness 238	contraction 66
prolactin 303	Motivation, limbic system 330	afterloaded 66, 202
sugar (lactose) 258	Motor activity 56 ff., 316 ff., 324	auxotonic 66 ff.
Milli (submultiple of a unit) 373 Mineralcorticoids (→ also	basal ganglia 324 cerebellum 326 f.	isometric 66 isotonic 66
aldosterone) 182, 296	influence on circulation 74	role of Ca ²⁺ 64
production 294	pyramidal tract 324	dilator 344
Minerals, intake 226	voluntary motor function 324	energy supply 72, 254, 283
intestinal absorption 262	postural motor control 328	extensibility 66
nutrition 226	aphasia 370	titin 68
Miniature endplate current 56,	cortex 324 ff.	fatigue 76
57 B2	end-plate 56	fiber 60 ff.
Mini-pill 303	function, control center 326	force-velocity diagram 68
Miosis 344, 358 f.	supportive 326	hypertrophy 76
MIT (monoiodotyrosine residue)	neuron 42, 58	inspiratory 108
286	α- 320, 324	intercostal 108
Mitochondria 12, 288	γ- 316, 320	length-force curve 68
critical O ₂ pressure 130	paralysis 322	length, regulation 316
skeletal muscle 58, 63 A	proteins 58, 62	mechanical features 66
structure and function 12 T ₃ /T ₄ effect 288	system 324 unit 58, 66	metabolism 282 middle ear 364
13/ 14 CHECK 200	uiiit 30, 00	middle car Jua

thermal balance 222

recruitment 58

Muscle myofibrils 60	Myosin I 30	transport 44, 162, 170, 266
O ₂ extraction 72	Myosin-II 58, 60	paracellular 236
pump 204	striated muscle 60 ff.	tubular resorption 162
puporectal 264	smooth muscle 70	Na+-Ca ²⁺ antiport carrier 36, 194
reflexes 316		-kidney 178, 182
relaxants 56		-myocard 194
respiratory 108, 132	N	-photosensors 350
resting length 68		Na+-2 Cl ⁻ -K+-cotransport 236
force 68	n (submultiple of a unit) 373	inhibition 172
tension curve 66	N (newton), unit 374	kidney 162
single-unit type 70	N (nitrogen), balance 226	saliva 236
short-term high performance	N ₂ dissolved in plasma 134	$Na^+-K^+-ATPase (\rightarrow also ATPase)$
72 skeletal 56 ff.	role in diving 134	26, 28, 46, 180, 242,
	toxicity 134	288
smooth 70	Na+ 56, 158, 162, 170, 182, 184, 258, 378	cardiac glycosides 26, 194
action potential 59 A caldesmon 70	absorption in intestine 262	electrogenicity 46 hyperpolarizing afterpotential
contraction 70	antiport carrier 26f.	46
M ₃ -receptors 82	balance, effect of aldosterone	heart muscle 194
softening effect of ATP 64	182	ouabain 26
soreness 76	body content, total 170	phosphorylation 26
spindle 316	channels 46, 56, 172, 262, 348	renal collecting duct 180
sphincter 344	activation 46	tubule 154, 156, 162
stapedius 364	collecting duct 162, 180	resting potential 44
stiffness 76	conductance 46	salt reabsorption, intestine
striated (→ also skeletal	action potential 46	262
muscle, heart muscle)	inactivation 46	stomach, parietal cells 242
59 ff.	intestine 262	T ₃ /T ₄ effect 288
contraction, molecular	kidney 162, 172	transport cycle 26
mechanism 62	photosensors 348	Na+-taurocholate cotransporting
cycle 64	resting potential 46	polypeptide (NTCP)
summation of excitation 66	voltage-gated, heart 192	248
tension, regulation 316	Cl- symportcarrier 26	NaCl, homeostasis 170
tensor tympani 364	concentration, cytosolic 26, 44	disturbances 172
tone 66	co-transport 258	regulation 170
types of contractions 66	distribution 93	deficiency 170
weakness 56	excretion, renal 162	excess, counter regulation 170
Musculus bulbocavernous 308	feces 262	hypertension 216
ischiocavernous 308	/H+ antiport carrier (= Na+/H+	reabsorption, salivary glands
Myasthenia gravis 118 Mycobacteria, immune defense	exchanger, NHE) 26,	236 sense of taste 338
96	28, 162, 174, 176	
Mydriasis 344	exchange carrier → Na+/H+ antiport carrier	uptake 170 NADH 39 C
Myelin 42, 48	kidney 174	NaP _i (Na ⁺ -phosphate symport
sheath, nerve fibers 42, 48	PKC 276	carrier) 178
Myelopoiesis 88	stomach, parietal cells 242	nano- (submultiple of a unit)
Myelosis, funicular 260	influx, motor end plate 56	373
Myenteric plexus 244	intake, hypertension 216	Natriuretic hormone 170
Myocard → heart	renal reabsorption 156	Natural killer cells 94, 96
Myocardial function 190 ff.	handling 162	Nausea 238, 328
infarction 198, 218	retention 216	Near point, eye 322, 346
metabolism 210	saliva 236	sightedness 346
O ₂ consumption 210	symport carrier 26 f.	vision 344
oxygen supply 210	amino acids 258	response 359, 360
Myofibrils 60	bile salts 248	Necrosis 98
Myogenic tonus 70	Cl ⁻ 28, 162, 172, 236	Neck reflex, tonic 328
Myoglobin 58, 72, 90, 128	glucose 156 f., 258	Neocerebellum 326
Myometrium → uterus	HCO ₃ - (hNBC) 162, 174	Neostigmine 56
Myopia 346	intestine 262	Nephrine 148
Myosin 58, 60	iodide (NIS) 286	Nephrocalcin 178
light chain kinase (MLCK) 36	phosphate (NaPi) 178	Nephron 148, 150
smooth muscle 70	vitamins 260	cortical 150

juxtamedullary 150	dopaminergic 326, 330	renal cellular secretion 156,
sections 148	GABAergic 326	174 f.
structure 148	glutamatergic 326	renal excretion 174
transport processes 154	Ia- 316, 320, 324	secretion, renal tubular 176
Nernst equation 32, 44, 388	Ib- 316	NH ₄ ⁺ 145 B2, 174 ff.
Nerve (→ also neuron and ner-	II- 316	excretion 176 ff.
vus) 78, 84, 310 antidromic conduction 48	instestinal 244	nonionic transport 174
cell 42	internal longitudinal resistance 48	production 176 NHE3 → Na⁺/H⁺Antiportcarrier
cholinergic 78, 82	membrane capacity 48	Niacin, intestinal absorption 260
conduction velocity 48	α-motor 320	Niacinamide 226
endings, free, smell 340	γ-motor 320	Nickel (Ni) 226
fiber 42, 48	motoric 324, 328	Nicotine 82
diameter 42, 49 C	neurosecretory 280	Nidation 298, 303
myelinated 42, 48	nitrogenergic 278	NIDDM (non-insulin dependent
unmyelinated 42, 48	parasympathetic 78, 82	diabetes mellitus) 284
fibers Ia 316	postganglionic 78, 82	Night blindness 226, 350, 352
Ib 316	preganglionic 78, 82	Nipples, erection 308
Iia 316	sensoric 312 ff.	NIS (2 Na+-Isymport carrier)
glossopharyngeal 132	serotoninergic 330	286
neurosecretory 280	soma 42	Nitric oxide → NO
optical 344	summation, spatial 52	synthase (NOS) 82, 278
pelvic splanchnic 308	temporal 52	Nitrogen \rightarrow N and N ₂
stimulation 50	structure 42, 43 A1	N _M -receptors 82
structure 42	sympathetic 78, 82, 84 terminal buttons 42	N _N -receptors 82
trigeminal 318, 322, 340, 359 Nerve growth factor (NGF) 42,	action potential 48	NO 74, 86, 94, 210, 212, 214, 238, 278
278, 338	transmission, electrotonic 48	coronary vasodilatation 210
Nervous system 42	visceral afferent 78, 234, 266	erection 308
autonomic 78 ff., 80/81 A, 82,	Neuropeptid Y (NPY) 84, 86, 230,	immune defense 94
266	280, 300	synthase 82, 278
centers 78	cotransmitter 84	Nociception 318, 320, 322
cotransmitter 86	receptor types 55 F	tract 318
innervation, organs 78	second messenger 55 F, 274	Nocisensors 318, 322
peripheral 266	Neurosecretion 280	Nodal point, eye 346
central 266, 310	Neurotensin 55 F	Nodes of Ranvier 42, 48
enteric 234, 266	second messenger 276	Noise suppression, auditory
somatic 266	Neurotransmitter 42, 50, 52, 55 F,	pathway 368
Nervus opticus 344, 359	234, 266	Non-bicarbonate buffers 126,
vagus 132, 242, 248, 370	autonomic nervous system 78 ff.	138, 140, 144f.
Net diffusion 20 Neuroendocrine system 266 ff.,	excitatory 52	Non-ionic diffusion 22, 156, 176 Norepinephrine (→ also cate-
280 ff.	exozytosis50	cholamines) 52, 78,
Neurofibrils 42	function 42	84, 194, 230, 236, 268,
Neurofilaments 14	inhibitory 52	280, 300
Neurogenic tonus 70	ionotropic 34, 55 F	adrenal medulla 86
Neurohypophysis 268, 280	metabotropic 34, 55 F	cerebral cortex 332
Neuron (→ also nerve) 42, 50	release 50	coronary vasodilatation 210
Αδ 318	re-uptake 52	extraneuronal uptake 86
adrenergic 78 ff., 330	termination of action 54 E	heart 194
axolemma 42	Neurotubuli 42	inactivation 86
axon 42	Newborn 93 D, 94, 118, 128, 136,	insulin secretion 282
axon hillock 42	220, 224, 250, 288,	intestinal tract 234
axonal transport 42	305	neurons 330
Ca ²⁺ conductance 44	distress syndrome 118	pheochromocytoma 216
cholinergic 78 ff.	Newton (N), unit 374	receptor types 55 F release 84
Cl ⁻ -conductance 44 collaterals 42	NF-αB (necrosis factor) 276 NGF (nerve growth factor) 42,	release 84 re-uptake 86
conduction velocity 48, 49 C	278, 338	saliva secretion 236
cortex area 332	NH ₃ 22, 156, 174, 176	second messenger 55 F, 274,
dendrites 42	diffusion 22	276
diameter 49 C	production 176	synthesis 84

Norepinephrine thermoregula- tion 222	0	radicals, immune defense 94 role in autoregulation 212
NOS (nitric oxide synthase) 82,	Ω (ohm), unit 375	saturation (S_{0_2}) 126, 128
278	O ₂ (Oxygen) 72, 106	fetus 220, 221 A
NPY (Neuropeptid Y) 84, 86, 230,	arterio-venous difference 132,	influence on CO ₂ binding
280, 300	210	curve 126
cotransmitter 84	in myocard 210	solubility coefficient 128
receptor types 55 F	artificial respiration 132	in plasma 126
second messenger 55 F, 274	binding curve	supply, fetus 220
NREM (non-REM) sleep 334	in blood 126, 128	myocard 210
NTCP (Na+ taurocholate	fetal hemoglobin 220, 129 C	therapy 136
cotransporting poly-	myoglobin 129 C	toxicity 134, 136
peptide) 248	breathing 136	diving 136
Nuclear bag fibers 316	capacity of blood 126, 128	transport in blood 128
chain fibers 316	chemosensors 132, 136	uptake, exercise 74
envelope 10	concentration, blood, maxi-	maximum 72, 76, 77 C
pores 10	mum 128	endurance athlet 77 C
function 16	difference, heart muscle 210	OAT1 (organic anion transporter
signal sequences 10	consumption 106, 120, 150,	type 1) 160, 268
receptors, calcitriol 292	228, 288	Obesity 230
Nucleolus 8, 10	heart muscle 210	OCT (organic cation transporter)
Nucleotides 8	maximum 74, 76	160
Nucleus/nuclei, accessory 368	measurement 112, 120	Ocular chamber 344
amygdaloid 330	organs 130, 187 A	muscle, nuclei 342
anterior olfactory 340	renal 150	pressure, internal 320
arcuate 230 caudate 310, 326	T ₃ /T ₄ effect 288 debt 72	Oculomotor control 326, 329, 342
cuneatus 322	deficiency 72,130, 136, 210, 212	Ocytocin → Oxytocin
Deiter 326, 328	ventilation 136	Odorant molecules 340
dentate 328	demands 72, 74, 130	OFF-bipolar cells 354
dorsomedial 230	heart muscle 210	field (central) 354
emboliformis 328	increased 130	ganglion cells 354, 358
fastigial 328	organ difference 130	OH ⁻ -Ionen 138 ff.
gracilis 322	difference, alveolar-arterial 122	Ohm (Ω) , unit 375
lateral 230	arteriovenous 74, 106, 130	Ohm's law 32, 116, 188
lemniscus 368	diffusion 21 A, 130	ion transport 32, 388
localisation signal 10	lung 22	circulation 188, 388
olivaris superior lateralis 364	tissue 130	ventilation 116
paraventricular 230	dissociation curve 128	Oil-and-water partition coeffi-
pedunculus pontinus 326	fetal hemoglobin 220	cient 20
RNA synthesis 8	dissolved in plasma 128	Olfactory epithelium 340
red 328	extraction 72 f., 130, 210	pathway 340
septal 330	coefficient 210	sensor cells, primary 340
subthalamic 326 suprachiasmatic (SCN) 334	heart muscle 210 exercise 74	tract 340 Oligodendrocytes 42, 338
thalamic, anterior 330	organs, difference 130	Oligopeptides, digestion 258
tractus solitarii 338	skeletal muscle 72	renal handling 158
ventromedial 230	half saturation pressure 128	Oligosaccharides 258
vestibular 342	high altitude 136	Oliguria 164
Nuel's spaces 364	Krogh's diffusion coefficient	shock 218
Nursing mother, Ca ²⁺ require-	22, 120	Olive inferior 328
ment 290	partial pressure 20	superior 368
Nutrition 226	alveolar 120, 128	Oncotic pressure 24, 94, 152, 166,
integration 234, 330	diving 134	208, 378
passage time, stomach 233, 240	arterial blood 128	influence on capillary fluid
reflex 320	chemical respiratory stimu-	exchange 208, 379
vegetarian 142	lant 132	plasma 152
Nystagmus 360	critical, mitochondria 130	One-half maximum velocity con-
caloric 342	high altitude 136	stant (K _M) 28
optokinetic 360	hyperventilation 120 mixed venous blood 120, 128	ON-bipolar cells 354
pathological 360	plasma 132	field (central) 354
postrotatory 342	piasiild 132	ganglion cells 354, 358

