Autonomic Nervous System (ANS)

Study Objectives
- To define receptors, autonomic neurotransmitters and blocking drugs, homeostasis, receptors and related concepts.
- To describe the anatomy and the physiology of the sympathetic and the parasympathetic nervous system, the visceral afferent system, the enteric nervous system, transmitter mechanisms in autonomic ganglia and at peripheral receptors, the bladder emptying, the pupillary reflexes.
- To explain the central autonomic control, the autonomic control of temperature, appetite, thirst, the subsynaptic autonomic mechanisms, emotional disorders, the Kluver–Bucy-syndrome.
- To use the above concepts in problem solving and case histories.

Principles
- The autonomic nervous system mediates neural control of the internal milieu despite substantial environmental changes.
- Cannon's law: The peristalsis in the small intestine is polarised, so it always proceeds in the oral-aboral direction.

Essentials
This ANS deals with 1. The autonomic system in general, 2. The sympathetic system, and 3. The parasympathetic system.

1. The Autonomic System In General
The autonomic system directly influences smooth muscles, glands and the heart through its two subdivisions, the sympathetic and the parasympathetic system. The two subdivisions function in a dynamic balance aiming at homeostasis. The enteric nervous system is lying within the walls of the gastrointestinal tract and includes neurons in the pancreas, liver and gallbladder, thus being an entity in itself. However, the enteric nervous system is clearly an important part of the autonomic nervous system that controls gastrointestinal motility, secretion and bloodflow.

The central autonomic system
The central autonomic nervous system outflow arises in the hypothalamus, the brainstem, and the spinal cord. The motor and premotor cortex, the cingulate gyrus and the hypothalamus can modulate the function of the autonomic medullary control neurons in the lateral horn of the grey matter. Circulatory changes during exercise and in various stressful situations are influenced or governed by the cortex and deeper brain nuclei. The central autonomic system also modulates release of certain peptides and catecholamines that affect both blood volume as well as the total peripheral vascular resistance.

The cerebral cortex assimilates all inputs of visual, olfactory, labyrinthine, locomotor origin, as well as from other specialised sensors (stretch receptors, chemo-, baro-, osmo-, and thermo-receptors).

The integration of these inputs into an appropriate response takes place in the hypothalamus and in the ponto-medullary centres. From here the efferent signals pass to the periphery via the sympathetic and the parasympathetic pathways.

The primary afferent projections from the baroreceptors reach the solitary tract nucleus (STN), and from here we have connections to the important dorsal motor nucleus of the vagus. A high baroreceptor activity stimulates the DMNV, so that
the vagal inhibition of the heart is increased. More importantly, the high baroreceptor activity inhibits the sympathetic drive to the heart and vessels thus reducing blood pressure.

The central autonomic structures co-operate in situations of survival character: Fright, flight or fight response, feeding and drinking in starvation, reproduction and sexual satisfaction for continuation of life, thermoregulation at extreme temperatures and emotional behaviour in crises.

**The Fright -Flight Or Fight Response**

Aggression and defence responses are elicited in emergency situations. The sympatho-adrenergic system gives rise to the fright, flight or fight-reactions in acutely stressful situations. The sympathetic reactions dominate over the parasympathetic and the subject is aggressive or anxious. The brain releases corticotrophin-releasing factor to the hypothalamic-pituitary portal system. The hypothalamic-pituitary axis secretes adencorticotrophic hormone, the cardiac rate and contractile force increases, the blood is distributed from viscera to the active skeletal muscles by visceral vasoconstriction and preferential vasodilatation. The subject hyperventilates, the gastrointestinal activity is reduced, and there is increased glycogenolysis and lipolysis. The airways dilatate, and the adrenal medullary (catecholamines) and cortical secretion (cortisol) increases. This response is seen in humans exposed to psychological-emotional stress. Stress in general is comprised of severe emotional and physical burdens (fear, pain, hypoxia, hypothermia, hypoglycaemia, hypotension etc).

Cannons emergency reaction is an immediate sympatho-adrenergic response to life-threatening situations, with both sympato-adrenergic and parasympathetic overactivity. The last phenomenon includes vagal cardiac arrest with involuntary defecation and urination.

