Regional localisation of left ventricular sheet structure: integration with current models of cardiac fibre, sheet and band structure

Stephen H. Gilbert*, Alan P. Benson, Pan Li, Arun V. Holden

Computational Biology Laboratory, Institute of Membrane and Systems Biology & Cardiovascular Research Institute, Worsley Building, Faculty of Biological Sciences, University of Leeds, Leeds LS2 9JT, UK

Received 22 January 2007; received in revised form 12 March 2007; accepted 13 March 2007; Available online 25 April 2007

Summary

The architecture of the heart remains controversial despite extensive effort and recent advances in imaging techniques. Several opposing and non-mutually compatible models have been proposed to explain cardiac structure, and these models, although limited, have advanced the study and understanding of heart structure, function and development. We describe key areas of similarity and difference, highlight areas of contention and point to the important limitations of these models. Recent research in animal models on the nature, geometry and interaction of cardiac sheet structure allows unification of some seemingly conflicting features of the structural models. Intriguingly, evidence points to significant inter-individual structural variability (within constrained limits) in the canine, leading to the concept of a continuum (or distribution) of cardiac structures. This variability in heart structure partly explains the ongoing debate on myocardial architecture. These developments are used to construct an integrated description of cardiac structure unifying features of fibre, sheet and band architecture that provides a basis for (i) explaining cardiac electromechanics, (ii) computational simulations of cardiac physiology and (iii) designing interventions.

Keywords: Ventricle; Sheet structure; Laminar myocardium; Continuum; Syncytial mesh; Helical ventricular myocardial band

1. Introduction

A heated debate is ongoing in the Journal and the wider literature on the morphology of the ventricular myocardium. The current controversy stems from longstanding disagreements—a complete and consistent anatomical description of cardiac structure has proved to be difficult. The heart is a contradiction between the symmetry of helically coursing fibres and regularly stacked laminae with the asymmetry of anisotropic fibre branching and merging of laminae. This complexity has led to the proposition of several incompatible structural models.

One current controversy concerns Torrent-Guasp’s concept of the helical ventricular myocardial band (HVMB)—a structural and functional hypothesis that the ventricles are a single band of muscle, coiled into two helices with insertions on the pulmonary trunk and aorta. This model has gained some favour in the medical community, particularly among cardiac surgeons. Indeed, the HVMB has influenced surgical procedures, surgical research and the understanding of the dynamics of heart contraction. However, the HVMB concept has received vigorous criticism of the objectivity and repeatability of the dissection method, its philosophical basis and the failure by its proponents to find or adequately look for independent corroborating evidence. At least eight reviews, one original paper and five letters supporting the HVMB, and six reviews, two original papers and seven letters opposing the HVMB have been published since January 2003. Until recently the argument has been polarised, but Criscione et al. [1] have widened the discussion to a more complex and subtle view of cardiac structure: that there may be regional specialisations of the myolaminar architecture within a continuum meshwork, and that the HVMB may or may not represent a simplification of these regions.

A second area of controversy is on the nature, orientation and degree of anisotropy of the laminar structure of the heart. A non-partisan consideration of both these controversies is presented here. New imaging evidence integrates some features from the opposing models. Based on published evidence, along with our revisualisation of data from diffusion tensor magnetic resonance imaging (DT-MRI), a unifying concept of cardiac structure is proposed which is based on the groundbreaking work of Grant [2].

Excellent historical reviews of heart structural research have been compiled [3,4] and are not recapitulated here.
2. Fundamentals of cardiac structure

Along with controversy on macroscopic heart structure, there is some disagreement and confusion on microstructural histology and the terminology used to describe this. This is chiefly a result of attempts to fit a simple classification scheme derived from skeletal muscle morphology to the more complicated structure of the heart. We first review and define the terminology.

2.1. Myocytes

Ventricular muscle cells (myocytes) are long, thin cells; their precise dimensions are variable depending on species, age and cardiac location, but are within the approximate range of 50—150 μm in length and 10—20 μm in diameter [3,5—8]. They are joined to their neighbours via intercalated discs at their ends. Terminal abutments are usually with more than one cell [3,9]. Being long extended cells, myocytes possess a principal direction given by the long axis of the cell, and the ellipsoidal myocardial nuclei are aligned to this axis [10—13].

