Histological architecture of cardiac myofibers composing the left ventricle of murine heart

Zienab A. Gouda¹, Yaser Hosny Ali Elewa²* and Assmaa O. Selim¹

Abstract

Background: Despite the fact that the exact architecture and orientation of left ventricular myocardial fibers are critical to cardiac functions either in health or disease, it is still debated.

Aim of the work: Histological demonstration of the transverse alignment of the myofibers making the left ventricular mass was applied to validate different anatomical speculations of cardiac myofibers organization; mainly Streeter’s conjecture and the ventricular myocardial band model of Torrent-Guasp.

Material and methods: Six healthy adult male C57BL/6N mice were utilized. Their hearts were separated into atria and ventricles. Then, both ventricles were divided into four levels [base, upper mid, lower mid and apex]. Paraffin step-serial transverse sections were stained with H&E and Masson’s trichrome stains. Morphometrical measurements of the thickness of the anterior, posterior, and lateral walls of the left ventricle (LV) as well as the inter-ventricular septal wall were realized with ImageJ software. For data analyses among the four levels of LV, Scheffé’s method was applied for multiple comparisons when a significant difference was observed by Kruskal-Wallis test (p<0.05).

Results: Examination of the LV from its base to its apex revealed obvious changes in the thickness of its walls and inter-ventricular septum as well as in its cavity shape. Interestingly, the myocardial fibers showed different running patterns [longitudinal, oblique and circular] among different levels as well as within the same level from sub-epicardial to the endocardial region. Throughout all levels, the sub-epicardial myofibers showed longitudinal orientation while that of the intermediate wall revealed either oblique [in upper mid-level] or circular orientations in other levels. The sub-endocardial fibers were mostly longitudinal with the presence of some circular fibers in the base and upper-mid regions. Masson’s trichrome sections revealed a trivial amount of collagen fibers just around each individual myocyte without any bundle formation.

Conclusion: The presence of different running patterns of the myocardial fibers among different levels of LV as well as within the same level indicates multiple rolling of the myocardial fibers. Thus, we suggested that the band model by Torrent-Guasp accounts for the patterns of myocardial fiber architecture forming the left ventricle.

Keywords: Left ventricle, myocardial band, Torrent-Guasp, C57BL/6N mice

Introduction

The structure and function of organs are inseparable both in health and disease states. Many questions in cardiovascular medicine are still pending due to the insufficient insight in the basic science [1]. In particular, the elucidation of left ventricular myocardial architecture is critical to understand the exact mechanisms of cardiac functions. The cardiac myofibers orientation plays an important role not only in mechanical contraction but also in the electrical conduction and energy metabolism [2]. Although there are numerous anatomical works
Many anatomical speculations had been documented about the exact ventricular myocardial arrangement. The organization of cardiac myofibers as following: “Myocardial fibers run like geodesics on a nested set of toroidal bodies of revolution” was described [4]. Further investigation showed that the musculature of the heart is arranged on the basis of a modified blood vessel rather than in the fashion of a skeletal muscle which has discrete origin and insertion [5]. The heart was described as a muscular band that begins in the insertion of the pulmonary artery and ends at the level of the aorta, forming a double helix, which limits both ventricles [6]. It is noteworthy, however, that although Torrent-Guasp [6] also de-emphasized the concept of the fibrous skeleton as a potential anchorage of the fibers, he offered no explanation as to the nature of the anatomical structures which permitted him to dissect out the myocardial band with the consistency he claimed to demonstrate [7].

Recently, by blunt dissection of the epicardium, or epimysium, that surrounds the ventricular mass, the cardiac myofibers forming the left ventricle (LV) had been reported to be organized in branching layers separated by cleavages, referred to as the “sheet architecture” [8,9].

By using diffusion tensor magnetic resonance imaging on the ventricular epicardium, the fibers are shown to extend obliquely. While, in the middle layer of the anterior, the posterior and lateral walls of the left ventricle, the fibers are nearly circumferential. In the endocardium, the fiber orientation is more oblique on the anterior wall than the posterior wall. Finally, from the apex to the base of the heart, the epicardial fibers are arranged clockwise, and the endocardial fibers are counter-clockwise [2].