Oocyte stage, primary 298	Osteolysis, malignant 292	ventricular 192, 200
Oogenesis 298	Osteomalacia 292	intestine, motility 244
Oogonia 298, 306	Osteoporosis 302	potential, heart 192
Open-probability, ion channels	Otoliths 342	stomach 240
34	Ouabain 26, 170	Pacinian corpuscles 314
Open system, thermodynamics 40	Ounce, conversion into SI unit 374	PAF (platelet-activating factor) 100, 102
Opioids 274, 280	Outer ear 364	PAH → p-aminohippurate
endogenous 318	Oval window 364, 366	Pain 318, 320, 322
exogenous 318 Gn-RH secretion 300	Ovaries 254, 268, 294, 298, 306 HDL receptor 254	assessments 318
receptor types 55 F	menstrual cycle 298	components 318 post-exercise muscle ache 77 D
gastointestinal tract 234	pregnancy 304	related behavior 318
second messenger 55 F, 274	production, fertiliziable egg	Palaeocerebellum 326
Opponent color channel 356	298	Pallidum 310, 326
Opsin 348, 350	testosterone production 306	Pancreas 232, 246
Opsonization 94, 96	Overshoot, action potential 46	cell types 282, 284
Optic chiasm 358	Overtone 362	enzymes 246
nerve 358	Overweight 226, 230	exocrine 246
lesion 358	OVLT (organum vasculosum	gastrin release 252
tract 354, 358	laminae terminalis)	hormones 282, 284
Optical apparatus 344	280, 310, 330	islets 268
nerve 344	Ovulation 298, 300, 302	juice 246, 252
system, simple 346	antiovulatory effect 303	somatostatin 284
Optokinetic nystagmus 360	inhibitors 300	Pancreatic lipase 252
Orexin 230 Organ of balance 360	Ovum 298, 302	necrosis, acute 246
Corti 364	first meiotic division 300 second meiotic division 308	polypeptide 282 secretions 246
Organelles 8	implantation 298, 303	Pantothenic acid 226
Organs, blood flow 187 A, 212,	Oxalate 102, 158, 160, 262	Papilla nervi optici 344, 358
214	inhibition, blood clotting 102	Para-aminohippurate (PAH) 150,
fetus 221 A	renal secretion 158, 160	160
transplanted 98	Oxidation, biological molecules	Paracellular transport 154
Organum vasculosum laminae	41	Paracrine action of hormones
terminalis (OVLT) 280,	of glucose, aerobic 72	282
310, 330	β-oxidation 256	Paraflocculus 326
Orgasm 308	2-oxoglutarate 176	Parallel fibres, cerebellum 328
Orgasmic cuff 308	β-oxybutyric acid (284	Paralysis, dissociated 322
Ornithine 258	Oxygen \rightarrow O ₂	Paraplegia 320, 328
Orthopnea 108	Oxygenation 128	Parasites, defense against 94
Orthostasis 5 C, 6 f., 204, 216	Oxyntic cells → parietal cells	Parasympathetic fibers, genital
Orthostatic hypotension 182 reflex 7 E, 204, 216	Oxytocin 269, 274, 280, 303, 304 receptors 55 F	tract 214 heart 194
Oscillation, unstable 6	second messenger 55 F, 274,	innervated organs 82
Osmol 377	276	ganglia 78 ff.
Osmolality, blood pressure 92	uterus 304, 308	nervous system, gastrointesti-
unit 377		nal tract 234
urine 164, 172		saliva secretion 236
Osmolarity 377	P	stimuli, salivary glands, blood
saliva 236		flow 214
unit 377	$P \rightarrow progesterone$	Parathormone → parathyrin
Osmometer 377	P _{0.5} (half saturation pressure) 128	Parathyrin (PTH) 36, 37 C3, 178,
Osmoregulation 170	P _A (alveolar pressure) 108	290
Osmosensors 170, 272, 330	P sensors 314	calcitriol 292
Osmosis 24	P (multiple of a unit) 373	chemistry 290
Osmotic coefficient 377 ff. diuresis 172, 176	p (submultiple of a unit) 373 P wave, ECG 196	deficiency 290 effects 290
effect on K+ excretion 182	Pa (Pascal), unit 374	influence on Ca ²⁺ and
pressure 24, 377	Pacemaker 70, 240	phosphate excretion 178
colloidal 92, 377	heart 192, 194, 200	regulation 290
Ossicles auditory 364	artifical 200	renal Ca ²⁺ reabsorption 178

Osteoclasts 290, 292

tertiary 192

Parathyroid glands , 268 290	Periglomerular cells 340	blood buffer 138
hormone → parathyrin	Perilymph 364, 366	calcium complex former 178
Paravertebral ganglionic chain	Perimeter 358	concentration, serum 290
78	Peripheral resistance 188, 206	deficiency 178, 292
Parkinson's disease 326	influence on heart function	DNA 8
Parietal cells, gastric 240, 242	206	excess 178
Parotid glands 236	Peristalsis 234	excretion 176, 178, 292
Pars recta, renal tubule 148	esophagus 238	H ⁺ secretion 176
Partial pressure 106, 120	intestine 244	homeostasis 290
Dalton law 106	large 264	intake 290
Parvalbumin, muscle fibers 64	stomach 240	intestinal absorption 292
Pascal (Pa), unit 374	ureter 148	metabolism 290
PAS domains 334	Permeability, Ca ²⁺ 82	plasma 290
Passive immunization 94	Cl- 44	renal reabsorption 158
Patch-clamp technique 34	coefficient (P) 22	solubility 290
Pause, compensatory 200	K+ 44, 56, 82	Phosphadidyl choline (lecithin)
post-extrasystolic 200 PBI → iodine	Na+ 44, 56, 82	14, 248, 252 bile 248
PCT (proximal convoluted	Peroxidase thyroid (TPO) 286 Peroxisomes 14	
tubule) 164	Perspiratio insensibilis 222	inositol-4,5-bisphosphate (PIP ₂) 276
PDGF (platelet-derived growth	Perspiration 222	Phosphatidylethanolamine 14
factor) 102, 278	water losses 168	Phosphatidylserine 14
PD-Sensors 312, 314	Pertussis toxin 276	Phosphaturia 178
Peak expiratory pressure 116	Peta- (multiple of a unit) 373	Phosphodiesterase 276, 348, 350
inspiratory pressure 116	Peyer's patches 232	cGMP-specific 278
Pendular movements, intestine	pH 138, 146, 378	Phospholipase A2 246, 252, 268,
motility 244	blood 132, 138	276
nystagmus 328	buffer 138	Phospholipase Cβ (PLCβ) 37 C1,
Penicillin, tubular secretion	normal range 142	82, 84, 276
156	cerebrospinal fluid 126, 132	Phospholipids 252, 254
Penis 308	clearance, esophagus 238	blood clotting 102
erection 278	erythrocytes 126	lipoproteins 254
REM sleep 334	esophagus 238	cell membrane 14
Pepsin 238, 242, 258	homeostasis, kidney 174	Phosphoric acid (→ also
Pepsinogens 242, 258	role of liver 176	phosphate) 142, 174
PepT1 (peptide transporter 1),	influence on diffusion 22	Phosphorylation 274
intestine 258	on protein bound Ca ²⁺ 290	Photochemistry, eye 348
PepT2, kidney 158 Peptidase 158, 242, 246, 258	measurement 146 plasma 126, 136, 138, 142	Photosensors 344, 346, 348 retinal distribution 348
Peptide(s), catabolism, renal	K ⁺ metabolism 180	sensor potential 354
148	saliva 236	Photopic vision 348
carrier PepT1 258	tubule lumen 174	Phyllochinone 260
PepT2 158	urine 156	Physical activity 72 ff.
digestion 258	Phagocytes 94	energy reserve 282
hormones 268, 274, 304	Phagocytosis 12, 28, 96	exercise capacity 76
placenta 304	Phase, vulnerable, heart 200	core temperature 224
messenger 274	Phenol red, tubular secretion	energy supply 226
renal reabsorption 156, 158	160	heat production 222
handling 128	Phenprocoumon 104	measure 74
transmitter 55 F	Phentolamine 84	measurement 76
Peptide-H+-symport carrier 28,	Phenylalanine 226	work 72 ff., 142, 282
158, 258	Phenylephrine 87 B	activation of the sympathetic
PER 334	Phenylethanolamine-N-methyl-	nervous system 74
Perception, visual 312 color 356	transferase 84 Pheochromocytoma 216	heat production 222
form 314	Phon 362	O ₂ consumption 74 threshold, aerobic 76
shape 314	Phosducin 350, 352	anaerobic 76
spatial nature 314	Phosphatase 276	ventilation 74
Perfusion, cerebral 186	alkaline 250	respiratory control 74, 133
pulmonary 120	Phosphate 138, 158, 175, 178,	A5
imbalance 120, 122, 130	290, 292	unit 374
Performance limit 76	absorption, intestine 262	Physiologic integration 266