2.2. Myofibres and myolaminae

In skeletal muscle morphology, myofibre has an absolute definition. In cardiac morphology it has different meanings to different researchers. Streeter [3] uses the term as a synonym for the myocytes. Other groups [4] use the term to indicate groups of three or more myocytes surrounded by a perimysial weave. The term myolaminae has been introduced [9] to describe sheet-like cleavage planes of attached myocytes (Fig. 1d—g). Some controversy exists regarding the form of laminae within the living heart. Some researchers [4,8,14], although recognising the existence of laminae, have played down their significance, emphasising instead the mesh-like nature of cardiac structure. The evidence presented in this review points to myolaminae and the groups of myocytes surrounded by a perimysial weave being a continuum of structure; therefore, we use myofibre to refer to individual myocytes and myolaminae to refer to these groups of myocytes. The term fibre is used by some groups to denote a continuous axial sequence of individual myocytes [15]. We use this term to denote the net axial direction of myocytes at a specific cardiac location, always with consideration of the important caveats in defining such a continuous axial sequence (as discussed in Sections 4 and 6). Likewise, the term sheet is related to the myolaminae but has a looser definition — describing the planar features of the myocardial wall, both the myolaminae and the cleavage planes between myolaminae [16].

2.3. Interlinking and merging of cardiac microstructure

Simplified connective tissue structure diagrams are often presented in reports of cardiac microstructure, but they can be misleading as they suggest a rigidity of structure. The key feature of cardiac microstructure is the interlinking and merging of structural elements; fibres link with several others at their poles, myolaminae coalesce and divide (visible in careful inspection of Fig. 1e) and fibres run from one myolaminae to another, all surrounded and supported by

---

Fig. 1. Microscopic cellular and intercellular (a—f) and macroscopic (g) tissue structure of the LV wall. (a) Lower magnification and (b) higher magnification optical images of a rat LV midwall stained with picrosirius red. Endomysial collagen surrounding individual myocytes is evident. M, myocytes; C, collagen cords; V, vascular structures; P, cleavage planes. (c) Honeycomb connective tissue skeleton revealed by scanning electron microscopy of human heart. The perimysium (P) envelops groups of myocytes. The endomysium, as final arborisation of the perimysium, supports and connects individual cells. The endomysial weave (W) envelops each individual myocyte and is connected to adjacent myocytes by lateral struts (S) presenting branches of variable size and extension. The range of length and diameter of these struts is very wide. Collagen struts also connect myocytes to interstitial microvessels (arrow 1) or perimysium (arrow 2). (d—f) Scanning electron micrographs showing three orthogonal surfaces of a typical left ventricular midanterior midwall specimen: (d) oblique view of tangential (TN), axial transmural (AT) and transverse (TR) faces of specimen; (e) view of TN and TR surfaces; (f) TR surface. Perimysial connective tissue weave surrounding myocardial sheets is evident and covers surface capillaries (C). (g) Micrograph of longitudinal—transmural section of left ventricle showing the radial alignment of myolaminae [a and b] from Young et al. [20] with kind permission from Blackwell Publishing; (c) from Rossi et al. [93] used with permission; (d—g) from LeGrice et al. [56] used with permission.
a complex collagen and elastin network. As with fibre nomenclature, connective tissue terminology is derived from skeletal muscle morphology. In its application, attention must be paid to the multi-level complexity of ventricular microstructure.

2.4. Endomysium

Each myocyte is wrapped in a prominent sheath of endomysium which supports the Z-bands and basal laminae [17]. The endomysium is seen in picrosirius red-stained confocal images (Fig. 1b), high-power scanning electron micrographs (not shown) and sodium hydroxide digestions (Fig. 1c).

2.5. Perimysium

The perimysium unites the myolaminae as units, from within, via collagen struts linking adjacent myocytes (120–150 nm) (Fig. 1b), and externally as a weave of connective tissue (Fig. 1c and f). Long perimysial collagenous tendons link connective tissue of adjacent myolaminae (Fig. 1a, b, e and f).

2.6. Epimysium

This term is of little value in cardiac morphology. In skeletal muscle it represents the outer smooth collagen layer of the muscle belly which makes no fibrous connections from its external surface — the cardiac equivalent is only present at endocardial and epicardial surfaces.

