The mammalian heart – towards an understanding of the intrinsic antagonistic function of the ventricles

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Abstract

The mammalian myocardium making up the ventricles exhibits, on a microscopic scale, a lamellar structure. Lamellae consist of end-to-end connected myocytes in the form of long strains that are densely crosslinked by bridging myocytic offsprings and strongly bound by the endomysium. In contrast, between lamellae only weak myocytic crosslinking and sparse connective tissue is found. The latter property enables reorientation of lamellar segments during systole. Due to the overall crosslinking, however, the myocardium represents a continuum, albeit with significant local variation. While most lamellar segments are oriented and contract during systole in a mostly surface-parallel direction, enabling the ejection of blood, there is a notable presence of transversely oriented lamellar segments. Such segments develop a partly antagonistic function (with respect to the one of the surface parallel segments), in that they counteract systolic wall thickening, the major mechanism for the ejection of blood. This so far underestimated intrinsic antagonistic organisation is substantiated by anatomic analyses; furthermore, significant consequences with respect to cardiodynamics and diseases of the heart are found.

Introduction

Antagonism is a basic feature in biology manifesting itself ubiquitously, e.g., coagulation of blood is antagonized by hemolysis, efferent and afferent nerves act antagonistically, etc. In particular, muscles, acting in essence unidirectionally, are usually associated with an antagonist.

In contrast, the four-chamber structure heart muscle seems to be void of an antagonistic counterpart at first sight. Its apparent function consists of ejecting blood, an antagonistic function seems to be unnecessary. Yet, an in-depth analysis of anatomy and function of the mammalian ventricles reveals that a particular form of antagonism is intrinsically present and exhibits important properties in view of physiologically well-ordered cardiodynamics.

Due to its physiological (also clinical) dominance, the left ventricle is in the foreground of our considerations. The anatomy of the human ventricles, along with their mechanical function (cardiodynamics) has been subject of research through centuries. Yet, there are still unresolved issues in this field. Well established is the fact that the human myocardium exhibits on a microscopic scale a lamellar structure [1]. Lamellae are thereby composed of long strains of multi-branching cardiomycocytes and are highly diverse with respect to dimensions. While lateral dimensions are generally in the sub-millimeter range, the length of individual lamellae is virtually unlimited; lamellae can be followed around the entire ventricle along numerous paths generated by inter-lamellar branching. Lamellae are intrinsically strongly bound by the dense endomysium and cardiomycocytes may have up to 7 offsprings connecting adjacent strains. Frequent inter-lamellar branching connections also exist, but bondage is quite loose [2-4]. The inter-lamellar perimysium and cross-bridges thereby allow for appreciable mutual displacement during the heart cycle (cleavage planes). Further constituents of the myocardium consist of the electrical conduction system, the intracellular matrix and the vasculature.

Likewise, well established is the general global helical architecture of the left ventricle: A helical arrangement of the lamellar units from the base to the apex is dominating.

There exists some controversy however with respect to the intermediate structure of the myocardium, i.e. between the global and macroscopic and microscopic scale. This bears enormous significance for cardiodynamics. Two main conceptual architectures have been proposed in the past. Most often put forward is the notion that all cardiomycocytes have a spatial orientation that is in essence surface-parallel [5,6]. The systolic action of the heart seems to support this view as a natural premise since the cardiomycocytes act essentially as uniaxial contraction elements. Within the framework of a further concept, also apparently suggested by the systolic constriction property of the ventricles, a band-like structure of the myocardium has been postulated [7]. Yet, both concepts can not be supported by either anatomical facts or cardiodynamics’ considerations. The myocardium is best understood as a netted continuum with substantial local variability; in fact, due to lamellar crosslinking by myocytic offsprings, every location in the ventricular wall can be reached without leaving myocytic strains.

Ventricular anatomy

Gentle pneumatic distention of the excised ventricles allows to visualize the cleavage system by CT [8] and outline the myocardial architecture (Figure 1).

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Quantification of the spatial orientation of lamellar segments in the mm range shows that deviations of more than 30° degrees in diastole and 40 degrees at end-systole from a surface-parallel orientation are observed in up to 40% of all discernable segments [9-12] (the well-known helical ± 60° distribution of the myocytes’ deviation from a plane perpendicular to the long axis is also confirmed). A strong stochasticity is furthermore associated with the local orientation distribution as well as with the ventricular inner and outer surface. At first sight, some similarity with a bird’s nest becomes apparent (Figure 2).

Cardiodynamics

Antagonistic action within the ventricularis wall

There are two major cardiodynamic causes that underline the significance of the findings outlined above. First, lamellar segments, more precisely cardiomyocytes, having a circular, surface-parallel orientation provide in essence the thrust for the systolic ejection of blood. As a result of circular constriction, mural thickening is induced that in fact is one major mechanism causing ejection. Contractile elements, however, that exhibit a strong deviation from a surface-parallel direction, produce a constrictive force component in radial direction, thereby counteracting mural thickening [13]. The effect consists of a significant antagonistic (with respect to the circular constriction) action. The two components generate different types of force tracings when measured by miniaturized intramural force-probes (Figure 3). In agreement with Starling’s law, lamellar segments that are oriented in a surface parallel alignment, unload during systolic shortening. In contrast, the forces produced by lamellar segments in a transmural arrangement increase as such segments elongate in the course with mural thickening.