Phytin 262	activating factor (PAF) 100, 102	maintaining 324
Pico- (submultiple of a unit) 373	activation 102	Potassium → K ⁺
PIF → prolactostatin	aggregation 102	Potentia coeundi 306
Pigmented epithelium 344	inhibitors 104	Potential, action 46 ff., 48
PIH (prolactin-release inhibiting	derived growth factor (PDGF)	unit 375
hormone) →	102, 278	difference, inner ear 366
dopamine	PLC_{β} (Phospholipase C_{β}) 37 C1,	diffusion 44
Pineal body (= pineal gland =	82, 84, 276	electrochemical 32
epiphysis) 334	Plethysmography 114	endocochlear 366
Pinocytosis 28	Pleura 108	end-plate 56
PIP ₂ (inositol-4,5-bisphosphate)	pressure (P _{pl}) 108	equilibrium 44
276	Plexus myentericus (Auerbach)	excitatory postsynaptic (EPSP)
Pirenzepine 82	234, 244	52, 56, 320
Pituitary gland 280, 286	submucous (Meissner) 234,	inhibitory prostynaptic (IPSP)
anterior 268, 280	244	52, 82, 332
influence of neurotrans-	PMA (premotor area) 324	maximum diastolic, heart
mitters 280	Pneumothorax 110	pacemaker 192
TSH secretion 286	diving 134	resting membrane 44
TRH receptors 286	types 110	reversal 47 B, 56
hormones 268	Podocytes, glomerulus 148	threshold 46, 48
posterior 180, 268, 280	Poikilothermy 222	transepithelial 156, 180
ADH secretion 170	Point, low 72	lumen-neagtive (LNTP) 162,
hormone secretion 280	Polkissen 184	236, 262
testosterone effect 306	Polyethylene glycol, intestine 262	lumen-positive (LPTP) 162,
pK _a value 138, 140, 378 f.	Polypeptide, pancreatic 282	172, 178, 180
PKA (proteinkinase A = A kinase)	Polyribosomes 10	Pound, conversion into SI unit
84, 274	Polysaccharides, chemical struc-	374
	ture 227 B	Power, unit 374
PKC (proteinkinase C) 36, 37 C1,	digestion 258	Powers of ten, calculation with
70, 84, 276	Polysomes 10	380 f.
PKG (proteinkinase G) 278 PKK (prekallikrein) 102	Polysynaptic reflexes 320	PP (pulse pressure) 194, 206
Placenta 100, 220, 292, 302, 304		ppb (parts per billion), unit 376
	Polyuria 164	
Placental barrier, function 304	POMC (pro-opiomelanocortin)	ppm (parts per million), unit 376
hormones 304	230, 280	P _{pl} (pleural pressure) 108
immunoglobulins 92	placenta 304	PQ interval, ECG 196
transfusion 220	Pond, conversion into SI unit 374	PQ segment, ECG 196
Plasma 88, 92	Pons 132, 310	Prazosin 87 B
albumin 92	eye movement 360	Prealbumin, thyroxin-binding
cells 94, 98	Pontocerebellum 326	(TBPA) 288
CO ₂ 124	Porphyrin 128	Precapillary sphincter 188
components 88, 92	Portal circulation 232, 248	Pregnancy 90, 260, 290, 292, 300,
factors 100	hypothalamus 280	302
globulins 92	vein 208, 232	Ca ²⁺ 290
osmolality 92, 171	Positive pressure ventilation	central venous pressure 204
proteins 88, 92, 154, 158, 377 f.	continuous (CPPV) 110	hormon concentrations 302
binding 24, 25 C, 88, 154	intermittent (IPPV)	hormonal control 304, 330
Ca ²⁺ 178, 290	Postcentral gyrus 322	nausea 238
blood puffer 138	Posterior funiculus, nuclei 322	Rh system 100
function 88, 92, 377 f.	Postextrasystolic pause 200	tests 294, 304
types 92	Postsynaptic inhibition 320	vitamin D-binding protein 292
pH 126	membrane 42, 50	vomiting 238
thromboplastin antecedent	Posttranslational modification	Pregnanediol 294, 303, 304
(PTA) 102	10, 12	Pregnenolone 294, 303
volume 170	Posttransscriptional modification	17-OH 294
measurement 168	12	Prekallikrein (PKK) 102
salt deficiency 170	Postural control 328	Preload, heart 202, 204
Plasmin 104	motor function 324, 326, 328,	Prepotential, heart 192
Plasminogen 104	342	Preproinsulin 282
Plasticity of smooth muscle 70	system 324	Presbyacusis 362, 366, 370
pyramidal cells 332	reflex 328	Presbyopia 346
Platelet(s) (→ also Thrombo-	labyrinthine 328	Presentation of antigens 96
cytes) 88, 102	Posture 324, 328	Pressure (unit) 374

Pressure arterial sensors 214 receptor 278 energy supply 228 filtrability, glomerulus 154 capillary 208 release inhibiting hormone → central venous (CVP) 190, 204 prolactostatin 268 ff.. functional minimum 226 280, 298, 303 colloidal osmotic 378 hormones 268 diuresis 170, 172 secretion, pulsatile 298 kinase A (PKA) 84, 274 hydrostatic 208 TRH effect 303 kinase C (PKC) 36, 37 C1, 70, 84, feet 208 Prolactostatin (PIH = dopamine) 276 internal ocular 344 268 ff., 280, 298, 303 kinase G (PKG) 278 intrapleural 108 Proliferation 272 kinase II (calmodulin-dependintrapulmonic 108 lymphocytes 94 ent protein) 36, 50 intrathoracic 108 Proliferative phase, menstrual kinase, tyrosine specific 282 oncotic 24, 208, 166, 378 cycle 298 minimum intake 226 peritubular capillaries 166 Prolipase 246 nuclear, nuclear pores 10 plasma 152 Pro-opiomelanocortin (POMC) nutrition 226 osmotic 24, 378 230, 280, 304 phosphorylated 274 Prophospholipase A2 252 transmural 188 plant 226 ventricular 190, 202 Proportional sensors 312, 314, plasma 92, 158 volume curve, heart 68 316 renal reabsorption 156 Proprioception 314, 316. 322 lung and thorax 116 synthesis 10, 13 F wave, pulse wave 190 Propriosensors 316, 324 influence of cyclic AMP 274 Pressosensors, arterial 314 neck 328 Proteinuria 158, 208 Prostacyclin (PGI₂) 104, 214, 268, Proteolysis 246 Presynaptic inhibition 320 Prothrombin 102, 104 membrane 42, 50 Prostaglandin(s) (PG) 162, 232, Prothrombinase 104 Pretectal region 359 234, 242, 268, 274, Protons -> H+ Previtamin D 292 Proximal stomach 240 Primary, response, antigen conautoregulation 212 P-Sensors 312, 314, 316 tact 94 E₂ 214, 318 Pseudohypoparathyroidism 290 saliva 236 fever 224 Psychotropic drugs 330 urine 164 effects 269 PTA (plasma thromboplastin Principal cells, kidney 162, 180 F_{2a} - 214 antecedent) 102 point, optical apparatus 346 fetal circulation 220 Pteroylglutamic acid 260 PRL → prolactin HCO₃- secretion, stomach 242 Pteroylpolyglutamate hydrolase Proaccelerin 102 I₂ (Prostacyclin) 104, 214, 268, 260 PTH → parathyrin Probenecid 160 Procarboxypeptidase 246 coronary vasodilatation 210 Ptyalin 236, 258 Process, ciliary 344 intestine 262 Puberty 302 Procolipase 246, 252 prostate 308 spermatogonia 306 Proconvertin 102 Puborectal muscles 264 second messenger 274, 276 Pudendal nerve 264 Procreative capacity 306 synthesis 269 Proelastase 246 inhibition 269, 318 Pulmonary (→ also lung) artery Progesterone 132, 294, 298, 300, uterus 304, 308 122, 186, 190 303, 304 Prostate 306, 308 partial pressures 122 actions 292, 303, 304 Protanomaly 356 pressure 120, 190, 206 chemistry 303 Protanopia 356 blood flow 122 degradation 303 Proteases, pancreatic juice 246, capillaries 106, 122, 124 esophagus sphincter238 258 circulation 186, 188 menstrual cycle 298 Protective reflexes 238, 320 resistance 206 17-OH-294 Protein C 104 edema 118, 120, 122, 132, 144, placenta 304 S 104 172, 208 plasma concentration 302 Protein 154, 148, 226, 228 infarction 120 transport 303 absorption, intestine 258 stretch receptors 132 production 294, 303 binding 24, 154 valves 190 respiration 132 bound iodine 286, 288 ventilation, exercise 74 secretion rate 303 caloric equivalent 228 ventilation/perfusion ratio 122 Proglucagon 284 capillary permability 208 Pulse pressure (PP) 190, 206 intestinal 282 catabolism, renal 148 receptors 214 Proinsulin 280, 282 chemical structure 227 B wave, velocity (PWV) 190 Prolactin 268, 269, 274, 280, 303, concentration in cerebrospinal Puncts, lacrimal 344 306 fluid 144 Pupilla 344, 346 lactation reflex 303 digestion 246, 258 contraction 344 menstrual cycle 298 enzymes 242, 246 dilatation 344, 352, 358

role in adaptation to light 352	equilibrium constant 40	Recurrent inhibition 316, 320
Pupillary reflex 352, 358	exergonic 38	Red blood cells → erythrocytes
Purine, receptor types 55 F	exothermic 38	5α -Reductase, testosteron 306
second messenger 55 F	hypersensitivity (allergy) 100	Reentry, excitation, heart 194,
Purkinje cells, cerebellum 326,	rate of 40	200
328	constant 40	Reference values 384 ff.
fibers 192	Reactive hyperemia 212	Reflection coefficient (σ) 24, 208,
action potential 200	Reading 336	377
Pursuit eye movement 360	presbyopia 346	Reflex 78, 316 ff.
Putamen 314, 324, 326	Rebound effect 272	abdominal 320
P wave (ECG) 190, 196	phenomenon 328	absence of 320
PWV (pulse wave velocity) 190	Receptive field 314, 354	accomodation 240
Pylorus 240	retina 354	arc 78
Pyramidal cells, motosensory	relaxation 238, 240	autonomic 320
cortex 332 tract 324	Receptor (sensory receptors →	conditioned 236, 242
	sensors) 6, 55 F, 268,	saliva secretion 236
Pyridoxal 260	312	consensual 359
Pyridoxamine 260	acetylcholin → cholinoceptor	corneal 320
Pyridoxine 260	second messenger 274 f. ADH 55 F, 212	crossed 320
Pyrogens 224	adrenalin → adrenoceptors	diagnostic 320
Pyruvate 73 B2 Pyruvic acid 73 B2	adrenarii → adrenoceptors	endogenous 234 extensor 320
Fyruvic acid 73 B2	angiotensin II 212	flexion 320
	CCK 55 F	gastrocolic 264
Q	gallbladder 248	gastroconc 204 gastrointestinal tract 234
Q	stomach 242	Henry-Gauer 218
Q10 value 40	cobalamine 28	Hoffmann's 316
QRS axis, ECG 198	cholinergic 82	knee jerk 316
QRS complex, ECG 196	dopamin 248	labyrinthine postural 328
QRS vector, mean 196, 198	epinephrine → adrenoceptors	load compensation 316
QT interval, ECG 196, 198	endothelin 212	locomotor 320
Quinidine, tubular transport 160	G protein-dependent 52	monosynaptic 316
Quinine 338	histamin 55 F, 212, 242, 274,	neck, tonic 328
Quotient, respiratory 120, 136,	276	nutrition 320
228	hormone 266, 268, 274	orthostatic 204
Q wave, ECG 196, 198	immunoglobulins 96	peristaltic 234, 244
abnormalities 198	insulin 268, 282	plantar 320
	internalization 52	polysynaptic 320
	ionotropic 52, 55 F, 82	postural 328
R	LDL 28	proprioceptive 316
	mediated endocytosis 12 f.	protective 238, 320
R (gas constant) 20, 32, 34, 112	metabotropic 52, 55 F, 82	pupillary 359
R wave, ECG 196	noradrenaline → adrenocep-	spreading 320
Radiation 222, 224	tors	statokinetic 328
Rahn valve 114	norepinephrin \rightarrow adrenocep-	time 316, 320
Raphe nuclei 328	tors	tone 66
RAS (renin-angiotensin system)	proteins 268, 278, 296	labyrinth 328
170, 184, 218	recycling 12, 13 F, 29 C	vagovagal, esophagus 238
RA sensor (rapidly adapting pres-	synapse 50, 55 F	vestibulo-ocular 342
sure sensor) 314	transferrin 90	withdrawal 320
Rate constant 40	tyrosinkinase 282	Refraction, eye 346
RBC (red blood cell) → erythro-	Receptor guanylyl cyclase 278	Refractive power 346
cyte	Receptor serine/threonine kinase	Refractory period (action poten-
RBF (renal blood flow) → kidney	278	tial) 46
RCC (red cell count) 90	Receptor tyrosine kinases 278,	absolute 46
Reabsorption, capillaries 208 renal tubule 148, 152 ff., 162,	282	relative 46
	Receptor tyrosine phophatases	heart 200
174 ff., 180 ff. Reaction, acrosomal 308	278	Regulation 4, 266
coupled 41	Recoverin 350, 352	Reissner's membrane 366
coupicu +1	Recruitment, motoric unit 58	Relaxation, receptive 238, 240

