Distribution of NADPH-diaphorase and AChE activity in the anterior leaflet of rat mitral valve

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Abstract

The mitral valve, as an active flap, forms the major part of the left ventricular inflow tract and therefore plays an important function in many aspects of left ventricular performance. The anterior leaflet of this valve is the largest and most ventrally placed of two leaflets that come together during ventricular systole to close the left atrioventricular orifice. Various neurotransmitters are responsible for different functions including controlling valve movement, inhibiting or causing the failure of impulse conduction in the valve and the sensation of pain. Nitric oxide acts as a gaseous free radical neurotransmitter, neuromediator and effective cardiovascular modulator. Acetyl-choline is known to function as a typical neurotransmitter. Histochemical methods for detection of nitric oxide-synthase marker, and method for detection of acetylcholinesterase (ACHE) were used. Both methods were performed on the same valve sample. A widespread distribution of nerve fibres was observed in the anterior leaflet of the mitral valve. The fine NADPH-d positive (nitrergic) nerve fibres were identified in all zones of valve leaflet. AChE positive (cholinergic) nerve fibres were identified forming dense network and fibres organized in stripes. Endocardial cells and vessels manifested heavy NADPH-d activity. Our observations suggest a different arrangement of nitrergic and cholinergic nerve fibres in the anterior leaflet of the mitral valve. The presence of nitrergic and cholinergic activity confirms the involvement of both neurotransmitters in nerve plexuses and other structures of mitral valve.

Introduction

Heart valves are critical structures in the function of the heart, and the surgical repair of diseased valves has become a major form of treatment for patients with valve disease. Paradoxically, little is known about the structural components that are critical factors in the function of the heart valves. However, the evidence linking valve innervation with functional integrity of the heart valves or stabilizing correlations between innervation and valve dysfunction is meagre. From the anatomical and physiological points of view, the anterior leaflet of the mitral valve (anterior mitral leaflet, or the anterior cusp of the left atrioventricular valve) forms an important boundary of the left outflow tract. This might imply that this structure is a possible candidate for a special sensory area among the leaflets and it is shown to be a highly active structure in the beating heart. The two leaflets of the mitral valve are actively contractile; the physical forces generated in the valve itself may stabilize and add precision to the sum of forces regulating valve movement. This is a critical significance both in the moments preceding and during the valve opening and closing.

Although the function alteration of the cardiac autonomic nervous system has been proved to be a powerful predictor of cardiac death or serious arrhythmia in patients with cardiac disease, little is known yet about the mechanisms regulating this system. Despite the increasing evidence about the importance of cardiac autonomic function abnormalities in heart disease, the controlling mechanisms remain poorly understood.

The mammalian and human hearts contain a variety of morphologically distinct nerve fibres known to influence cardiac function. Just the cardiac valve tissue has been shown to be innervated by nerve fibres containing active sympathetic, parasympathetic and peptidergic neurotransmitters. Recent studies suggest that nitric oxide (NO) may play an important role in the control of this regulatory system and that neural regulation of coronary blood flow and cardiac conductance is essential for the maintenance of cardiac function. NO as a gaseous free radicals acts like a neurotransmitter and effective cardiovascular modulator. This gas plays a fundamental role in cardiovascular physiology and pathophysiology. Within the cardiovascular system, NO participates in the regulation of coronary blood flow and tension of vessel wall. It is released not only from endothelial cells (ECs), but also may serve as a non-adrenergic, non-cholinergic neurotransmitter in cardiac innervation. The regulators of NO production are physical and chemical stimulators transmitted by the vessel wall and by endocardium as well. NO influences all neural substrates that contribute to the generation of the autonomic nerves activity in the central and peripheral nervous system. NO is synthesized from its precursor L-arginine by the enzyme NO synthase (NOS), which occurs in three distinct isoforms. These consist of the constitutive calciun-dependent enzymes endothelial NOS (eNOS, originally identified in vascular endothelium, but also present in platelets, myocardium and endocardium) and neuronal NOS (nNOS, present in neuronal tissue) and the calcium-independent, cytokine-inducible NOS (iNOS, expressed in macrophages and other tissues during immune stimulation). In the vascular and cardiac tissue it is thus constitutively produced by eNOS or nNOS, which has been shown to be identical with NADPH-diaphorase under condition of tissue fixation. Reduced NADPH-d may be used as a marker for NOS, since NOS and NADPH-d are extensively co-localized. Although the NADPH-d reaction is not completely specific, it is nonetheless highly selective for NOS activity in the heart. There are no other diaphorases to confound localization in this organ as an evidence of the reduction of nitro blue tetrazolium (NBT) to the formazan insoluble in water.