**Feeding and drinking**

Bilateral destruction of the ventromedial hypothalamic nuclei leads to hyperphagia and failure of body weight control. Such animals become obese, and they have high plasma insulin.

Bilateral lesions in the lateral hypothalamic regions cause a temporary hypophagia.

The cells of the ventromedial nuclei have a special affinity for glucose, and these cells are responsible for insulin secretion from the pancreatic β-cells. Signals from the dorsal motor nucleus of the vagal nerve increase insulin secretion, and sympathetic stimulation inhibits the release of insulin. The ventromedial nuclei seem to function like a glucostat.

Stimulation and ablations of the limbic system affect food intake. Obviously from clinical practice, psychological factors, emotional disturbances, motivations and conditioned behaviour are all affecting our drive for food intake.

**Sexual behaviour**

Hypothalamic and other limbic system co-operation are responsible for a wide variety of autonomic and somatic phenomena associated with emotions. Stimulation of the midbrain septum yields pleasurable sensations and sexual drive in patients. The dorsomedial nucleus of the hypothalamus is probably a major sex centre responsible for the sexual act. Stimulation of the ventromedial and preoptic regions also releases sexual activities.

**The Thermocontrol**

Thermoreceptors can initiate generalised reactions to heat and cold. The signals from both superficial and deep thermoreceptors must act through the hypothalamus to arouse appropriate, generalised reactions.

Cooling or heating the denervated lower extremities of spinal men evoked vasoconstriction and shivering or vasodilatation and sweating of the innervated upper body shortly after cooled or warmed arterial blood reached the brain. The anterior hypothalamus is responsible for sensing blood temperature variations. The anterior hypothalamus, in particular the preoptic area, has been shown to contain numerous heat-sensitive cells and less cold-sensitive receptors. Such central thermoreceptors are also found at other levels of the CNS. After destruction of the hypothalamus, the midbrain reticular formation takes over the temperature control. Sections eliminating both the hypothalamus and the mesencephalon leave the medulla and spinal cord to control temperature. The posterior hypothalamus does not contain thermoreceptors. Concerning thermocontrol.

**The brain and the immune defence system**
Internal and external stress affects the prefrontal cortex, whereby the limbic system with the hypothalamus is activated. Hypothalamic nuclei release corticotropin-releasing hormone (CRH) to the portal blood. The blood reaches the adenohypophysis, where CRH triggers the release of adenocorticotropic hormone (ACTH), endorphins and met-enkephalin. ACTH works through different pathways in order to protect the body. ACTH stimulates the adrenal cortex to release corticosteroids, which produce immuno-suppression. Immuno-suppression reduces the number of inflammatory effector cells, including helper T cells and killer cells.

On the other hand, cancer therapists assume that relaxed lifestyle and positive reinforcement may have stimulated the immune defence in some patients with malignant diseases, and explain miraculous remissions. Higher brain centres may even affect the reticuloendothelial production of killer cells through the peripheral nerves to the lymph nodes and bone marrow.

*Autonomic nerves* are composed of two neurons termed the preganglionic and the Postganglionic neuron based on anatomical location relative to the ganglion. A preganglionic neuron has its cell body in the spinal cord or brainstem and is modulated by higher centres and by spinal reflexes. The preganglionic axon leaves the CNS from the cranial, thoracic, lumbar or sacral regions and synapse in the autonomic ganglia with the cell body of the postganglionic neuron. The postganglionic neurons innervate the effector organs (Viscera).

Viscera function involuntarily and their activity must be modulated by the autonomic nervous system with excitatory or inhibitory signals. All autonomic nerves have ganglia outside the CNS in contrast to the somatic nervous system, where neural connections are located entirely within the CNS. Most somatic nerves that control motor function are myelinated and have a high conduction velocity, whereas most postganglionic neurons are unmyelinated with a low conduction velocity. However, the preganglionic neurons are mostly myelinated with a high conduction velocity.

*Receptors for neurotransmitters* are specific cellular components, whose interaction with the neurotransmitter, a hormone or a drug produces a biological response in the cell.