endergonic 38

endothermic 38

Rectum 232, 264

acoustic neurons 368

Release-inhibiting hormones

268, 269, 280

Releasing hormones 268, 269,	equivalent 74, 106	Rigidity, decerebrate 328
280	exchange rate 120	Right axis (heart) 198
REM sleep (rapid eye movement)	muscles 132	Rigor complex 64
334	quotient (RQ) 120, 136, 228	mortis 64, 326
Remnants of chylomicrons 256	rate 106, 118	Rinne's test 364
of VLDL 254	sensors 132	Riva-Rocci (blood pressure
	stimulants 132	measurement) 206
Renal (→ also kidney) artery,		-
stenosis 184	volumes 112	RNA (ribonucleic acid) 8 ff.
blood flow 150	conversion 112	RNA polymerases 8
failure 142, 176	standardization 112	RNAses, pancreas 246
function, glucocorticoid effect	ways, flow resistance 116	Rods 344, 348, 350, 356
296	Response, local 46	adaptation 352
Renin 148, 184	Resting expiratory level 112	monochromatism 352
hypertension 216	membrane potential 44	light absorption maximum 356
Renin-angiotensin system (RAS)	metabolic rate 226	Roots, spinal cord 310
170, 184, 218	position, respiration 112	Rotation, sensors 342
control 184	pressure-volume curve, heart	Round window, ear 364, 366
Renshaw cells 316, 320, 324	202	RPF (renal plasma flow) → kid-
Replication, cell 8	lung and thorax 116	nev
Repolarisation (action potential)	skeletal muscle 66	RQ → respiratory quotient
46	tidal volume 112, 114, 118	rRNA (ribosomal ribonucleic
Repressor protein 8	tremor 226	acid) 8, 10
	Resuscitation 110	rTr ₃ (reverse-T ₃) 288
Reproduction 298 ff., 330		
RES (Reticuloendothelial system)	Reticular zone, adrenal cortex	Ruffini's corpuscle 314
(→ also macrophage	296	RV (residual volume) 112, 114
and mononuclear	Reticuloendothelial system (RES)	R wave (ECG) 196
phagocytotic system)	(→ also macrophage)	Ryanodine receptors (RYR) 62, 64
96	96	RYR 65 D
Reserve volume, expiratory 112	Reticulocytes 88	RYR1 63 B
inspiratory 112	Reticulum, endoplasmic 10, 12,	RYR2 63 B
Residual capacity, functional	sarcoplasmic 60, 194	
(FRC) 112, 114, 116	Retina (→ also visual and eye)	
	Retina (→ also visual and eye) 344, 346, 356, 358	S
(FRC) 112, 114, 116 volume (RV) 112, 114		S
(FRC) 112, 114, 116 volume (RV) 112, 114 Resistance peripheral 186, 206	344, 346, 356, 358 contrast of a stimulus 354	_
(FRC) 112, 114, 116 volume (RV) 112, 114	344, 346, 356, 358 contrast of a stimulus 354 corresponding points 360	σ (reflection coefficient) 24, 377
(FRC) 112, 114, 116 volume (RV) 112, 114 Resistance peripheral 186, 206 hypertension 216 unit 375	344, 346, 356, 358 contrast of a stimulus 354 corresponding points 360 ganglion cells 354, 358	σ (reflection coefficient) 24, 377 S hormones → glucocorticoids
(FRC) 112, 114, 116 volume (RV) 112, 114 Resistance peripheral 186, 206 hypertension 216 unit 375 vessels 186, 188	344, 346, 356, 358 contrast of a stimulus 354 corresponding points 360 ganglion cells 354, 358 processing of visual stimuli	σ (reflection coefficient) 24, 377 S hormones → glucocorticoids and cortisol
(FRC) 112, 114, 116 volume (RV) 112, 114 Resistance peripheral 186, 206 hypertension 216 unit 375 vessels 186, 188 Respiration (→ respiratory) 106	344, 346, 356, 358 contrast of a stimulus 354 corresponding points 360 ganglion cells 354, 358 processing of visual stimuli 354	o (reflection coefficient) 24, 377 S hormones → glucocorticoids and cortisol S wave, ECG 196
(FRC) 112, 114, 116 volume (RV) 112, 114 Resistance peripheral 186, 206 hypertension 216 unit 375 vessels 186, 188 Respiration (→ respiratory) 106 artificial 110	344, 346, 356, 358 contrast of a stimulus 354 corresponding points 360 ganglion cells 354, 358 processing of visual stimuli 354 receptive field 354	o (reflection coefficient) 24, 377 S hormones → glucocorticoids and cortisol S wave, ECG 196 S (siemens), unit 375
(FRC) 112, 114, 116 volume (RV) 112, 114 Resistance peripheral 186, 206 hypertension 216 unit 375 vessels 186, 188 Respiration (→ respiratory) 106 artificial 110 control 132	344, 346, 356, 358 contrast of a stimulus 354 corresponding points 360 ganglion cells 354, 358 processing of visual stimuli 354 receptive field 354 sensors 344, 348	σ (reflection coefficient) 24, 377 S hormones → glucocorticoids and cortisol S wave, ECG 196 S (siemens), unit 375 SA → sinoatrial
(FRC) 112, 114, 116 volume (RV) 112, 114 Resistance peripheral 186, 206 hypertension 216 unit 375 vessels 186, 188 Respiration (→ respiratory) 106 artificial 110 control 132 dead space 110, 114	344, 346, 356, 358 contrast of a stimulus 354 corresponding points 360 ganglion cells 354, 358 processing of visual stimuli 354 receptive field 354 sensors 344, 348 potential 354	o (reflection coefficient) 24, 377 S hormones → glucocorticoids and cortisol S wave, ECG 196 S (siemens), unit 375 SA → sinoatrial SA1 (slow adapting pressosensor)
(FRC) 112, 114, 116 volume (RV) 112, 114 Resistance peripheral 186, 206 hypertension 216 unit 375 vessels 186, 188 Respiration (→ respiratory) 106 artificial 110 control 132 dead space 110, 114 effects of diving 134	344, 346, 356, 358 contrast of a stimulus 354 corresponding points 360 ganglion cells 354, 358 processing of visual stimuli 354 receptive field 354 sensors 344, 348 potential 354 11-cis-Retinal 348	o (reflection coefficient) 24, 377 S hormones → glucocorticoids and cortisol S wave, ECG 196 S (siemens), unit 375 SA → sinoatrial SA1 (slow adapting pressosensor) 314
(FRC) 112, 114, 116 volume (RV) 112, 114 Resistance peripheral 186, 206 hypertension 216 unit 375 vessels 186, 188 Respiration (→ respiratory) 106 artificial 110 control 132 dead space 110, 114 effects of diving 134 functions 106	344, 346, 356, 358 contrast of a stimulus 354 corresponding points 360 ganglion cells 354, 358 processing of visual stimuli 354 receptive field 354 sensors 344, 348 potential 354 11-cis-Retinal 348 Retinol 260, 350	o (reflection coefficient) 24, 377 S hormones → glucocorticoids and cortisol S wave, ECG 196 S (siemens), unit 375 SA → sinoatrial SA1 (slow adapting pressosensor) 314 SA2 314
(FRC) 112, 114, 116 volume (RV) 112, 114 Resistance peripheral 186, 206 hypertension 216 unit 375 vessels 186, 188 Respiration (→ respiratory) 106 artificial 110 control 132 dead space 110, 114 effects of diving 134 functions 106 high altitude 132, 136	344, 346, 356, 358 contrast of a stimulus 354 corresponding points 360 ganglion cells 354, 358 processing of visual stimuli 354 receptive field 354 sensors 344, 348 potential 354 11-cis-Retinal 348 Retinol 260, 350 Return, venous 204	o (reflection coefficient) 24, 377 S hormones → glucocorticoids and cortisol S wave, ECG 196 S (siemens), unit 375 SA → sinoatrial SA1 (slow adapting pressosensor) 314 SA2 314 Saccadic eye movement 360
(FRC) 112, 114, 116 volume (RV) 112, 114 Resistance peripheral 186, 206 hypertension 216 unit 375 vessels 186, 188 Respiration (→ respiratory) 106 artificial 110 control 132 dead space 110, 114 effects of diving 134 functions 106 high altitude 132, 136 influence on venous return 110	344, 346, 356, 358 contrast of a stimulus 354 corresponding points 360 ganglion cells 354, 358 processing of visual stimuli 354 receptive field 354 sensors 344, 348 potential 354 11-cis-Retinal 348 Retinol 260, 350 Return, venous 204 Reversal potential 47 B, 56	o (reflection coefficient) 24, 377 S hormones → glucocorticoids and cortisol S wave, ECG 196 S (siemens), unit 375 SA → sinoatrial SA1 (slow adapting pressosensor) 314 SA2 314 Sacadic eye movement 360 Saccharin, taste 338
(FRC) 112, 114, 116 volume (RV) 112, 114 Resistance peripheral 186, 206 hypertension 216 unit 375 vessels 186, 188 Respiration (→ respiratory) 106 artificial 110 control 132 dead space 110, 114 effects of diving 134 functions 106 high altitude 132, 136 influence on venous return 110 internal 106	344, 346, 356, 358 contrast of a stimulus 354 corresponding points 360 ganglion cells 354, 358 processing of visual stimuli 354 receptive field 354 sensors 344, 348 potential 354 11-cis-Retinal 348 Retinol 260, 350 Return, venous 204 Reversal potential 47 B, 56 Reverse-T ₃ (rTr ₃) 288	o (reflection coefficient) 24, 377 S hormones → glucocorticoids and cortisol S wave, ECG 196 S (siemens), unit 375 SA → sinoatrial SA1 (slow adapting pressosensor) 314 SA2 314 Saccadic eye movement 360 Saccharin, taste 338 Saccharase 258
(FRC) 112, 114, 116 volume (RV) 112, 114 Resistance peripheral 186, 206 hypertension 216 unit 375 vessels 186, 188 Respiration (→ respiratory) 106 artificial 110 control 132 dead space 110, 114 effects of diving 134 functions 106 high altitude 132, 136 influence on venous return 110 internal 106 mechanical 110	344, 346, 356, 358 contrast of a stimulus 354 corresponding points 360 ganglion cells 354, 358 processing of visual stimuli 354 receptive field 354 sensors 344, 348 potential 354 11-cis-Retinal 348 Retinol 260, 350 Return, venous 204 Reversal potential 47 B, 56 Reverse-T ₃ (rTr ₃) 288 Rhesus system 100	o (reflection coefficient) 24, 377 S hormones → glucocorticoids and cortisol S wave, ECG 196 S (siemens), unit 375 SA → sinoatrial SA1 (slow adapting pressosensor) 314 SA2 314 Saccadic eye movement 360 Saccharin, taste 338 Saccharase 258 Saccharose 258
(FRC) 112, 114, 116 volume (RV) 112, 114 Resistance peripheral 186, 206 hypertension 216 unit 375 vessels 186, 188 Respiration (→ respiratory) 106 artificial 110 control 132 dead space 110, 114 effects of diving 134 functions 106 high altitude 132, 136 influence on venous return 110 internal 106 mechanical 110 mechanics 108, 110, 112, 144,	344, 346, 356, 358 contrast of a stimulus 354 corresponding points 360 ganglion cells 354, 358 processing of visual stimuli 354 receptive field 354 sensors 344, 348 potential 354 11-cis-Retinal 348 Retinol 260, 350 Return, venous 204 Reversal potential 47 B, 56 Reverse-T ₃ (rTr ₃) 288 Rhesus system 100 Rhinencephalon 340	σ (reflection coefficient) 24, 377 S hormones → glucocorticoids and cortisol S wave, ECG 196 S (siemens), unit 375 SA → sinoatrial SA1 (slow adapting pressosensor) 314 SA2 314 Saccadic eye movement 360 Saccharin, taste 338 Saccharase 258 Saccharose 258 Saccule 342
(FRC) 112, 114, 116 volume (RV) 112, 114 Resistance peripheral 186, 206 hypertension 216 unit 375 vessels 186, 188 Respiration (→ respiratory) 106 artificial 110 control 132 dead space 110, 114 effects of diving 134 functions 106 high altitude 132, 136 influence on venous return 110 internal 106 mechanical 110 mechanics 108, 110, 112, 144, 116	344, 346, 356, 358 contrast of a stimulus 354 corresponding points 360 ganglion cells 354, 358 processing of visual stimuli 354 receptive field 354 sensors 344, 348 potential 354 11-cis-Retinal 348 Retinol 260, 350 Return, venous 204 Reversal potential 47 B, 56 Reverse-T ₃ (rTr ₃) 288 Rhesus system 100 Rhinencephalon 340 Rhodopsin 348, 350, 352, 356	o (reflection coefficient) 24, 377 S hormones → glucocorticoids and cortisol S wave, ECG 196 S (siemens), unit 375 SA → sinoatrial SA1 (slow adapting pressosensor) 314 SA2 314 Saccadic eye movement 360 Saccharin, taste 338 Saccharose 258 Saccharose 258 Saccule 342 Salbutamol 87 B
(FRC) 112, 114, 116 volume (RV) 112, 114 Resistance peripheral 186, 206 hypertension 216 unit 375 vessels 186, 188 Respiration (→ respiratory) 106 artificial 110 control 132 dead space 110, 114 effects of diving 134 functions 106 high altitude 132, 136 influence on venous return 110 internal 106 mechanical 110 mechanics 108, 110, 112, 144,	344, 346, 356, 358 contrast of a stimulus 354 corresponding points 360 ganglion cells 354, 358 processing of visual stimuli 354 receptive field 354 sensors 344, 348 potential 354 11-cis-Retinal 348 Retinol 260, 350 Return, venous 204 Reversal potential 47 B, 56 Reverse-T ₃ (rTr ₃) 288 Rhesus system 100 Rhinencephalon 340	σ (reflection coefficient) 24, 377 S hormones → glucocorticoids and cortisol S wave, ECG 196 S (siemens), unit 375 SA → sinoatrial SA1 (slow adapting pressosensor) 314 SA2 314 Saccadic eye movement 360 Saccharin, taste 338 Saccharase 258 Saccharose 258 Saccule 342
(FRC) 112, 114, 116 volume (RV) 112, 114 Resistance peripheral 186, 206 hypertension 216 unit 375 vessels 186, 188 Respiration (→ respiratory) 106 artificial 110 control 132 dead space 110, 114 effects of diving 134 functions 106 high altitude 132, 136 influence on venous return 110 internal 106 mechanical 110 mechanics 108, 110, 112, 144, 116	344, 346, 356, 358 contrast of a stimulus 354 corresponding points 360 ganglion cells 354, 358 processing of visual stimuli 354 receptive field 354 sensors 344, 348 potential 354 11-cis-Retinal 348 Retinol 260, 350 Return, venous 204 Reversal potential 47 B, 56 Reverse-T ₃ (rTr ₃) 288 Rhesus system 100 Rhinencephalon 340 Rhodopsin 348, 350, 352, 356	o (reflection coefficient) 24, 377 S hormones → glucocorticoids and cortisol S wave, ECG 196 S (siemens), unit 375 SA → sinoatrial SA1 (slow adapting pressosensor) 314 SA2 314 Saccadic eye movement 360 Saccharin, taste 338 Saccharose 258 Saccharose 258 Saccule 342 Salbutamol 87 B
(FRC) 112, 114, 116 volume (RV) 112, 114 Resistance peripheral 186, 206 hypertension 216 unit 375 vessels 186, 188 Respiration (→ respiratory) 106 artificial 110 control 132 dead space 110, 114 effects of diving 134 functions 106 high altitude 132, 136 influence on venous return 110 internal 106 mechanical 110 mechanics 108, 110, 112, 144, 116 mouth-to-mouth 110	344, 346, 356, 358 contrast of a stimulus 354 corresponding points 360 ganglion cells 354, 358 processing of visual stimuli 354 receptive field 354 sensors 344, 348 potential 354 11-cis-Retinal 348 Retinol 260, 350 Return, venous 204 Reversal potential 47 B, 56 Reverse-T ₃ (rTr ₃) 288 Rhesus system 100 Rhinencephalon 340 Rhodopsin 348, 350, 352, 356 kinase 350	o (reflection coefficient) 24, 377 S hormones → glucocorticoids and cortisol S wave, ECG 196 S (siemens), unit 375 SA → sinoatrial SA1 (slow adapting pressosensor) 314 SA2 314 Saccadic eye movement 360 Saccharin, taste 338 Saccharose 258 Saccharose 258 Saccule 342 Salbutamol 87 B Saliva 232, 236, 258, 260