There is no anatomic evidence supporting the concept of the “ventricular myocardial band” [7]. Moreover, previous histological studies that described the ventricular myocardium were very shallow as most studies use cross-sectional or longitudinal areas of one level only. Transversally, the myocardium wall is made up of millions of myocytes set in a matrix of connective tissue. Each individual cardiac myocyte appears as a long thin cell, possessing multiple side branches, joined to its neighbors by intercalated discs [10]. Longitudinally, the myofibers in subendocardial and subepicardial parts of the LV wall run parallel to the ventricular equator [5].

The previous histological data concerning the architecture and orientation of cardiac myofibers in the LV are scarce. Furthermore, the previously mentioned anatomical speculations of the cardiac myofibers organization remain a highly contentious topic and have not yet been histologically confirmed.

Thus the aim of the present study was to validate different anatomical speculations of cardiac myofibers organization by histological examination of step-serial transverse sections over all the different levels of murine LV. We hypothesized that the observation of the running patterns of the cardiac myofibers in mouse LV at different levels could clarify the exact cardiac myofibers organization. In this study, we expected the presence of different running patterns of the myocardial fibers among different levels of LV which could be an indication of the existence of multiple rolling and helical arrangements of cardiac myofibers.

Material and methods
Animal and tissue preparation
Six healthy adult male mice, eight to ten weeks old were purchased from Japan SLC (Shizuoka, Hamamatsu, Japan). These animals were maintained in a controlled environment [room temperature of 22±4°C, a relative humidity of 55±20%, and a 12-h light and dark cycle] in the animal facility of the Graduate School of Veterinary Medicine, Hokkaido University. The mice were allowed free access to tap water and an adequate diet and were used at 4 months of age. In handling the experimental animals, the investigators adhered to the Guide for the Care and Use of Laboratory Animals, Hokkaido University, Graduate School of Veterinary Medicine [approved by the Association for Assessment and Accreditation of Laboratory Animal Care International].

After the mice had been killed by deep inhalation anesthesia, hearts were removed. The atria were separated from ventricles, and the total length of both ventricles was measured [average, 10 mm]. Then, as there are variations in the septal wall thickness and the size of left ventricular cavity, we divided both ventricles transversally into four levels of equal thickness [average, 2.5 mm each]. The first level started just beneath the atria and characterized by narrow cavity and we considered it as “base level”. The second and third parts were “upper and lower mid levels” characterized by wide cavity and thick septal wall and it represented the middle levels of the ventricles. The fourth cone-shaped level was characterized by narrow slit shaped cavity and thicker septal wall, and it represented “apex level”.

Then, the cardiac tissues at four levels were fixed with 4% paraformaldehyde. For histological examination, the specimens were dehydrated in graded alcohol and embedded in paraffin. Subsequently, step serial sections [3-μm-thickness] from different levels of the ventricles with 15 μm interval after every 10 serial sections were cut. These sections were deparaffinized, rehydrated rehydrated, stained with Hematoxylin and Eosin (H&E) or Masson’s trichrome (MT), and observed under light microscope [11].

Morphometric study
By using Image J software [National Institute of Health; NIH, Bethesda, MD, USA], the thickness of the anterior, posterior, and lateral walls of the LV as well as the inter-ventricular septal wall was measured, analyzed, and compared among the four levels of heart [base, upper and lower mid and apex]. To measure the thicknesses of LV walls and inter-ventricular...
The lateral ventricular wall was the thickest at the upper mid level. The mouse heart was divided into four main levels: base, intermediate, posterior, sub-endocardial regions at the base level (Figures 1A). Step-serial HE stained cross sections of the LV revealed obvious changes in the thickness of anterior, lateral, posterior ventricular walls and inter-ventricular septum, as well as, in the shape of cavity among different levels of the heart. The results of histoplanimetrical analysis were summarized in Table 1.