*Acetylcholine* (ACh) is the transmitter between the pre- and the post-ganglionic neurons, not only in the sympathetic nervous system, but also in the parasympathetic system. The cholinergic receptors are nicotinic or muscarinic. The cholinergic receptors of the ganglia and in the somatic motor endplate are *nicotinic*. Nicotine and acetylcholine activate nicotinic cholinergic receptors. When the action potential arrives at the preganglionic fibre, acetylcholine is released from
its terminals and diffuses across the synaptic cleft to bind to the specific nicotinic receptors on the membrane of the postganglionic neuron. Nicotinic receptors are linked to cation channels lined with negative charges. These channels open enough to allow mainly hydrated Na\(^+\) to enter the cell rapidly (for about 1 ms) and depolarise the membrane.

**The nicotinic cholinergic receptor**

The resulting current elicits an *excitatory postsynaptic potential* (EPSP). Repolarisation is also fast (ms).

Acetylcholine is also the neurotransmitter for the sympathetic innervation of sweat glands, and they are completely blocked by *atropine*. The acetylcholine receptors of the sweat glands are *muscarinic*, since acetylcholine and muscarine activate them.

**The muscarinic cholinergic receptor**

These slowly working surface-receptors are linked to a long lasting cascade of events starting with binding of the hormone to the receptor, activation of G-proteins, enzyme activation, production of second-messengers, protein kinase g of K\(^+\)-channels, with efflux of K\(^+\), so the membrane is hyperpolarised. In this example, acetylcholine is an inhibitory transmitter.

**2. The Sympathetic Nervous System**

The preganglionic sympathetic nerve fibres originate in small multipolar neurons in the lateral horn of the grey matter in the thoracic and lumbar spinal cord. The central sympathetic outflow converges on these preganglionic neurons. Their axons are thin myelinated fibres that leave the spinal cord through the ventral root. The preganglionic fibres then leave the spinal nerve forming myelinated white rami communicantes, through which they reach the nearest ganglion in the paravertebral ganglia of the paired sympathetic trunk. Typically, each fibre will end here forming synapses with up to 20 postganglionic neurons. A few preganglionic fibres pass the sympathetic trunk without interruption to form the splanchnic nerves that reach the three unpaired prevertebral ganglia (coeliac = solar plexus, superior mesenteric and inferior mesenteric) of the lower intestinal and urinary organs. Most sympathetic ganglia are remote from the organ supplied. The postganglionic fibres are all unmyelinated, and they leave the sympathetic trunk through the grey rami communicantes and thus reach the effectors supplied by the sympathetic system. The effectors are the smooth muscles of all organs (blood vessels, viscera, lungs, hairs, pupils), the heart and glands (sweat glands, salivary and other digestive glands). In addition, the sympathetic postganglionic fibres innervate adipocytes, hepatocytes and renal tubular cells.

The sympathetic system is a functional and phylogenetic unit of the sympathetic system and the adrenal medulla. The adrenal medulla is a modified sympathetic ganglion. Any increase in sympathetic activity increases the secretion of adrenaline and noradrenaline from the medulla into the circulation. The preganglionic fibres to the adrenal medulla pass all the way to the special postganglionic cells in the adrenal medulla. The synapse is cholinergic (nicotinic) as it is for all preganglionic synapses. The postganglionic cells of the adrenal medulla have developed to cells filled with chromaffine granules, and are called chromaffine cells. These cells do not conduct signals, but synthesise adrenaline (and noradrenaline) which is released into the blood. Sympathetic stimulation triggers the conversion of tyrosine to dihydroxyphenylalanine (DOPA). A non-specific decarboxylase catalyses the conversion of DOPA to dopamine, which is taken up by the chromaffine granules in the cells. The granules contain the crucial enzyme, dopamine \(\beta\)-hydroxylase. This enzyme is activated by sympathetic stimulation, and catalyses the formation of noradrenaline from dopamine.

A few granules store noradrenaline (NA), while the remaining granules liberate NA to the cytosol, where NA is methylated by phenylethanolamine N-methyltransferase to adrenaline. Adrenaline is taken up by chromaffine granules and stored as the predominant adrenal hormone.

**Adrenergic Receptors**

The sympathetic system exerts either excitatory or inhibitory actions through adrenergic receptors. Adrenergic receptors are *membrane-receptors*. The dual response to adrenergic stimulation was known before Ahlquist in 1948 proposed that adrenergic receptors could be divided into two groups, \(\beta\) - and \(\alpha\) - receptors, on the basis of *blocking drugs* (Box 6-1). The basic idea of Ahlquist is that noradrenaline (NA) act predominantly on *vasoconstricting* \(\alpha\)-receptors, and isoprenaline (Iso) predominantly on *vasodilatating* \(\beta\)-receptors. Both types of receptors are stimulated by adrenaline (Ad).