(FRC) 112, 114, 116 volume (RV) 112, 114 Resistance peripheral 186, 206 hypertension 216 unit 375 vessels 186, 188 Respiration (→ respiratory) 106 artificial 110 control 132 dead space 110, 114 effects of diving 134 functions 106 high altitude 132, 136 influence on venous return 110 internal 106 mechanical 110 mechanics 108, 110, 112, 144, 116 mouth-to-mouth 110 muscles 108	344, 346, 356, 358 contrast of a stimulus 354 corresponding points 360 ganglion cells 354, 358 processing of visual stimuli 354 receptive field 354 sensors 344, 348 potential 354 11-cis-Retinal 348 Retinol 260, 350 Return, venous 204 Reversal potential 47 B, 56 Reverse-T ₃ (rTr ₃) 288 Rhesus system 100 Rhinencephalon 340 Rhodopsin 348, 350, 352, 356 kinase 350 Rhythm, circardian 296, 334	o (reflection coefficient) 24, 377 S hormones → glucocorticoids and cortisol S wave, ECG 196 S (siemens), unit 375 SA → sinoatrial SA1 (slow adapting pressosensor) 314 SA2 314 Saccadic eye movement 360 Saccharin, taste 338 Saccharase 258 Saccule 342 Salbutamol 87 B Saliva 232, 236, 258, 260 primary 236
(FRC) 112, 114, 116 volume (RV) 112, 114 Resistance peripheral 186, 206 hypertension 216 unit 375 vessels 186, 188 Respiration (→ respiratory) 106 artificial 110 control 132 dead space 110, 114 effects of diving 134 functions 106 high altitude 132, 136 influence on venous return 110 internal 106 mechanical 110 mechanics 108, 110, 112, 144, 116 mouth-to-mouth 110 muscles 108 pressure-volume relationships	344, 346, 356, 358 contrast of a stimulus 354 corresponding points 360 ganglion cells 354, 358 processing of visual stimuli 354 receptive field 354 sensors 344, 348 potential 354 11-cis-Retinal 348 Retinol 260, 350 Return, venous 204 Reversal potential 47 B, 56 Reverse-T ₃ (rTr ₃) 288 Rhesus system 100 Rhinencephalon 340 Rhodopsin 348, 350, 352, 356 kinase 350 Rhythm, circardian 296, 334 diurnal 334 generator 334	o (reflection coefficient) 24, 377 S hormones → glucocorticoids and cortisol S wave, ECG 196 S (siemens), unit 375 SA → sinoatrial SA1 (slow adapting pressosensor) 314 SA2 314 Saccadic eye movement 360 Saccharin, taste 338 Saccharase 258 Saccharose 258 Saccharose 258 Saccule 342 Salbutamol 87 B Saliva 232, 236, 258, 260 primary 236 secondary 236
(FRC) 112, 114, 116 volume (RV) 112, 114 Resistance peripheral 186, 206 hypertension 216 unit 375 vessels 186, 188 Respiration (→ respiratory) 106 artificial 110 control 132 dead space 110, 114 effects of diving 134 functions 106 high altitude 132, 136 influence on venous return 110 internal 106 mechanical 110 mechanics 108, 110, 112, 144, 116 mouth-to-mouth 110 muscles 108 pressure-volume relationships 116 work of breathing 116	344, 346, 356, 358 contrast of a stimulus 354 corresponding points 360 ganglion cells 354, 358 processing of visual stimuli 354 receptive field 354 sensors 344, 348 potential 354 11-cis-Retinal 348 Retinol 260, 350 Return, venous 204 Reversal potential 47 B, 56 Reverse-T ₃ (rTr ₃) 288 Rhesus system 100 Rhinencephalon 340 Rhodopsin 348, 350, 352, 356 kinase 350 Rhythm, circardian 296, 334 diurnal 334 generator 334 Rhythmicity, heart 192	o (reflection coefficient) 24, 377 S hormones → glucocorticoids and cortisol S wave, ECG 196 S (siemens), unit 375 SA → sinoatrial SA1 (slow adapting pressosensor) 314 SA2 314 Saccadic eye movement 360 Saccharin, taste 338 Saccharose 258 Saccharose 258 Saccule 342 Salbutamol 87 B Saliva 232, 236, 258, 260 primary 236 secondary 236 Salivary glands 236 blood flow 214
(FRC) 112, 114, 116 volume (RV) 112, 114 Resistance peripheral 186, 206 hypertension 216 unit 375 vessels 186, 188 Respiration (→ respiratory) 106 artificial 110 control 132 dead space 110, 114 effects of diving 134 functions 106 high altitude 132, 136 influence on venous return 110 internal 106 mechanical 110 mechanics 108, 110, 112, 144, 116 mouth-to-mouth 110 muscles 108 pressure-volume relationships 116 work of breathing 116 Respiratory acidosis 144	344, 346, 356, 358 contrast of a stimulus 354 corresponding points 360 ganglion cells 354, 358 processing of visual stimuli 354 receptive field 354 sensors 344, 348 potential 354 11-cis-Retinal 348 Retinol 260, 350 Return, venous 204 Reversal potential 47 B, 56 Reverse-T ₃ (rTr ₃) 288 Rhesus system 100 Rhinencephalon 340 Rhodopsin 348, 350, 352, 356 kinase 350 Rhythm, circardian 296, 334 diurnal 334 generator 334	o (reflection coefficient) 24, 377 S hormones → glucocorticoids and cortisol S wave, ECG 196 S (siemens), unit 375 SA → sinoatrial SA1 (slow adapting pressosensor) 314 SA2 314 Saccadic eye movement 360 Saccharin, taste 338 Saccharase 258 Saccule 342 Salbutamol 87 B Saliva 232, 236, 258, 260 primary 236 secondary 236 Salivary glands 236 blood flow 214 Salt (→ also Na* and Cl⁻), balance
(FRC) 112, 114, 116 volume (RV) 112, 114 Resistance peripheral 186, 206 hypertension 216 unit 375 vessels 186, 188 Respiration (→ respiratory) 106 artificial 110 control 132 dead space 110, 114 effects of diving 134 functions 106 high altitude 132, 136 influence on venous return 110 internal 106 mechanical 110 mechanics 108, 110, 112, 144, 116 mouth-to-mouth 110 muscles 108 pressure-volume relationships 116 work of breathing 116 Respiratory acidosis 144 alkalosis 142, 144	344, 346, 356, 358 contrast of a stimulus 354 corresponding points 360 ganglion cells 354, 358 processing of visual stimuli 354 receptive field 354 sensors 344, 348 potential 354 11-cis-Retinal 348 Retinol 260, 350 Return, venous 204 Reversal potential 47 B, 56 Reverse-T3 (ГТ3) 288 Rhesus system 100 Rhinencephalon 340 Rhodopsin 348, 350, 352, 356 kinase 350 Rhythm, circardian 296, 334 diurnal 334 generator 334 Rhythmicity, heart 192 Riboflavin, intestinal absorption 260	o (reflection coefficient) 24, 377 S hormones → glucocorticoids and cortisol S wave, ECG 196 S (siemens), unit 375 SA → sinoatrial SA1 (slow adapting pressosensor) 314 SA2 314 SA2 314 SAccadic eye movement 360 Saccharin, taste 338 Saccharase 258 Saccharase 258 Saccharose 258 Saccharose 258 Saccule 342 Salbutamol 87 B Saliva 232, 236, 258, 260 primary 236 secondary 236 Salivary glands 236 blood flow 214 Salt (→ also Na* and Cl⁻), balance 148, 162, 170
(FRC) 112, 114, 116 volume (RV) 112, 114 Resistance peripheral 186, 206 hypertension 216 unit 375 vessels 186, 188 Respiration (→ respiratory) 106 artificial 110 control 132 dead space 110, 114 effects of diving 134 functions 106 high altitude 132, 136 influence on venous return 110 internal 106 mechanical 110 mechanics 108, 110, 112, 144, 116 mouth-to-mouth 110 muscles 108 pressure-volume relationships 116 work of breathing 116 Respiratory acidosis 144 alkalosis 142, 144 air, purification 110	344, 346, 356, 358 contrast of a stimulus 354 corresponding points 360 ganglion cells 354, 358 processing of visual stimuli 354 receptive field 354 sensors 344, 348 potential 354 11-cis-Retinal 348 Retinol 260, 350 Return, venous 204 Reversal potential 47 B, 56 Reverse-T₃ (rTr₃) 288 Rhesus system 100 Rhinencephalon 340 Rhodopsin 348, 350, 352, 356 kinase 350 Rhythm, circardian 296, 334 diurnal 334 generator 334 Rhythmicity, heart 192 Riboflavin, intestinal absorption 260 Ribonucleic acid → RNA	o (reflection coefficient) 24, 377 S hormones → glucocorticoids and cortisol S wave, ECG 196 S (siemens), unit 375 SA → sinoatrial SA1 (slow adapting pressosensor) 314 SA2 314 Saccadic eye movement 360 Saccharin, taste 338 Saccharose 258 Saccharose 278 Salvarose 278 Sal
(FRC) 112, 114, 116 volume (RV) 112, 114 Resistance peripheral 186, 206 hypertension 216 unit 375 vessels 186, 188 Respiration (→ respiratory) 106 artificial 110 control 132 dead space 110, 114 effects of diving 134 functions 106 high altitude 132, 136 influence on venous return 110 internal 106 mechanical 110 mechanics 108, 110, 112, 144, 116 mouth-to-mouth 110 muscles 108 pressure-volume relationships 116 work of breathing 116 Respiratory acidosis 144 alkalosis 142, 144 air, purification 110 chain 12, 73 B3	344, 346, 356, 358 contrast of a stimulus 354 corresponding points 360 ganglion cells 354, 358 processing of visual stimuli 354 receptive field 354 sensors 344, 348 potential 354 11-cis-Retinal 348 Retinol 260, 350 Return, venous 204 Reversal potential 47 B, 56 Reverse-T₃ (rTr₃) 288 Rhesus system 100 Rhinencephalon 340 Rhodopsin 348, 350, 352, 356 kinase 350 Rhythm, circardian 296, 334 diurnal 334 generator 334 Rhythmicity, heart 192 Riboflavin, intestinal absorption 260 Ribonucleic acid → RNA Ribose 8	o (reflection coefficient) 24, 377 S hormones → glucocorticoids and cortisol S wave, ECG 196 S (siemens), unit 375 SA → sinoatrial SA1 (slow adapting pressosensor) 314 SA2 314 Saccadic eye movement 360 Saccharin, taste 338 Saccharase 258 Saccharose 258 Saccharose 258 Saccule 342 Salbutamol 87 B Saliva 232, 236, 258, 260 primary 236 secondary 236 secondary 236 Salivary glands 236 blood flow 214 Salt (→ also Na⁺ and Cl⁻), balance 148, 162, 170 disturbances 170, 172, 173 B regulation 170
(FRC) 112, 114, 116 volume (RV) 112, 114 Resistance peripheral 186, 206 hypertension 216 unit 375 vessels 186, 188 Respiration (→ respiratory) 106 artificial 110 control 132 dead space 110, 114 effects of diving 134 functions 106 high altitude 132, 136 influence on venous return 110 internal 106 mechanical 110 mechanics 108, 110, 112, 144, 116 mouth-to-mouth 110 muscles 108 pressure-volume relationships 116 work of breathing 116 Respiratory acidosis 144 alkalosis 142, 144 air, purification 110 chain 12, 73 B3 compensation 142	344, 346, 356, 358 contrast of a stimulus 354 corresponding points 360 ganglion cells 354, 358 processing of visual stimuli 354 receptive field 354 sensors 344, 348 potential 354 11-cis-Retinal 348 Retinol 260, 350 Return, venous 204 Reversal potential 47 B, 56 Reverse-T₃ (rTr₃) 288 Rhesus system 100 Rhinencephalon 340 Rhodopsin 348, 350, 352, 356 kinase 350 Rhythm, circardian 296, 334 diurnal 334 generator 334 Rhythmicity, heart 192 Riboflavin, intestinal absorption 260 Ribonucleic acid → RNA Ribose 8 Ribosomal ribonucleic acid →	o (reflection coefficient) 24, 377 S hormones → glucocorticoids and cortisol S wave, ECG 196 S (siemens), unit 375 SA → sinoatrial SA1 (slow adapting pressosensor) 314 SA2 314 Saccadic eye movement 360 Saccharin, taste 338 Saccharase 258 Saccule 342 Salbutamol 87 B Saliva 232, 236, 258, 260 primary 236 secondary 236 Salivary glands 236 blood flow 214 Salt (→ also Na* and Cl⁻), balance 148, 162, 170 disturbances 170, 172, 173 B regulation 170 excretion 162, 170
(FRC) 112, 114, 116 volume (RV) 112, 114 Resistance peripheral 186, 206 hypertension 216 unit 375 vessels 186, 188 Respiration (→ respiratory) 106 artificial 110 control 132 dead space 110, 114 effects of diving 134 functions 106 high altitude 132, 136 influence on venous return 110 internal 106 mechanical 110 mechanics 108, 110, 112, 144, 116 mouth-to-mouth 110 muscles 108 pressure-volume relationships 116 work of breathing 116 Respiratory acidosis 144 alkalosis 142, 144 air, purification 110 chain 12, 73 B3 compensation 142 control 132, 214	344, 346, 356, 358 contrast of a stimulus 354 corresponding points 360 ganglion cells 354, 358 processing of visual stimuli 354 receptive field 354 sensors 344, 348 potential 354 11-cis-Retinal 348 Retinol 260, 350 Return, venous 204 Reversal potential 47 B, 56 Reverse-T₃ (rTr₃) 288 Rhesus system 100 Rhinencephalon 340 Rhodopsin 348, 350, 352, 356 kinase 350 Rhythm, circardian 296, 334 diurnal 334 generator 334 Rhythmicity, heart 192 Riboflavin, intestinal absorption 260 Ribonucleic acid → RNA Ribose 8 Ribosomal ribonucleic acid → rRNA	o (reflection coefficient) 24, 377 S hormones → glucocorticoids and cortisol S wave, ECG 196 S (siemens), unit 375 SA → sinoatrial SA1 (slow adapting pressosensor) 314 SA2 314 SA2 314 SA2 314 SA2 314 SA2 324 SA2
(FRC) 112, 114, 116 volume (RV) 112, 114 Resistance peripheral 186, 206 hypertension 216 unit 375 vessels 186, 188 Respiration (→ respiratory) 106 artificial 110 control 132 dead space 110, 114 effects of diving 134 functions 106 high altitude 132, 136 influence on venous return 110 internal 106 mechanical 110 mechanics 108, 110, 112, 144, 116 mouth-to-mouth 110 muscles 108 pressure-volume relationships 116 work of breathing 116 Respiratory acidosis 144 alkalosis 142, 144 air, purification 110 chain 12, 73 B3 compensation 142	344, 346, 356, 358 contrast of a stimulus 354 corresponding points 360 ganglion cells 354, 358 processing of visual stimuli 354 receptive field 354 sensors 344, 348 potential 354 11-cis-Retinal 348 Retinol 260, 350 Return, venous 204 Reversal potential 47 B, 56 Reverse-T₃ (rTr₃) 288 Rhesus system 100 Rhinencephalon 340 Rhodopsin 348, 350, 352, 356 kinase 350 Rhythm, circardian 296, 334 diurnal 334 generator 334 Rhythmicity, heart 192 Riboflavin, intestinal absorption 260 Ribonucleic acid → RNA Ribose 8 Ribosomal ribonucleic acid →	o (reflection coefficient) 24, 377 S hormones → glucocorticoids and cortisol S wave, ECG 196 S (siemens), unit 375 SA → sinoatrial SA1 (slow adapting pressosensor) 314 SA2 314 Saccadic eye movement 360 Saccharin, taste 338 Saccharase 258 Saccule 342 Salbutamol 87 B Saliva 232, 236, 258, 260 primary 236 secondary 236 Salivary glands 236 blood flow 214 Salt (→ also Na* and Cl⁻), balance 148, 162, 170 disturbances 170, 172, 173 B regulation 170 excretion 162, 170