NO is described also as a neurotransmitter and neuromodulator in the vagus nerve, similarly to an acetylcholine. It has additional effects on the level of parasympathetic post-ganglionic nerve terminals that may be particularly important within the heart. Many studies suggest that NO modulates cardiac vagal control, increase the activity of central vagal motoneurons and, more contentiously, contribute to the bradycardic effects of vagal stimulation. Overall, a picture of NO as a sympatholytic and a vagotonic agent does emerge from the animal data.

Acetylcholine (ACH) is known to function as a typical and primary chemical neurotransmitter. Although both divisions of the autonomic nervous system are crucial for the heart regulation, the anatomic distribution of cholinergic

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(parasympathetic) nerves is less precisely known than that of sympathetic innervation. AChE is an important component of the heart’s cholinergic system; it is known to regulate the cardiac parasympathetic responses by controlling acetylcholine levels. According to Williams and Jew, the rat mitral valve shows that the contractions and relaxations of valves are nerve-mediated.

Therefore, the purpose of the present study was to confirm and compare the distribution of both NADPH-d and AChE enzymes in the anterior mitral leaflet of the rat heart. The presence and especially the mutual relationship of both enzymes can contribute to understanding of NO and ACh neurotransmitters in functioning of mitral valves.

Materials and Methods

Groups of animals used in experiment

Twenty male white Wistar rats, aged 3 months, with a mean weight of 250-350 g were used in this study. Rats were housed in cages and maintained under standard conditions at 20°C, with 12 h light/dark cycle and water and food ad libitum. All experiments were conducted in accordance with European Directive for the Protection of Vertebrate Animals Used for Experimental and Other Scientific Purposes (86/609/EU) and approved by the State Veterinary and Food Administration of Slovak Republic by decision 1696/07-221a.

Samples of tissue and fixation

Animals were anaesthetized with pentobarbital (30 mg/kg). A midline abdominal incision, followed by two lateral costal margin incisions exposed the diaphragm, which was then incised. The thoracic cavity was opened and the heart was perfused with a saline solution and 4% paraformaldehyde with 0.1% glutaraldehyde in 0.1 M phosphate buffer (PB), pH 7.4. The heart was removed, the left ventricle rapidly opened and the valves were cut from the atrioventricular ring. The sample material was placed in the same fixative for 2 hours. The fixed tissue was transferred overnight into 30% saccharose in the same PB at 4°C. For double staining of NADPH-d and AChE, the valves were first stained for NADPH-diaphorase and then selectively blocked the enzymatic activity of AChE. The doubly stained valves were rinsed in distilled water, mounted on glass slides by a whole-mount stretch technique, dried overnight and covered with Entellan (Merck, Germany). The preparations were observed under a light microscope Optica B600 TI with photo Moticam 2300 and Motic images Plus, 2.0 ML.

Results

For descriptive purposes, the innervation of the mitral valve (anterior mitral leaflet) was divided into 3 zones. The basal zone lies close to the fibromuscular (atrioventricular) ring. The intermediate zone is in the middle third of the mitral valve. The distal zone lies close to the free edge of the valve and attachment of the chordae tendineae; it is a free moving zone of the mitral valve (Figures 1, 2).

In whole-mount preparations of the mitral valves, the majority of NADPH-d staining was seen in the anterior leaflet only. Under physiological condition with various intensity, vessels (Figures 3 A,C), fine nerve fibres (Figures 3C; 4C; 5A,B) and endothelial cells of the endocardium (Figures 3A,C; 4A,B; 5A,B) in the examined leaflet were stained from light blue to dark blue colour.

NADPH-d positive arterioles and capillaries were localized in the attachment (basal) zone of the anterior mitral leaflet (Figures 3 A,C). The fine NADPH-d positive (nitrergic) nerve fibres were identified in basal zone running close to the vessels and in the tunica adventitia as well (Figure 3C). Most endocardial (endothelial) cells (EECs) were only lightly NADPH-d stained in basal zone, but a small

Figure 1. The mitral valve of the rat heart. The picture shows the zone adjacent to the fibromuscular ring (arrow on the left) and the free moving zone (arrow on the right), (original magnification x 5).

Figure 2. Scheme of the mitral valve. Drawing of the valve segment depicting the basal, intermediate and distal zones (arranged according to Ahmed et al.)
amount of EECs and nerve fibres were stained moderately in this zone (Figures 3A,B,C).

In the intermediate zone and especially in the distal zone, nitrergic nerve fibres were running along margins of EECs or crossing them only in the direction of blood flow (Figures 4C; 5A,B). Blue stained endothelial cells are seen in different colour on the background (3A: small arrow down; 3C). AChE positive nerve fibres are identified forming both a dense network and fibres organized in stripes (3A: big arrow; 3B). Calibration bars: 3A, B 20 μm, 3C 10 μm.