The rank order of sensitivity of a series of chemically similar compounds for activating a receptor (agonists) or inhibiting the receptor response (antagonists) is considered diagnostic of the receptor subtype. More and more closely related
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Dr. Salah

3. The β-receptors are distinguished, so there is already three subtypes for each of the following receptors: $\alpha_1$ABC-receptors, $\alpha_2$ABC-receptors, and $\beta_23$-receptors.

1. The $\alpha_1$-receptors are blocked by Phenoxybenzamine and Phentolamine.

The $\alpha_1$-receptors are located on the surface of target cells (vascular smooth muscle, sphincter muscles of the gastrointestinal tract and bladder, and radial iris muscles). They are highly sensitive to NA, less sensitive to Ad, and almost insensitive to isoprenaline.

The $\alpha_1$-receptors act through phospholipase C and through intracellular [Ca$^{2+}$] elevation. Ca$^{2+}$ binds to calmodulin in the cytosol. The complex activates protein kinase, which catalyses the phosphorylation of proteins. They become enzymatically active, and trigger vasoconstriction.

In contrast, the presynaptic $\alpha_2$-receptors are located on the presynaptic membrane (sympathetic end bulbs). NA released into the synaptic cleft diffuses to the $\alpha_1$-receptors on the target cells, but part of the NA diffuses back to the $\alpha_2$-receptors on the presynaptic nerve terminals. Here, NA activates membrane adenylcyclase, reducing [cAMP] in the cells, and thus inhibiting release of more NA from the vesicles by negative feedback. Hence, a function of $\alpha_2$-receptors is auto-inhibitory feedback. These receptors are also found in gastric smooth muscle cells and the $\beta$-cells of pancreatic islets. Stimulation decreases gastric motility and attenuates insulin secretion.

2. The $\beta$-receptor is blocked by propranolol.

The $\beta$-receptors are located on effector cells that are most sensitive to isoprenaline, but less so to Ad and NA. All $\beta$-receptors act through activation of adenylcyclase and cAMP. $\beta_1$-receptors are equally sensitive to NA and Ad, whereas $\beta_2$-receptors are more sensitive to Ad than to NA.

$\beta_1$-receptors are located in the myocardium - primarily on pacemaker cells. The $\beta_1$-receptors of the heart are stimulated by NA which increases cAMP production with increased chronotropic (increased heart rate) and inotropic effect (increased force). Heart patients use Cardioselective $\beta_1$-blockers such as Metoprolol, because Metoprolol decreases cardiac arrhythmias and tachycardia.

$\beta_2$-receptors are found primarily on bronchiolar smooth muscle cells, vascular smooth muscle, uterine smooth muscle, salivary glands, the intestine and the liver. When NA binds to $\beta_2$-receptors, it causes inhibition of the target organ. Therefore, NA causes vasodilatation, bronchodilatation and uterine relaxation. Similarly, sympathomimetics such as $\beta_2$-stimulators (salbutamol) increase cAMP production, resulting in bronchodilatation, increased salivary secretion, uterine relaxation and enhanced hepatic glucose output. $\beta_2$-stimulators are used to eliminate bronchial asthma attacks.

Butoxamine is a selective $\beta_2$-blocker.

The near-vision response is also called the convergence response. Near vision - even with only one eye - triggers accommodation and pupillary contraction. The ciliary muscle and the pupillary sphincter muscle are innervated of the parasympathetic oculomotor nerve, and the two muscles (with M-receptors) contract simultaneously for near vision. This leads to increased refractive power or accommodation, and to pupillary contraction (miosis).

When a person closes his eyelids the pupils enlarge, and when he opens the pupils again the pupils become smaller. This is due to the pupillary light reflex, where retinal ganglion cells are stimulated by light, send signals through the optic nerve to the olivary pretectal nucleus neurons. These light-sensitive neurons are connected to the parasympathetic preganglionic neurons in the oculomotor Edinger-Westphal nuclei on both sides. The light reflex contracts the pupillary sphincter muscle. Argyll-Robertson’s pupillary syndrome refers to small, light-refractive pupils with maintained convergence response to near vision. The syndrome is seen in neurosyphilis, when the pupillary light reflex is spoiled by interruption of the fibres from brachium to the olivary pretectal nucleus neurons.