There is evidence, furthermore, that the antagonism represents an important determinant for keeping an overall internal balance of forces and deformations (inertial effects can largely be disregarded), avoid excessive deformations and monitor cardiac performance into early diastole (antagonistic forces were found to extend into early diastole thereby supporting rapid early diastolic filling). Animal experiments involving inotropic drugs and clinical observations (below) in connection with fibrosis confirm the antagonistic concept [14].

Second, an apparently “optimal”, regular arrangement of cardiomyocytes has been analysed in mathematical heart models; such arrangements, however, are found to be highly sensitive and unphysiologic: even a slight deviation from the “optimal” architecture results in a dramatic decrease of cardiac performance. Stochasticity, in turn, enables a stable cardial performance over a wide variety of anatomical aberrations [15,16].

Asymmetry of the action of inotrops

It is a specific feature of the oblique transmural netting component to react more sensitive to inotropic interventions than the tangential netting component. The answer of the oblique structure to positive inotropics is at least double that of the tangential alignment. The answer to negative inotropics is 5 times more pronounced (Figure 4). In any case this asymmetry in answer only apply to low dosage of medication. At high doses the answer to positive as well to negative inotrops more or less is comparable in both netting components [13,17,18].

Clinical consequences

The derailment of the equilibrium provided by the antagonistic organisation of myocardial function might explain various cardiological diseases. Thereby, the clinical understanding of ventricular afterload has to be extended. Besides the extrinsic hemodynamic afterload, the heart contracts against an intrinsic myostructural, stromatogenic and hydraulic resistance. Within the framework of this complex afterload environment that the heart has to cope with, it is prone to derail each in a characteristic way, causing each a particular pattern of clinical pathology and requiring each its particular therapy [14].

Hemodynamic Afterload

The hemodynamic afterload is given by the systolic ventricular pressure, typically increases as a consequence of arterial hypertension,
aortic stenosis or contractile or fibrotic constriction of the ventricular outflow tract. Obviously, those disturbances can well be diagnosed and treated under control of intracavitary pressure.

Myostructural Afterload

The myostructural afterload which results from myocardial transmural netting, derails in all kinds of myocardial hypertrophy. The particular feature of hypertrophy is that the intrinsic antagonism sustains the process of hypertrophy such the disease is self-amplifying. This vicious circle can be interrupted by tempering separately the contractile forces in the oblique transmural deviation of myocardial netting, which can be achieved by use of low dose negative inotropics. The clinician’s problem is that there are no means to evaluate the degree of derailment unless by measuring wall stiffness, which recently promises to become measurable in terms of “Cardiac Shear Wave Velocity Detection” [19-21].

Another manifestation of derailment of myostructural derailment most probably is Takotsubo-syndrome [22-24], which goes along with an active systolic dilation of the ventricular apex. Concommitant increase in proximal wall tension, which again escapes clinical evaluation, restricts perfusion of the intramural microvasculature and ends up in most painful myocardial ischemia. Running investigations are focused on the question, whether the dominance of dilating forces is caused by an atrophy of tangential netting components in physically undertrained elderly patients, while the oblique transmural netting component persists because it is kept fighting against systolic wall thickening. An alternative option is an excessive increase in sensitivity of transmural netting components for stress-induced positive inotropic stimulation. In any case the therapeutic way to go is low dose β-Blockade [25].

Stromatogenic afterload

The stromatogenic afterload derives from the effect of connective tissue netting which in the normal heart most probably has no measurable effect. It increases however dramatically in case of transmural or diffuse myocardial fibrosis (Figure 5) which hinders systolic wall thickening. At the time being there is no effective therapy described.

Hydraulic afterload

The hydraulic afterload is caused by intramural fluid pressure, which increases in case of myocardial edema in myocarditis and in myocardial storage diseases. Intramural pressure acts on the smaller endocardial surface plane as well as on the larger epicardial surface plane which means that in any case intramural pressure has a prevailing dilatory effect (Figure 6). With any kind of accumulation of intramural fluids, wall thickness increases, which event expands the surface difference between epicardium and endocardium. Furthermore, intramural pressure increases. All effects contribute to an increase in hydraulic afterload hence dilating forces increase. Therapy must focus on reduction in wall thickness and intramural pressure by eliminating fluid and stored semifluid masses.

Conclusions

The ventricular architecture is characterized by a lamellar structure that is ubiquitously crosslinked. While during systole constrictive...
(essentially surface parallel) forces causing the ejection of blood dominate, there exists a notable antagonistic, dilating activity (opposing mural thickening) caused by lamellar segments that are oriented in a transmural direction. This antagonistically organized setting, along with some stochasticity with respect to lamellar segment orientation, is well adapted to keep local balance in that excessive deformation patterns are prevented; furthermore, a ventricle working properly over a wide range of physiological conditions is supported.

Deraliment of such antagonistic equilibration is prone to cause diseases that manifest themselves in various forms. In particular, in myocardial hypertrophy, disturbances in fibre-netting like obstructive hypertrophic cardiomyopathy, fibrosis, edema, storage diseases and, as described only recently, in Takotsubo-syndrome, which causes critical increase in antagonistic wall stress. This new pattern of pathophysiology requires new diagnostic means by which to measure wall stiffness as an indicator of wall tension.

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