Discussion

It is well known that the mammalian heart displays regional distribution of autonomic innervation, whereby branches of sympathetic and parasympathetic nerves influence on certain cardiac regions in a different extent,6,27 also depending on individual age and health status.28-30 The occurrence and distribution of different neurotransmitters and neuromediators were investigated in the heart in different species of animals by investigating the presence of acetylcholine,23,31 nitric oxide, neuropeptide Y, tyrosine hydroxylase and calcitonin gene related peptide.1,31-34

The co-localization of NADPH-d and nNOS in mitral valve leaflets was characterized also by Olsen et al.15 Reports by Ursell and Mayes,22 Shoba and Tay,35 Kukanova and Mravec 27 have studied the localization, histology and function of cardiac neurones that contain a marker for NADPH-d and NOS. There is also strong histo-
chemical evidence for the presence of NOS throughout cardiac autonomic neurones. Cardiac innervation is further complicated by the presence of intrinsic neurones, which are known to be capable of significant interaction with both efferent and afferent (sensory) neurones.2 The NADPH-d activity in subpopulation of cardiac neurones was reported by Maifrino et al.26 NADPH-d positive neurones in the heart could provide another source of NO, in addition to endothelial and endocardial cells, and contribute to the actions of NO on the heart. This is an evidence that NO supports and plays a role in the local control of the heart by means of intrinsic neurones.27

The most relevant physiological stimuli for the release of NO from endocardial endothelial cells may be flow-induced shear stress and the cyclical mechanical deformation of the endocardium that occurs during the cardiac cycle.4,14 The most obvious structural response of endothelium to shear stress is to change cell shape and its orientation.3,9,23 Endocardium of rat is characterized by a very thin subendoendocardial layer of connective tissue. As stated above, this may cause possible alteration in shape and orientation of EECs. The blood flow can function as a stress factor to the valve surface, especially in the free moving part of the valve leaflet.

The presence of strongly positive NADPH-d reaction was determined in the blood vessels of rat heart by Ursell and Mayes.22 NADPH-d positive fibres were relatively abundant also in the sinus node, but were very sparse in the atrioventricular node and in the right bundle branch.4 Nevertheless, it is possible that NADPH-d reactive nerve fibres have direct influence on the function of the valves, the frequency and force of valve closure and the cardiac contraction; they may play a role in precipitating arrhythmias and premature extrasystoles.3,18 An accurate and full understanding of the precise movements of the valve leaflets and the mechanisms regulating these movements is likely to provide the information to understand and development of treatment for many different cardiac valve problems, including mitral valve diseases such as prolapse and myxomatous degeneration.15

It is generally known that previous studies,3,21,22,23 have used acetylcholinesterase (AChE) histochemistry to identify cholinergic neurones in the heart. Cholinergic nerve terminals were localized predominantly in endocardial plexuses of the atria and in the left ventricle. According to Williams et al.,6 the nerve plexus of the atrioventricular valve is possessing primary, secondary and tertiary components and this terminology was applied in more recent studies.1,20

At this time, we have no complete information of both nitrergic and cholinergic innervation, nitrergic and cholinergic co-expression in rat heart valves. However, the main functional role of nitric oxide in peripheral cholinergic modulation is known.4,14 The study of Yoshida and Todt4 in the intracardiac plexus of monkey and canine presents also the co-localization of NOS-positivity and AChE in neurones. Some small neurones were intensely stained for NADPH-d activity, but they were not co-localized with AChE. Our observations confirmed both different localization of NADPH-d and AChE positive nerve fibres and their different course in regions of rat valve leaflet. These findings indicate different targets and functions of neurotransmitters in these nerve structures. The demarcation between regions of the valve innervated by cholinergic versus nitrergic nerve fibres may also correlate with the line of leaflet closure. The anterior leaflet of mitral valve, as a neurally-controlled tissue, provides thus a clinically relevant model for the investigation of functional role of nervous system in normal and abnormal heart valve action.3,18

Our observations confirmed different arrangement of nitrergic and cholinergic nerve fibres in the anterior leaflet of mitral valve in rat. The presence of nitric and cholinergic activity in nerve plexuses of atrioventricular valve confirms their involvement in modulation of neurotransmission. These findings can contribute to further study of valve innervation in some cardiac diseases (valves stenosis and insufficiency). Our results suggest that nitric oxide and acetylcholine may also play a role as neurotransmitters and/or neuromediators in neural control of the cardiac blood flow and impulse conduction.

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