3. Models of heart structure

Seven conceptual models of cardiac structure are summarised in Fig. 2. The variety of proposed structures is visually striking — this is due, in a large part, to the different levels of cardiac structure described (discussed in detail in Section 4). Model 6 (Fig. 2, part 6) groups the myocardium into regional functional units analogous to skeletal muscles. Models 1–5 (Fig. 2, parts 1–5) are continuum concepts — stressing to differing degrees the anisotropic interconnectivity inherent in cardiac structure. Models 1 and 2 (Fig. 2, parts 1 and 2) describe fibre orientation. Model 3 (Fig. 2, part 3) describes changes through layers from epicardium to endocardium. Models 4 and 5 (Fig. 2, parts 4 and 5) examine the myolaminar structure, with less emphasis on fibre orientation. Some descriptions of Model 7 (Fig. 2, part 7), the HVMB, have emphasised discrete structural bundles [18] while other reports have proposed the band within a continuum framework [8].

4. Philosophical approach for description of a complex structure

How can so many models co-exist? Some must be wrong — or is it possible that different models are true representations of the heart when considered from different perspectives? One 1965 review on cardiac structure stands out as a philosophical foundation upon which all future studies should be based. Robert P. Grant, M.D., in his Notes on the Muscular Architecture of the Left Ventricle [2], puts forward the following principles (along with an excellent description of cardiac structure):

1. as evidenced from numerous dissections, a given fibre segment has branching connections with other fibre segments in several different directions — as such the myocardial structure is a syncytium-like arrangement [19],
2. the cardiac structure problem is therefore a three-dimensional network problem,
3. structure understood from study of the network depends on the level on which the structure is approached; whether an attempt is made to describe the predominant behaviour in the entire ventricle(s) or at the other extreme, to describe the average branching from a specific location,
4. many models of heart structure can therefore be proposed depending on the conceptual approach,
5. statistical study of the branching must be added to geometry for an accurate picture of myocardial architecture,
6. in consideration of such statistical problems one approach is to construct models for different degrees of generalisation of the problem,
7. no single model gives the whole story but together they provide a schema,
8. the existence of a syncytium-like arrangement does not in itself dictate that no separate ‘bundles’ are present; however, due to the complex structure it is possible to construct by dissection bizarre arrangements which have no underlying anatomical reality,
9. any dissection of the myocardium may represent (a) a valid schema from an infinite set of valid schema, (b) a bizarre and meaningless pathway or (c) a grouping of fibre paths within the syncytium of such general shared fibre direction that it can be considered a physiological or anatomical bundle,
10. a unique anatomical bundle is not the same as a unique physiological bundle — a connection between separate anatomical structures may produce one physiological structure,
11. a statistical approach is required to demonstrate any independent anatomical entity.

When viewed from this perspective it is not surprising that many models exist, that these are not all mutually compatible and that argument continues. Further evidence for these principles is presented in a dissection study by Fox and Hutchins [19], where principle (1) is re-stated and emphasised: ‘The only level of the network of cells that can be referred to accurately as a fibre is a single cell. The “fibre” is often only one cell in length before it splits and branches’.

When considering the structural debate it is logical to return to Grant’s principles — an approach adopted in this review. It should be noted that although Grant suspected regional variations in fibre branching within the left ventricle, he was sceptical whether these local prevalences would statistically warrant consideration as separate
bundled. One feature of the myocardial structure problem Grant did not consider was that significant structural differences may exist between individuals of the same species — a question discussed at length later in this review.

5. Traditional and novel techniques

Traditional anatomical techniques have several limitations which have made it difficult to apply Grant’s principles: (i) dissections and histological studies are subjective; (ii) the techniques are better suited to qualitative rather than quantitative descriptions, so the application of statistical analysis is difficult; (iii) the techniques are destructive and (iv) the reconstruction of three-dimensional fibre and sheet orientation from two-dimensional sections has been difficult [3].

Recent technical developments have greatly contributed to the available data on cardiac structure. Automated confocal microscopy methods have allowed extended imaging of three-dimensional tissue at high resolution, clearly demonstrating laminar structure [20,21]. Optical methods, including polarisation microscopy, have allowed demonstration of fibre orientation in the whole heart [22,23].