Briefly, the anterior wall was the thickest at the base level [383.68±0.29 μm] (Figure 1B). Then, it became gradually thinner towards the apex level [89.45±0.78 μm] (Figure 1E). The lateral ventricular wall was the thickest at the upper mid level [189.03±0.58 μm] (Figure 1C), and it was the thinnest at the apex [93.32±1.35 μm] (Figure 1E). At the base level, the lowest thickness of posterior wall [64.68±0.23 μm] and inter-ventricular septal wall [135.89±0.55 μm] was observed (Figure 1B). On the other hand, the highest thickness of the posterior wall [299.34±0.31 μm] and the inter-ventricular septal wall [204.93±1.12 μm] was observed at the apex level (Figure 1E).

At the base level just beneath the left atrium, the cavity of LV appeared as a small triangle shaped opening that represents the aortic orifice (Figure 1B). A gradual widening of its cavity ranging from nearly triangular to rectangular shapes at the upper and lower mid levels (Figures 1C and 1D), and it became a small slit at the apex level (Figure 1E).

The cross section of the LV at base level revealed the branch of coronary artery, paracanal interventricular branch, in the lateral surface of LV wall (Figure 1B). Its position changed from lateral to anterior position at the mid levels (Figures 1C and 1D), meaning oblique running patterns of this artery towards the apex of heart.

Histological characteristics of cardiac myofibers in mouse LV at the base level

Next, we compared the running pattern of cardiac myofibers in the lateral wall of LV at each level. The results were summarized in Table 2.

At the base level, the LV was composed of a thick wall (Figures 1B and 2A). The running directions of cardiac myofibers forming the lateral LV wall differed among sub-epicardial, intermediate, and sub-endocardial regions at the base level (Figures 2B and 2D). At both sub-epicardial and intermediate regions, most of cardiac myofibers showed a longitudinal running pattern, meaning they run along the axis between base and apex of heart (Figures 2B and 2C). However, at sub-
Table 1. Statistical analysis of the thickness of the anterior, lateral, posterior ventricular wall and septal wall among the different levels of the left ventricle.

<table>
<thead>
<tr>
<th>Level</th>
<th>Anterior wall</th>
<th>Lateral ventricular wall</th>
<th>Posterior wall</th>
<th>Septal wall</th>
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<tbody>
<tr>
<td>Base level</td>
<td>383.68±0.29abc*</td>
<td>112.36±0.55abc&lt;sup&gt;*&lt;/sup&gt;</td>
<td>64.68±0.23abc&lt;sup&gt;*&lt;/sup&gt;</td>
<td>135.89±0.55bd&lt;sup&gt;*&lt;/sup&gt;</td>
</tr>
<tr>
<td>Upper mid level</td>
<td>269.98±0.41b&lt;sup&gt;*&lt;/sup&gt;</td>
<td>189.03±0.58bd&lt;sup&gt;*&lt;/sup&gt;</td>
<td>90.63±0.22bd&lt;sup&gt;*&lt;/sup&gt;</td>
<td>193.69±1.03bd&lt;sup&gt;*&lt;/sup&gt;</td>
</tr>
<tr>
<td>Lower mid level</td>
<td>130.36±0.36c&lt;sup&gt;*&lt;/sup&gt;</td>
<td>149.62±0.62cd&lt;sup&gt;*&lt;/sup&gt;</td>
<td>191.31±0.24cd&lt;sup&gt;*&lt;/sup&gt;</td>
<td>156.14±1.50cd&lt;sup&gt;*&lt;/sup&gt;</td>
</tr>
<tr>
<td>Apex level</td>
<td>89.45±0.78bcd&lt;sup&gt;*&lt;/sup&gt;</td>
<td>93.32±1.35bcd&lt;sup&gt;*&lt;/sup&gt;</td>
<td>299.34±0.31bcd&lt;sup&gt;*&lt;/sup&gt;</td>
<td>204.93±1.12bcd&lt;sup&gt;*&lt;/sup&gt;</td>
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*Significant levels difference (Kruskal-Wallis test, P<0.05). a,b,c,d significant difference with apex, lower mid level, upper mid level and base, respectively (Scheffé’s method, P<0.05).

Table 2. The running patterns of cardiac myofiber in lateral wall of mouse LV at different levels of the heart.