Sympathetic preganglionic neurons in the intermediolateral cell column of segment T1-T2, send ascending axons to the superior cervical ganglion. Postganglionic axons follow the ciliary nerve into the eye. The nerve terminals end on $\alpha_1$-receptors on the dilator pupillae muscle, and noradrenaline is neurotransmitter. The sympathetic fibres also contain vasoconstrictors to the facial skin and stimulate facial sweat glands (see Horners syndrome).

Catecholamines are substances consisting of catechol (an aromatic structure with two hydroxyl groups) linked to an amine.
Catecholamines increase heart rate and cardiac output by stimulation of the adrenergic $\alpha_1$-receptors in the myocardium. Catecholamines, released by the adrenal medulla, support the sympathetic system by modifying the circulation during exercise. During exercise the blood is directed to the working muscles from other parts. Noradrenergic nerve fibres innervate blood vessels all over the body. Sympathetic innervation accounts for vascular tone and vasoconstriction.

The most important exercise response in humans is a tremendous vasodilatation in the vascular bed of muscles. The vasodilatation is probably due to a decrease in the $\alpha$-adrenergic tone of muscular arterioles, and to the action of adrenaline on $\beta_2$-receptors.

Catecholamines dilate the bronchial airways by stimulating adrenergic $\beta_2$-receptors. They increase both tidal volume and respiratory frequency. The result is increased ventilation. Catecholamines acting on $\beta_1$-receptors cause increased cardiac output. Catecholamines relax the smooth muscles of the digestive tract ($\beta_2$-receptors), but contract the sphincters. Catecholamines stimulate metabolism (by activation of the thyroid hormone, $T_3$) and lipolysis. Adrenaline stimulates hepatic glycogenolysis via $\beta_2$-receptors.

Finally, adrenaline stimulates the ascending reticular system (i.e., the reticular activating system or RAS) in the brain stem, thus keeping us alert and causing arousal reactions with desynchronisation of the EEG.

The resistance vessels of the striated muscles in hunting predators (and perhaps in humans) are also innervated by another system. This is the cholinergic, sympathetic vasodilator system. It is capable of a rapid and appropriate bloodflow response during hunting.

Acute stress activates the splanchnic nerves and liberates large amounts of adrenaline from the medulla. Diabetics who are developing acute hypoglycaemia, secrete large amounts of catecholamines. Acute muscular activity starts a large catecholamine secretion in exercising persons.

Besides catecholamines, ACTH is also released during stress by increasing hypothalamic signals. ACTH stimulates the glucocorticoid and to some extent the mineralocorticoid secretion through cAMP. Small amounts of glucocorticoids are permissive for the actions of catecholamines.

Plasma catecholamines are rapidly removed from the blood and have a half-life in plasma of less than 20 s. This is the combined result of rapid uptake by tissues and inactivation in the liver and vascular.

**The Autonomic Control Of The Cardiovascular System**

The brainstem is the primary site for the autonomic cardiovascular control.

High-pressure baroreceptors are distension-activated stretch receptors located in the walls of the carotid sinus and the aortic arch. Increased arterial blood pressure increases the signal frequency in the sensory baroreceptor neurons that project into the medullary cardiovascular centre (i.e., the solitary tract nucleus, nucleus ambiguus and the dorsal motor nucleus of the vagus, DMNV). Impulses generated in the baroreceptor neurons with increasing blood pressure, activate the vagal efferents to the heart and inhibit the sympathetic tone towards the heart. As a consequence the heart rates and force of contraction decreases. Impulses generated in cardiac baroreceptors by cardiac filling, also activate the force of contraction. The postganglionic fibres from the 3 upper cervical ganglia of the sympathetic trunk pass to the heart as the cardiac nerves to the cardiac plexus.