Saluretics 172	somatovisceral 314	Sex characteristics, female 302
Sarcolemma 56, 60	strength 316	male 306
Sarcomere 60 f.	taste 338	chromosomes 306
resting length 68	touch 314	genetic 306
Sarcoplasm 60	vision 344ff.	hormones, binding globulin
Sarcoplasmic reticulum 60	Sensibility, somatovisceral 314	268, 302, 306
Sarcosomes 60	Sensitization 94, 318	female 294, 298, 300
Satiety peptides 230	reflex pathways 336	male 294, 304, 306
Scala media 364	Rh blood groups 100	synthesis 294
tympani 364, 366	Sensor(s) 312, 337	Sexual arousal 308
vestibuli 364, 366	blood pressure 214	behavior 330
Scalene muscles 108	volume 214	development 306
Scavenger receptors, LDL 256	cold 224	delayed 288
Schlemm's canal 344	D- 314, 316	differentiation 306
Schwann cells 42	filling pressure 214	maturation 298
Sclera 344	heat 314	reflexes 308
Sclerosis 216	light sensitive 344	response 308
SCN (suprachiasmatic nucleus)	olfactory 340	SGLT1 (sodium glucose trans-
334	potential 312	porter, type 1) 158
Scotoma 358	hair cells 366	SGLT2 26, 158
Scotopic vision 348	photosensors 350, 354	SH2 (src[sarcoma]-homology)
Scurvy 226	primary 312, 340	domains 278, 282,
Second messenger 268, 274, 276	P- 314	284
adrenoceptors 84 f.	pressure 314	SHBG (sex hormone-binding
M-cholinoceptors 82	proprioceptive 316	globulin) 268, 302,
neurotransmitter receptors	pulse rate 214	306
52, 55 C	retina 344, 348	Shear force 92
Secondary active cotransport	secondary 312, 338, 342	Shivering, thermoregulation
with Na+ 26, 258	organ of corti 364	222 f.
of Cl ⁻ 262	skin 314	Shock 100, 186, 206, 218
response, immune defense 94	smell 340	anaphylactic 100, 218
saliva 236	touch 314	causes 218
Secretin 234, 240, 246, 274	vibration 314	compensation mechanisms
esophagus 238	warm 314	218
insulin secretion 282	Sensory aphasia 370	hypoglycemic 218, 284
pancreas secretion 246	impression 312	index 218
second messenger 274	input, central processing 322	irreversibel 218
stomach 242	modality 312	lung 122
Secretion, renal → kidney	physiology 312 ff.	manifest (progressive) 218
constitutive 13 F	receptors → sensors	Shunt 120
Secretory granules 12, 268	SERCA (sarcoplasmic endo-	alveolar 120
phase, menstrual 298	plasmic reticulum	arteriovenous 122
Segmentation (intestine) 244	Ca ²⁺ -transporting	blood flow 122, 220
Seizures 136	ATPase) 16, 64	extra-alveolar 120
Selectins 14, 98	Serotonin 102, 160, 230, 274, 276,	left-to-right 220
Selection, clonal 98	330, 332	right-to-left 220
Selenium (Se) 226	cerebral cortex 332	SI units 372 f.
Self-awareness 336	neurons 330	Sickle cell anemia 92
Semen emission 308	platelets 102, 103 A	Siemens (S), unit 375
production 306	receptor types 55 F, 274, 276	Sieving coefficient 24
Semilunar valves, heart 190	second messenger 55 F, 274,	glomerular (GSC) 154
Seminal vesicles, testosterone	276	Siggaard-Andersen nomogram
effect 306, 308	tubular tranport 160	146, 147 C
Seminiferous tubules 306	Sertoli cells 306	Signal recognition particle (SRP)
Senses 312, 314, 322, 338, 340,	Serum 88	12
342	disease 100	Signal sequence 12
balance 342	electrolytes, ECG 198	transduction 274
chemical 338	sickness 100	Sildenafilcitrate (Viagra®) 278
hearing 362 ff.	Servocontrol 6	Silibants, voice 370
movement 316, 342	Servomechanism 6	SIH → somatostatin