<table>
<thead>
<tr>
<th>Levels regions</th>
<th>Sub-epicardial region</th>
<th>Intermediate region</th>
<th>Sub-endocardial region</th>
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<tr>
<td>Base level</td>
<td>Longitudinal</td>
<td>Longitudinal</td>
<td>Circular</td>
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<tr>
<td>Upper mid level</td>
<td>Longitudinal to oblique</td>
<td>Oblique to circular</td>
<td>Longitudinal, circular</td>
</tr>
<tr>
<td>Lower mid level</td>
<td>Longitudinal and circular</td>
<td>Circular</td>
<td>Longitudinal</td>
</tr>
<tr>
<td>Apex level</td>
<td>Longitudinal to oblique</td>
<td>Longitudinal</td>
<td>Longitudinal</td>
</tr>
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endocardial region, cardiac myofibers showed mainly a circular running pattern along the cavity of LV.

**Histological characteristics of cardiac myofibers in mouse LV at the upper mid level**

At the upper mid level (Figure 3A), the cardiac myofibers were longitudinally arranged along the axis between base and apex of the heart at sub-epicardial region in lateral wall of LV (Figure 3B). They changed to oblique running patterns at the deep portion of sub-epicardial regions, indicating a spiral running pattern of them along the axis between base and apex of the heart. And then, cardiac myofibers changed from an oblique to circular running patterns towards deep intermediate regions.
At the sub-endocardial region, cardiac myofibers showed both longitudinal and circular running patterns (Figure 3D).

**Histological characteristics of cardiac myofibers in mouse LV at the lower mid level**

At the lower mid level (Figure 4A), the cardiac myofibers showed both longitudinal and circular running patterns along the axis between base and apex of the heart at the sub-epicardial region (Figure 4B). At the intermediate region, cardiac myofibers predominantly showed circular running patterns (Figure 4C). At the sub-endocardial region, cardiac myofibers were longitudinally arranged (Figure 4D).

**Histological characteristics of cardiac myofibers in mouse LV at apex level**

At the apex level (Figure 5A), the cardiac myofibers were arranged longitudinally beneath the epicardium and obliquely at intermediate region along the axis between base and apex of the heart (Figure 5B). Then most of cardiac myofibers were circularly arranged at the intermediate myocardial region (Figures 5B and 5C). At the sub-endocardial region, cardiac myofibers were longitudinally arranged (Figure 5D).

**MT stained sections**

**Connective tissues in the mouse LV**

We examined the amount of connective tissues by MT stained sections at each level and revealed that there was a very little amount of collagen fibers just around each individual cardiac myocyte (Figures 6A-C). They were observed around the long axis of cardiac myocytes (Figures 6B and 6C), and there was no difference in the amount of connective tissues among different levels [data not shown].

**Discussion**

The cardiac myocytes architecture in LV is complex. Their arrangement is as so critical as evidenced by a significant functional impairment seen in post-infarction remodeling and other cardiac diseases. Adequate understanding of the myocardial structure and function is indispensable for the management of patients with congenital and acquired heart diseases [12]. In mice, the cardiac myofibers are longer [~150 µm] than the thickness of the standard histological sections [13], thus in the present study, we applied step-serial cross sections of the LV but not longitudinal sections. Furthermore, the mouse was chosen because it is the most popular animal model for studying normal and abnormal cardiac development as it has a strong heart [600 beats/min] [14].

In the present study, we revealed changes in the thickness of LV wall and the shape of its cavity among studied levels. It had been reported that the thickest part of cardiac muscular strand is at the basal level of human LV [15]; however, this study showed that the thickest part of the lateral LV wall was observed at the upper mid level. Towards the apical
Figure 6. Photomicrographs of the connective tissue in the base level of mouse LV according to MT stained sections (A-C). Notice few collagen fibers (arrows) surrounding individual cardiac myocyte in the sub-epicardial region [panel A, MT], intermediate region [panel B, MT], and in the sub-endocardial region [panel C, MT]. Bars: 100 μm.

cone, the lateral LV wall became thinner, while the inter-ventricular septum became thicker than that in other levels. Similar features were recorded in bovine heart [16,17]. These differences might reflect the functional or developmental species-related differences of heart structure. Furthermore, the changes of the thickness of LV wall observed in this study were in accordance with the previous report by [18] who referred these changes to the presence of papillary muscles and trabeculae. These structural changes were due to the continuous cardiac myocytes loops forming the LV walls [13].