**The autonomic control of blood pressure and heart rate.**

The sympathetic effect is dominant. Increased sympathetic activity constricts the veins, which increases cardiac output by augmenting cardiac filling. Arteriolar constriction reduces cardiac output by increasing the arterial blood pressure (i.e., afterload). Other sensory inputs from skeletal muscles, lungs, gastrointestinal viscera, hypothalamus and forebrain help to co-ordinate the autonomic cardiovascular responses related to exercise, respiration, and feeding and temperature control. Hormones, such as angiotensin II, can also modulate the autonomic responses through neurons in the circumventricular organs of the brain. These organs (such as the area postrema) lack the blood-brain barrier.

The sympathetic system innervates the sinus node, the coronary vessels and the myocardial syncytiun. Each fibre ends in many terminals, and from the terminals the transmitter noradrenaline is released to the $\beta_2$-receptors of the smooth muscle cells of the coronary vessels and of the myocardium. As a result of increased sympathetic tone, the contractility of the myocardium is increased. Thus, the end systolic volume falls from its usual volume- as an example from 70 ml to 40 ml, and the end diastolic volume increases due to increased venous return of blood from 140 to 180 ml. Hereby, the
stroke volume is increased from 70 to 140 ml of blood in the example. A combination of the doubling of stroke volume with a threefold increase in heart rate, results in a 6-fold rise in cardiac output.

Sympathetic stimulation depolarizes the sinus node, so that the threshold potential is reached faster than normal. Hereby, the heart rate is increased, and may reach 220 beats/min in young persons. Such a high frequency is due to a maximal sympathetic activation of the heart combined with a reduction of the vagal tone.

**Sympathetic Activation**

Activation of noradrenergic fibres leads to peripheral sympathetic vasoconstriction, so that blood is shunted to central areas. The heart is stimulated through \(\beta_1\)-receptors so that its frequency and contractility is increased. Other organs are also stimulated to make the person fit for fight or flight in any stressful situation.

The postganglionic sympathetic fibres have noradrenaline and ATP containing vacuoles in their nerve terminals. Hence, they release noradrenaline and ATP. The noradrenaline is produced in the chromaffine granules of the neuron.

**A sympathetic ganglionic synapse with a small intensity fluorescent cell (SIF cell).**

Acetylcholine is released from the preganglionic cell and binds to nicotinic receptors on the postganglionic cell.

Acetylcholine also binds to small SIF cells with muscarinic receptors and vesicles that contain dopamine. Dopamine interacts with dopamine receptors (D2 and D4) on the postganglionic cell and modulates ganglionic transmission by increased permeability to small ions and hyperpolarisation.

Liberation of noradrenaline and ATP to the blood does not only lead to constriction of arterioles and arterial vessels, but also constriction of veins and venules. Without venous constriction, the large venous compliance would cause an inordinate amount of blood to be stored in the veins upon sympathetic arteriolar constriction. The consequence would be decreased venous return, which decreases cardiac output and perfusion of vital organs.

Activation of presynaptic purine receptors by adenosine inhibits adrenaline release from the postganglionic terminals innervating the blood vessels. This results in massive vasodilatation.

Exercise and stress demand mobilisation of energy to muscles and heart. Activation of \(\beta_2\)-receptors in the arteriolar wall by circulating catecholamines from the medulla also contributes to vasodilatation in the striated muscles. The total peripheral vascular resistance is reduced during exercise to 20-30% of resting values.

During stress the cutaneous circulation is reduced at first, but then the cutaneous bloodflow rises due to the increased heat production. The brain vessels are only modestly constricted by sympathetic stimulation.

**3. The Parasympathetic System**

The parasympathetic system has two subdivisions. The cranial division in the brainstem innervates the blood vessels of the head and neck and of many Thoraco-abdominal viscera. The sacral division in the sacral cord innervates the smooth muscles of the walls of the viscera and their glands (the large intestine, liver, kidney, spleen, the bladder and the genitals).

The parasympathetic system only innervates a small percentage of the resistance vessels. Only arteries in the brain and of the penis, the clitoris, and the labia minora receive parasympathetic innervation. Hence, the parasympathetic system has a minimal effect on the arterial blood pressure.

Parasympathetic fibres travelling in the vagus nerve are of utmost importance in affecting the cardiac rate. Vagal fibres innervate the sino-atrial- and the atico-ventricular-nodes as well as the atrial muscle walls.