position 316, 342

smell 340

Set point, variation 6

value 4

Silicon (Si) 226

Simultaneous contrast 354

Sleep-wake cycle 334

Cii 270	Clidia - filana - than - than - C2	
Singing 370 Single-unit, muscle typ 70	Sliding filament hypothesis 62, 64	release-inhibiting hormone → somatostatin
Sinoatrial (SA) node 190, 192,	Slit membrane, glomerular 148	releasing hormone → soma-
200	SMA (suplementary motor area)	toliberin
Sinus arrhythmia 200	324	Somatovisceral senses 314
bradycardia 200	Small intestine 232, 240	Sone 362
rhythm 192	Ca ²⁺ absorption 290	Sorbitol 262
tachycardia 200	cell replacement 244	Sorting 12, 13 F
Skeletal growth, STH 280	function 244	Sound 362
Skeletal muscle 56 ff., 324 ff.	motility 244	analysis 368
actin-myosin interaction 68	stucture 244	central processing 368
action potential 56, 59 A	Smell 340	conduction 364, 366
all-or-none response 66 blood flow 74	Smooth muscle 70 intestine 244	air 364 deafness 364
regulation 214		dearness 364 direction 368
Ca ²⁺ concentration, cytosolic	M ₃ -receptors 82 SNAP-25 (synaptosome-as-	distance 368
66	sociated protein 25)	frequencies 362, 368
contracture 66	50	intensity 362, 368
difference cardiac/smooth	SNARE (synaptosome-associated	loudness 362
muscle 68	protein receptor) 30	perception 362
fibers	Sneezing 132, 320	physics of 362
extrafusal 316	Snorkel diving 134	pressure 362, 366, 368
intrafusal 316	$S_{O_2}(O_2 \text{ saturation})$ 126	level 362
stimulation 62	Sodium → Na ⁺	sensors 364, 366
types 58	Softening effect, ATP 64	speech 370
glycogen concentration 58	Solubility coefficient, CO ₂ 126	stimulus 362
glycogenesis 282	O ₂ 128	Space travel 136
gradation of contraction	Solubility product, calcium	Spatial orientation 342, 368
force 66	phosphate 290	Spatial summation 52, 320, 352,
lactate production 76 length-force curve 68	Solutes, activity 24, 376 Solutions 376	354, 358 Specific dynamic action → diet-
mechanical features 66	Solvent drag 24, 156, 162	induced thermogene-
mitochondria 63 A	Soma, nerve cell 42	sis
motor units 58, 66	Somatic sensory functions 314	Spectrum, visible 356
myoglobin function 72, 128	Somatoliberin 230, 234, 268, 269,	Speech 236, 362, 370
persistant local depolariza-	280	fundamental frequencies 370
tion 66	Somatomedins 280, 284	Sperm 30, 302, 306
reflex tone 66	Somatosensory centers 314, 322	capacitation 302, 308
rigor 66	Somatostatin (SIH) 84 ff., 230,	emmision 308
complex 64	234, 240, 268 ff.,	estrogens 302
summation 66	280 ff., 284 f.	fertilization 308
superposition 66	actions 284	maturation 306
tetanus 58, 66	cotransmitter 84 effects 284	motility 30, 58 testosterone 306
Skin 74, 222, 224 α ₁ -adrenoceptors 214	influence on glucagon secre-	uterus 302 f.
blood flow 186, 222	tion 272, 284	Spermatids 306
physical work 74	on insulin release 272,	Spermatocytes 306
regulation 214	282, 284	Spermatogonia 306
formation of cholecalciferol	pancreas 282	Spermatogenesis 306
292	receptors 55 F	Spermatozoa 306
mechanosensors 314	second messenger 55 F, 274	Spherical aberration 346
nocisensors 318	secretion 284	Sphincter, esophagus 238
sensibility 314	stomach 234, 242	precapillar 188
sensory functions 222 ff., 314,	Somatotopic representation 322,	pupil 344
322	324	Spines, cerebral cortex 332
temperature 222, 224 thermosensors 222 ff.	Somatotropin (= STH = growth	Spinal cord 78, 310, 320, 322, 324, 326
Sleep 332, 334	hormone = GH) 269, 278, 280, 284, 288,	anterior roots 310
deprivation 334	303	dorsal roots 310
phases 334	influence on insulin release	grey matter 310
stages 334	284	hemisection 322

receptor 278

ipsilateral paralysis 322

reflexes 320	adequate 312	Substance P 52, 86, 248, 318, 324,
segments 310	contrast 354	326
transection 328	transduction 312	gallbladder 248
white matter 310	transformation 312	Substantia nigra 324, 326
ganglion 310, 318	Stokes-Einstein equation 20	Subsynaptic Potential 50
nerves 310	Stomach (\rightarrow also gastric) 240,	Successive contrast 352
shock 328	242, 252, 258, 264	color 354
Spinocerebellum 326	anatomy 240	Succinate, tubular transport 160
Sphingomyelin 14	chief cells 240	Suckling 236, 280, 303
Spinnbarkeit of cervix mucus 298	digestive activity 258	reflex 320, 330
Spiral ganglion 366, 368	distal 232, 240	Sucrose, digestion 258
Spirometer 112, 116	emptying time 232	Sugar hormones → glucocorti-
Spironolactone 172	rate 240	coids and cortisol
SPL (sound pressure level) 362	function 240	Sulci, brain 310
Spleen 232	gastric acid 242	Sulfate 156, 158, 160, 250, 262,
blood cell formation 88	gastrin production 234	294
Spliceosome 10	glands 242	conjugates 160, 250
Splicing 8	juice 242	carrier 250
variable 268	motility 240	tubular secretion 156, 160
Split-brain patient 337	mucus neck cells 240	steroids 294
Spot, blind 348, 358	parietal cells 240, 242	transcellular secretion 156,
Squalene 294	protein digestion 258	160
Src (sarcoma) protein 278	proximal 232, 240	intestinal absorption 262
SRF → somatoliberin	secretion 242	Sulfuric acid (\rightarrow also sulfate) 142,
SRH → somatoliberin	size 240	174
SRP (signal recognition particle)	ulcer 242	Summation 320
12	Stool 262, 264	skeletal muscle 66
Standard bicarbonate concentra-	color 250	spatial 320, 354, 358
tion 142	Stop codon 10	neuron 52
measurement 146	Storage in memory 336	vision 352
Standard temperature pressure	of energy 282	temporal 320
dry (STPD) 112	STPD (standard temperature	neuron 52
Stapedius muscle 364	pressure dry) 112	vision 352
Stapes 364	Strabismus 360	Superoxid dismutase 96
Staphylokinase 104	amblyopia 360	Superposition, skeletal muscle 66
Starch, digestion 246, 258	Strength, ionic 376	Suppository 264
Starling relationship 208	training 76	Suppression, saccadic 360
Start codon 10	Streptokinase 104	T helper cells 98
Starvation 142	Stress, CRH secretion 296	Supraventricular arrhythmias
Static (postural) work 74	cortisol secretion 296	200
Statoliths 342	hyperprolactinemia 303	Surface sensitivity 314
Steady state 41	Stretch sensors 214	tension 118
Stellate cells 328	rectum 264	Surfactant (surface-active agent)
Stem cells 88, 306	reflex 316, 328	118, 122
Step test, Margaria 76	Stria vascularis366	hyperoxia 136
Stercobilin 250	Striatum 326	Suxamethonium 46, 56
Stercobilinogen 250	Stroke volume (SV), heart 76,	SV (stroke volume), heart 76, 106,
Stereocilia 342	106, 186, 190, 202,	186, 190, 202, 204
Stereognosis 314	204	Swallowing 132, 236, 238, 240,
Steroid diabetes 296	increased afterload 204	320
Steroid hormones → steroids	preload 204	S wave (ECG) 196
Steroids 248, 250, 268, 304	physical work 74	Sweat 222, 224
biosynthesis 294	Strophantin 194	glands 214, 222
degradation 294	Struma 286	innervation 78, 79 ff., 224
metabolism 294	STS → somatostatin	secretion 224
placenta 304	ST segment (ECG) 196	SWS (slow-wave sleep) 334
producing glands 294	elevation 198	Symbiosis hypothesis, mitochon-
receptors 278	Stuart-Prower factor 102	dria 12
secretion 248, 250, 294	Subfornical organ 280	Sympathetic → also adrenergic
sex, male 306	Sublingual gland 236	and autonomic
STH → somatotropin	Submandibular gland 236	Sympathetic nervous system,
Stimulus 42		activation 74

	Sympathetic nervous system	second messenger 276	neural factors in 225 D
	constriction, veins 218	Tachypnea 108	Thermodynamics, laws 38 ff.
	flight or fight behavior 330	Tactile motor function 314	Thermogenin 222, 230, 288
	gallbladder 248	TAL (thick ascending limb) 162	Thermogenesis, non-shivering 224
	gastrointestinal tract 234	Tanning lamps, calcitriol synthe-	Thermoneutral zone 224
	heart 194	sis 292	Thermoregulation 222, 224
	physical work 74	Target value (control circuit) 4	TRH release 286
	saliva secretion 236	Taste 338	Thermosensors 224, 314
	regulation of circulation 214	Taurine 248	hypothalamus 330
	Symport, definition 26	Tawara's bundle branches 192	central 224
	Synapse 42, 50 ff., 266	TBG (thyroxine-binding globulin)	Thiamine, intestinal absorption
	axoaxonic 320	288	260
	electrical 18, 50	TBPA (thyroxine-binding pre-	Thiazide 162, 172
	latency 52	albumin) 288	Thick ascending limb (TAL) 162
	postsynaptic receptors 50, 55 F	TCT → calcitonin	Thiocyanate 286
	functions 42	Tears 344	Thiocyanate 200 Thiouracil 286
	reciprocal 340	Tectorial membrane 364	Thirst 168, 170, 224, 236
	valve-like function 48	Tectum 326	center 168
	Synaptic cleft 42, 50	TEE (total energy expenditure)	shock 218
	delay 52	226	Thorax 108 ff.
	facilitation 50	Telencephalon 310	Three-dimensional vision 360
	inhibition 320	Temperature 375	Threonin 226
	knobs 42	body 132, 222, 224	Threshold(s) 312, 340
	pontentiation 50	circadian variation 224, 381	aerobic 76
	transmission 42	C	anaerobic 72, 76
	termination 52, 56	regulation 222 ff.	audiogram 366
	Synapsin 50	skin 222	auditory 362 ff.
	Synaptobrevin 50	unit 375	potential 46, 48, 192
Į	Synaptosome-associated protein	Temporal lobe system 336	recognition, smell 340
	25 (SNAP-25) 50	Temporal summation, visual re-	taste 338
	Synaptosome-associated protein	ceptors 352	visual 352
	receptor (SNARE) 30	Tendon, Golgi organs 316	Thrombin 102, 104
	Synaptotagmin 50	Tension time index, heart muscle	Thrombocyte (→ also platelet)
	Synchronization to day-night	210	88. 102 f.
	cycle 334	Tensor tympani muscle 364	aggregation 102
	Syncytium 16, 18	Terminal button 42	inhibition 85, 104
	Synovial A-cells 96	Testis 254, 268, 294, 306	Thrombocytopathia 104
	System, closed 40	FSH effect 306	Thrombocytopenia 104 Thrombocytopenia 100, 104
	conicellular 358	HDL receptors 254	Thrombokinase 104
	endocrine 266 ff.	Testosteron, 5-α-dihydro 306	Thrombopoietin 88, 148
	limbic 310, 330	Testosterone 268, 300, 306	Thromboprotection 104
	magnocellular 358	receptor 306	Thrombosis 104
	opened 40	synthesis 294	Thrombosis 104 Thromboxane A ₂ 102, 214, 269,
	parvicellular 358	Tetanus 56	276
	Systole 190	skeletal muscle 66	second messenger 276
	Systole 150	smooth muscle 70	Thrombus 102, 104
		Tetany 46, 290, 292	Thryptophan 226
	Т	Tetrahydrofolate 260	Thymine 8
	•	Tetraiodothyronine (T ₄) 286	Thymosin 266
	t (ton), unit 374	Tetrodotoxin (TTX) 47 B	Thymus 88, 94
	T (multiple of a unit) 373	TGF(tubuloglomerular feedback)	Thyreotropic hormone-releasing
	T wave, ECG 196	172, 184	hormone (TRH) →
	$T_3 \rightarrow \text{triiodothyronine and thy-}$	TGF(transforming growth factor)-	thyroliberin
	roid hormones	β receptor 278	Thyrocalcitonin → calcitonin
	$T_4 \rightarrow$ thyroxine and thyroid hor-	Thalamus 310, 318, 322, 338, 340,	Thyroglobulin 286
	mones	368	Thyroid gland 36, 286
	Tachycardia 200, 218	auditory pathway 368	hormones 286
	atrial 200	disinhibition 326	nuclear receptors 278, 288
	in shock 218	visual pathway 358	actions 288
	sinus 200	Theca cells 300	control of release 288
	ventricular 200	Theophylline 276	parafollicular C cells 36, 286,
			r

ventricular 200 Tachykinin 55 F

Thermal balance 222

peroxidase (TPO) 286	neurogenic 70
Thyroid-stimulating hormone \rightarrow	Torr, conversion into SI unit 374
thyrotropin	Touch sensor 314
Thyroliberin 268, 269, 280, 286,	tPA (tissue plasminogen activa-
303	tor) 104
second messenger 274, 276	TPO (thyroid peroxidase) 286
Thyrotropic hormone (TSH) →	TPR (total peripheral resistance)
thyrotropin	186 f.
Thyrotropin (TSH) 269, 280, 286	Trace elements 226 Trachea 370
receptor, autoantibody 288 releasing hormone → thy-	Tractus, corticospinalis lateralis
roliberin	324
second messenger 274, 276	olfactoius 340
Thyroxine (T ₄) 268, 286	opticus 358
Thyroxine-binding globulin	reticulospinalis lateralis 328
(TBG) 288	medialis 328
prealbumin (TBPA) 288	retinohypothalamic 334
Tidal volume 106, 112, 114, 118,	rubrospinalis 328
120	spinocerebellaris anterior 328
alveolar part (V _A) 114, 120	spinoreticular 318
exercise 74	spinothalamicus ventralis 318,
maximum 77 C	322
physical work 74	vestibulospinalis 328
training 76	Training 76
Tiffeneau test 118	Transcallular transport 18, 154
Tight junction, function 18	Transcellular transport 18, 154 Transcobalamine 92, 260
renal tubule 154 Timbre 362	Transcortin 92, 296
Tin (Sn) 226	Transcription factor, hormon-ac-
Tip links 342, 366	tivated 278, 288
Tissue, hormones 268	Transcytosis 18, 28
injury, blood clotting 103 B1	Transducin 276, 348, 350, 352
plasminogen activator (tPA)	Transduction 312, 348
104	channels, hair cells 366
respiration 106, 130	photoelectric 348
thrombokinase 104	signal 274
thromboplastin 102	stimulus 312
inhibitor 104	Transfer ribonucleic acid
Titin 60, 66	\rightarrow tRNA
Titratable acid 174	Transferrin 90
Titration curve 380	Transformation 312
T-killer cells 94, 98, 232	Transforming growth factor
TLC (total lung capacity) 112, 113	(TGF)-β 278
A TNF(tumor necrosis factor)α, ef-	Transfusion, placental 220 Translation 10
fect on CRH secretion	Translocation, hormone-receptor
296	protein complex 278
Tocopherol 260	Translocator protein 12
Tolerance, immunologic, periph-	Transmembrane proteins 12
eral 98	Transmission, hereditary 8 f.
central 94	Transmitter → neurotransmitter
Ton, unit 374	Transplant rejection, primary 100
Tone (sound) 362	immunosuppression 98
Tongue 338	Transport, active 26
Tonicity 377 f.	axoplasmic 42, 58, 280
Tonotopicity, auditory pathway	convection 24
368	electrogenic 26, 28
Tonus, skeletal muscle 328	electroneutral 28
smooth muscle 70	energy-dependent 26 non-ionic 174
myogenic 70 neurogenic 70	paracellular 18, 154
incurogenic / 0	paracential 10, 134