The present work revealed obvious complexity of cardiac myofibers architecture in-between different levels, as well as, within the same level of LV. These identified features in the present study were also observed in human heart [13]. Characteristically, the cardiac myofibers just beneath of epicardium showed mainly a longitudinal running pattern along with the axis between base and apex of the LV, and they tended to be changed into a longitudinal or circular running pattern towards endocardium at all examined LV levels. Using diffusion tensor magnetic resonance imaging, at the ventricular epicardium, the fibers were extending obliquely [2].

The different orientations of cardiac myofibers throughout the LV wall at the same level might indicate the helical looping of cardiac myofibers forming the LV as mainly suggested by Torrent-Guasp [19,1]. In addition, our considerations might be also supported the previous report of other researchers [20]. They observed highly discontinuous laminae of myofibers that begin and end many times in-between the endocardium and the epicardium.

Furthermore, the running pattern of cardiac muscle is closely related to the blood vessels of the heart [21,22]. In fact, the left paraconal interventricular branch in the present study run obliquely. Moreover, some studies reported that there is a correlation between the spiral running pattern of cardiac myofibers and the ventricular function [augmentation the momenta of the blood flow] [23,24].

The cardiac vasculature might not necessarily correspond to the direction of the cardiac myofibers. By using two photons microtomy, the cardiac blood vessels were shown to move up, while the cardiac myofibers generally were moved down [13]. The spiral orientation of sub-epicardial myofibers to form the double helical muscular band in LV had been recorded anatomically [25]. Many congenital heart defects such as ventriculo-arterial discordance (complete transposition of the great arteries), encouraged the interruption of the spiral pattern of the heart [26]. In the upper mid level sections, the diverse running patterns of cardiac myofibers noted in the sub-epicardial, intermediate, and endocardial region, were in agreement with [27]. They showed that, this peculiar arrangement of cardiomyocytes and twisting appearance are considered the key factor of normal systolic and diastolic myocardial function.

In this study, the little amount of connective tissue observed
around each cardiomyocyte was in accordance with [7]. They mentioned that each individual cardiac myocyte is wrapped in an endomysium, which supports the intercalated discs in binding adjacent myocyte to one another, as well as forming a so-called ‘weave’ around the myocytes. However, another study attributed the separated and distinguishable myofibers to the difference in the fiber direction within their boundaries as the fibers are not outlined by a major connective tissue sheet [7]. They depicted this complex nested continuum of LV wall to form mechanical and electrical syncytium. Moreover, contraction of the ventricular myocardial band in this way creates a muscular force for suction to fill the ventricle during diastole [6]. Other researchers also suggested that the oblique oriented fibers of the septal wall are necessary for functional performance of LV [16]. Our results might emphasize the close correlation between histology and function of mouse LV.

**Conclusion**

In this study, we revealed obvious changes in LV wall thickness, inter-ventricular septum thickness, and also in its cavity shape from its base to apex in mouse heart. Furthermore, we revealed the running pattern of cardiac myofibers in LV wall at different levels from beneath atrium to apex of the heart with several histological characteristics. The different running patterns of the cardiac myofibers among different levels as well as, within the same level from sub-epicardial to the endocardial region might indicate multiple rolling of the cardiac myofibers. Thus, we suggested that the band model of Torrent-Guasp (Figure 7) accounts for the patterns observed in mouse LV. With the new cardiac imaging techniques, the general arrangement of the myofibers that make up the left ventricular walls is reviewed to provide a morphologic basis for diagnosis of several cardiac pathologies such as cardiac deformation, failure and myopathy. In addition, disarray of LV myocardial fibers could be a predictor of chronic kidney disease. However, further investigations are needed to concern both the orientation and coupling of cardiac fibers throughout all levels of the LV to strengthen the present data in the literature.

**Competing interests**

The authors declare that they have no competing interests.

**Authors’ contributions**

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**References**