The parasympathetic system also innervates the tear and the salivary glands, and the muscles within the eye.

Excitation of the vagus decreases heart rate and atrial contractile force, increases intestinal motility, contracts the gall bladder and bronchi, and relaxes the sphincters of the gastrointestinal tract. The vagal decrease in heart rate is due to the rhythm shift to special P cells, which have a slow rate of depolarisation. Acetylcholine (ACh) is liberated on the cardiac cell membranes, ACh-activated K⁺ - channels are opened (via cholinergic receptors and G-regulatory proteins), and K⁺ leaks out of the cells, thus opposing the pacemaker current. Vagal stimulation slows down the AV-conduction, causing the co-ordination of atrial and ventricular rhythm to be disrupted. Vagal stimulation can lead to death. Thus external massage of the carotid sinus can cause collar death by greatly increasing vagal stimulation.
The effect of acetylcholine released in the autonomic ganglia can be simulated by nicotine. Conversely, the effect of acetylcholine released by parasympathetic nerve terminals at the target organs can be simulated by muscarine. These observations suggest the presence of two different types of cholinergic receptors. Cholinergic receptors are activated by ACh and by metacholine (MeCH).

The most important ganglionic blocking drug for blockade of both sympathetic and parasympathetic transmission is hexamethonium.

Cholinergic receptors are located in all autonomic ganglia (nicotinic type), in postganglionic terminals at target organs with parasympathetic innervation (muscarinic type), and in the motor endplate (nicotinic type).

Nicotinic receptors are those activated by acetylcholine, nicotine and nicotine agonists (ex. dimethylphenylyperazine, DMPP). Nicotine stimulates all autonomic ganglia simultaneously. Hence, sympathetic vasoconstriction in the limbs and viscera is accompanied by increased gastrointestinal activity and slowing of the heart via the vagus. Nicotinic receptors are blocked completely by d-tubocurarine, and hexa- or decamethonium (Box 6-3). The motor endplate has a different type of nicotinic receptor than the ganglia, since its receptors are not blocked by hexamethonium, but are blocked by d-tubocurarine and decamethonium.

Acetylcholine, muscarine and muscarinic agonists (pilocarpine and carbacholine, CCh), activate muscarinic receptors. At least 5 different muscarinic receptor molecules have been identified (M₁, M₂, M₃ .). Activation of the M₁ type. Activation of the M₂ type activates an inhibitory G-protein, which inhibits adenylcyclase. Muscarinic receptor activation is linked to G-protein activation and second-messenger systems.

The dominating receptors in the heart are the M₂ receptors mainly in the coronary circuit and the M₁ receptors elsewhere. The typical effects of acetylcholine are hyperpolarisation and reduced spontaneous depolarisation of the sinusoidal cells leading to reduced pacemaker activity. Besides, the cardiac contractility and conduction velocity are reduced (Dhein et al, 2001).

Muscarinic receptors are blocked completely by atropine, and by antimuscarinic drugs such as homatropine and scopolamine (Table 6-3). These drugs do not block the nicotinic effect of ACh on the postganglionic neurons or on the motor endplate.

1. The sympathetic system consists of short preganglionic and long postganglionic nerve fibres. The parasympathetic system contains long preganglionic and short postganglionic fibres.
2. The chemical transmitter at the target organ is noradrenaline in the sympathetic and acetylcholine in the parasympathetic system.
3. The sympathetic system contains adrenergic receptors (ıı and ıı), whereas the parasympathetic system has cholinergic receptors (muscarinic or muscarinergic and nicotinic or nicotinergic).
4. Activation of the cholinergic system serves anabolic functions (ie, stay and play), whereas activation of the noradrenergic system serves catabolic functions (ie, fight, fright or flight).
5. Activation of ı₁-receptors increases intracellular [Ca²⁺], which leads to phosphorylation of protein kinases and thus to a response. Activation of ı₂-receptors triggers an inhibition of the membrane adenylcyclase, reducing [cAMP] in the cells. ı₁- and ı₂-receptors activate adenylcyclase, which increases cAMP production in the cell. Muscarinic receptors are completely blocked by atropine. Activation of M₁-receptors increases intracellular [Ca²⁺]. Activation of M₂ inhibits adenylcyclase, and through an inhibitory G-protein reduces the formation of cAMP.