primary active 26 rate, maximal (I_{max}) 383 f. rheogenic 26, 28 saturable 28 secondary active 26 specific 28 tertiary active 29 B transcellular 18, 154 transmembrane 16 all-trans-retinal 348 Treadmill ergometry 76 Trehalase 258 Trehalose 258 Tremor, intention 328 of resting 326 TRF → thyroliberin TRH → thyroliberin Triacylglycerol 252, 254 absorption 252, 254 chemical structure 227 B digestion 246, 254 hydrolase 252 lipoproteins 254 source 257 D fate 257 D synthesis, from free fatty ecids Triads 60, 62 Trigger effect, Ca2+, heart 62, 194 Trigger zone, chemosensory 238 Triglycerides → Triacylglycerol Triiodothyronine (T₃) 268, 278, 286 Trimetaphan 82 Tripalmitin 228 Tripeptides, intestinal absorption 258 Tricuspid valve 190 Tritanomaly 356 Tritanopia 356 tRNA (transfer ribonucleic acid) 8 Tropomyosin 60 Troponin 60, 62 Troponin-tropomyosin complex 64 Trypsin 246, 252, 258, 260 activation 247 D CCK secretion 246 colipase activation 252 phospholipase A2 activation 252 effects 247 D Trypsinogen 246 TSC (thiazide sensitive co-transporter) 162 TSH → thyrotropin T(ransversal tubules) system, skeletal muscle 60

TTX (Tetrodotoxin) 47 B Tube, uterine 308, 309 A Tuberculum olfactorium 340

Tubocurarine 56, 82

Tonus fibers 66

passive 20

Tubular excretion 154	urine concentration 166	Valence, ion 375
reabsorption 154, 158	Ureter 148	Valine 226
secretion 154	innervation 79 ff.	Valsalva's maneuver 116
Tubule (→ also Kidney), distal	Urethra 308	Vanadium (V) 226
148	Uric acid 158, 176	Vanilloid receptor Typ 1 (VR1)
convoluted (DCT) 148	excretion, renal 174	314
driving forces for resorption 162	reabsorption 156, 158 secretion 156,	Van't Hoff's law 24, 377, 388 Staverman 24
epithelium, structure 154	Urinary bladder 70, 148, 308	Vas afferens, glomerulus 148, 184
proximal 148, 162, 166, 174	innervation 79 ff.	deferens 308
convoluted (PCT) 164	calculi 158, 178	efferens, glomerulus 148, 184
Tubules, longitudinal, skeletal	Urine 148 ff., 184	Vasa recta, renal medulla 150,
muscle 60	Ca ²⁺ 290	164
seminiferous 306	concentration mechanism 164,	Vasoactive intestinal peptide
transverse, skeletal muscle 60	166	(VIP) 52, 86, 232, 234,
Tubulin 30, 58	disorders 166	236, 262, 280
Tubuloglomerular feedback (TGF)	flow rate 152	cerebral cortex 332
172, 184	17-ketosteroids 294	substances 212
Tumor cells, immune defense 96	NH+4 176	Vasoconstriction 122, 212, 213 B,
Tumor necrosis factor (TNF)α	osmolality 166	214, 220
296	pH 156, 174, 176	cortisol 296
Tuning, hearing 368	primary 164	hypoxic 122, 212, 220
T wave, ECG 196, 198	titratable acid 174	fetus 221 C
Two-point discrimination 314, 322	volume 152 Urobilinogen 250	shock 218
TXA ₂ (thromboxan A ₂) 102, 269	Urokinase 104	thermoregulation 224 veins 218
Tympanic cavity 364	UT1 (urea transporter Typ 1) 166	Vasodilatation 212, 213 B, 214,
Tyrosine 84	UT2 166	278
derivates, hormonal 268	Uterine tube 308, 309 A	NO 278
Tyrosine kinase-associated re-	Uterus 70, 303, 304	thermoregulation 224
ceptors 278	α_1 -adrenoceptor 304	Vaso-intestinal peptide (VIP) 52,
•	contractions 280	86, 232, 234, 236, 238
	erection 308	262, 274, 280
U	estrogens 302	cerebral cortex 332
	gap junctions 304	neurons 240
UCP (uncoupling protein) 230	menstrual cycle 298	rectum 264
UDP(uridindinucleotid)-glu-	muscle 303	second messenger 274
curonic acid 250	oxytocin 303	Vasopressin (→ also adiuretin)
Ulcer, gastric 242	progesteron 303	166, 274, 276
Ultrafiltrate 154 Ultrafiltration coefficient 152	prostaglandins 304, 308 UTP (uridine triphosphate) 250	Vegetarian diet 142 Vectorcardiogram 196
Umami 338	Utricle 342	Vein(s) 188, 204
Umbilical artery 220	UV light, exposure 292	pulmonary 186
vein 220	formation of vitamin D 292	umbilical 220
Unconsciousness, anoxia 130	deficiency 292	valves 204
Uncoupling protein (UCP) 230	Uvula, cerebellum 326	Veith-Müller horopter 360
Uniporter 28		Velocity, blood flow 188
Units, basic 372		of conduction, in nerve fibres
derived from SI units 372	V	48
information 312		detectors 314, 316
multiples 373	$V_{O_2} \rightarrow O_2$ utilization	linear, unit
SI 372	V _A (alveolar part of tidal volume)	volume 374
submultiples 373	114	Venae cavae 188
Unmyelinated nerve fibers 48 Uracil 8	V _D (dead space volume) 114	Venoles 189 A
	V _T (tidal Volume) 114	postcapillary 188
Urate → uric acid Urea 158, 166, 176	Vaccination 94 Vagina 302, 308	Venous central pressure 186, 190 204, 218
carrier 166	estrogens 302	circulation 204
excretion 166	pH 302	flow 204
production 176	Vagus nerve (→ also cholinergic)	pressure 204
reabsorption, renal 156, 158	242, 248, 370	curve 191 A

reabsorption, renal 156, 158 recirculation 166

val, unit 375

indifference point 204

influence on capillary fluid	Vestibulocerebellum 326	E (D- α -tocopherol) 226, 252,
exchange 208	Viagra® (sildenafilcitrate) 278	260
shock 218	Vibration, sensors 314	H (biotin) 226, 260
return 110, 204	Villi, intestinal 244	K (Phytonadion) 102, 104, 226,
artificial respiration 110	Vimentin 14	252, 264
Bainbridge reflex 216	VIP → vasointestinal peptide	antagonists 104
driving forces 204	Virilization 306	deficiency 104
respiration 204 artificial 110	Virus, immune defense 94, 96, 98 Visceral afferents 78, 234	K ₁ 226, 260 K ₂ 226, 260
Ventilation (→ also lung) 74, 106,	Viscosity, blood 92, 188	Vitamins 226
108, 120, 130	Viscous-isotropic phase, lipid	absorption 260
alveolar 106, 120	digestion 252	deficiencies 226
high altitude 136	Visible spectrum 356	fat-soluble 226, 252, 260, 302
peak expiratory pressure 116	Vision (→ also eye and retina)	toxiciticy 226
inspiratory pressure 116	346	water-soluble 260
pressure differences 108	binocular 358, 360	Vitrous body 344
dead space 106, 114	color 356	opacification 136
diving 134	day 348	VLA-4 (adhesion molecule) 98
driving force for 108	threshold 352	VLDL (very low density lipo-
exercise 74	depth 360	proteins) 254, 256,
high altitude 136	dim-light 348	260
mechanical 110	movement 358	estrogen effect 302
muscle 108, 132	night 348	remnants 254
/perfusion ratio 122	photopic 348	Vocal cord 370
regulation 126	scotopic 348	paralysis 118
chemosensors 126	three dimensional 360	Voice 370
rhythm generator 132 total 106	Visual (→ also eye and retina)	change 306
acidosis 142	acuity 348, 354, 358 cortex 358	Volume clearance, esophagus 238
hypoxia 136	field 358	deficit 170, 173 B
maximum 77 C	binocular 360	compensation 218
endurance athletes 77	pathways 358	shock 218
physical work 74	pigment 344, 348, 350	excess 170, 173 B
hypoxia 136	role in adaptation 350	extracellular → ECF 168
venous return 204	sensors, spatial summation	hypertension 216
Ventilation-perfusion ratio 122	352	interstitial 168
Ventricles, cerebrospinal fluid	temporal summation 352	intracellular → ICF 168
310	thresholds 352	ratio 376
Ventricular (→ also heart)	absolute 352	regulation 170
arrhythmias 200	Vital capacity 112	unit 372
fibrillation 200	forced 118	velocity 374
pressure 190, 202	Vitamin A 226, 252, 260, 350	Voluntary motor function 324
tachycardia 200	deficiency 350	inhibition 326
volume 202	intestinal absorption 260	Voltage clamp 34
work diagram 202 Veratridine 47 B	B ₁ (thiamin) 226, 260	Vomiting 78, 142, 238 center 238
Verbalization 336	B ₂ (riboflavin) 226, 260 B ₆ (pyridoxin) 226, 260	salt and water homeostasis 173
Vergence, eye movement 360	B ₁₂ (cobalamines) 90, 226	B
Vermis 326	absorption 260	von Willebrand factor 102, 103 A
Vertical type (heart) 198	C 90, 226, 260	Vowels 370
Vertigo 328, 342	intestinal absorption 260	V ₁ receptor (ADH), second mes-
Very low density lipoproteins	renal reabsorption 156	senger 276
(VLDL) 254, 256, 260	D 226, 252, 292	V ₂ receptor (ADH) 24
Vesicles 28	daily intake 292	second messenger 276
chromaffin 84	deficiency 262, 292	VR1 receptor (vanilloid receptor
secretory 12, 30	optimal 386	type 1) 314
Vessels, capacitance 188	binding protein (DBP) 158,	Vulnerable phase (myocard) 200
wall tension 188	292	
Vestibular ganglia 342	D ₂ (= ergocalciferol) 226, 292	

nuclei 328, 342

organ 342

reflex 342

 D_3 (= cholecalciferol = calciol)

226, 260, 292

W

W (watt), unit 374 f. Wakefulness 322, 332 Warfarin 104 Warm sensors 222 ff., 314 Water \rightarrow H₂O Watt second, unit 374f. Waves, EEG 332 paroxysmal 332 slow, intestine 244 stomach motiltity 240 W cells, retina 358 Weber's rule 352 Weber's test 364 Weight, unit 374 Wernicke's area 370 Whispering 370 White matter, spinal cord 310 Wilson leads (ECG) 196

Window, oval/round 364, 366 Wind space 370 Windkessel 188, 202 Withdrawal reflex 320 Work, muscle 74 negative dynamic 74 physical 74, 142 positive dynamic 74 pressure/volume, heart 202 static postural 74 unit 374 Writing 330, 336 Ws. unit 374

X

X cells, retina 358 X-chromosome 8, 307 B,C Xenobiotics 160

Υ

Yard, conversion into SI unit 372 Yawning 132 Y cells, retina 358 Y-chromosome 8, 307 B,C Yohimbin 87 B

Z

Zeitgeber, external 334
Zero-current potential 34
Z-plates, muscle 60
Zona fasciculata → adrenal cortex
glomerulosa → adrenal cortex
pellucida 308
reaction 308
reticularis → adrenal cortex
Zone, comfort 224, 226
thermoneutral 224
Zonular fibers 344