Whole heart fibre and laminar structure have been revealed by DT-MRI, a three-dimensional technique which uses MRI to provide the axes of diffusion of protons of water molecules in tissue. Detail of tissue architecture is imaged by its restriction of the random movement of the protons. The pattern of diffusion is represented mathematically by a symmetric second-rank tensor in three-dimensional space, which can be written as a 3 × 3 matrix. The three orthogonal eigenvectors of this tensor (ranked in order of their magnitudes of their corresponding eigenvalues) have been related to cardiac structure, with the eigenvalues representing the diffusion along three principal axes. Theory suggests that the direction of greatest proton diffusion (i.e. the eigenvector with the largest eigenvalue, the primary eigenvector) will be along the fibre long axis, intermediate diffusion (the secondary eigenvector) will lie in the myolaminar plane, orthogonal to the fibre long axis. The third and minor direction of diffusion (the tertiary eigenvector) is by definition orthogonal to the primary and secondary eigenvectors, so is normal to the myolaminar plane. The correspondence between the primary eigenvector and fibre orientation, and the secondary and tertiary eigenvectors and sheet orientation has been validated by combined DT-MRI and three-dimensional histological reconstruction of fibre and sheet structure [9,13,24—27].

An advantage of whole heart DT-MRI and optical methods is that they can provide digital datasets that can be analysed and compared algorithmically at high speed. They are, therefore, inherently suitable for statistical analysis. The bias, resolution and selectivity for the architectural characteristic measured apply equally across the entire heart. Examination of data from these techniques is therefore highly suited to identification of grouped fibre paths, which satisfy Grant’s criteria for uniqueness; the validity of structural models is discussed in this light.

6. Doughnut and pretzel models — Models 1 and 2 (Fig. 2, parts 1 and 2)

The fibre path models are based on the synthesis of many measurements of angles of fibre orientation through the ventricular wall and differ primarily in their scope (Model 1 is left ventricle (LV) only; Model 2 is LV and right ventricle (RV)). No argument is made for discrete traceable fibre paths — rather a recognisable general fibre trajectory [3]. These models reconstruct the long accepted helical pattern of fibre orientation, which has recently been confirmed by DT-MRI [30,35—40] (Fig. 3). DT-MRI fibre-tracing algorithms (Zhukov and Barr [28] and Kondratieva et al. [29], applied to canine data; Schmid et al. [30], applied to porcine data; Rohmer et al. [31], applied to human data) and automatic computational constructive visualisation methods (Chen et al. [32,33] and Fig. 15(B) in [8]), used to extract meaningful visual information from histologically recorded three-dimensional fibre orientation datasets [33,34] (recorded from the rabbit heart), produce models remarkably similar to that proposed by Streeter (Fig. 2, part 1).

Fig. 2. Models of mammalian ventricular architecture. (1) LV as nested doughnut (toroidal) geodesics (synthesised by Streeter [3] from work of Krehl [49], Hort [54] and reconstructed by Jouk et al. [23] (see Model 2)): assumes myocardium is a specialised form of vascular musculature, with myocardial fibres as a network of preferentially orientated and branched myocardial cells forming end-to-end cellular junctions. The fibres follow paths of shortest length (geodesics) nested on doughnut-shaped surfaces (toroidal bodies of revolution). (2) LV and RV as nested pretzel geodesics (Jouk et al. [23]): extension of Model 1 to describe both left and right ventricles, based on data from quantitative optical polarisation microscopy of the foetal human heart. Fibres follow geodesic trajectories on nested warped pretzels, where a pretzel is two side-to-side joined doughnuts. (3) Three-layered ventricle (proposed by Rushmer et al. [48] and developed by Anderson et al. [4] and others): the ventricles are composed of superficial, middle and deep layers. The fibres follow a continuum or functional spongylous mesh (FSM) with multiple branching and cannot be split down further into functional units. There is a net helical fibre orientation across and within the ventricular walls [4]. Some debate has occurred between proponents of this view regarding the degree of fibre penetration between layers. (4) Simple laminar structure (proposed by LeGrice et al. [56]): the myocardial wall consists of an ordered laminar structure, separated by extensive cleavage planes that run approximately radially from endocardium to epicardium. In tangential sections the plane of the laminae coincides with local fibre orientation. Layers are about 4 ± 2 myocytes across and 48,4 ± 20.4 μm thick (in the dog). Components of the connective tissue matrix connect adjacent laminae. (5) Complex laminar structure: strain modelling, histological and DT-MRI studies [9,11,13,44,45,58] have revealed evidence of two approximately normal myolaminae populations. More complex features of cardiac structure have been described to account for this evidence, but have not been formalised or organised into complete conceptual structural models. However, one early model has been proposed by Harrington et al. [11] and is shown here. (6) Distinct muscle bundles (DMBs): Robb and Robb [61], following from Hall [94], consider cardiac structure analogous to skeletal muscle — fibres gathered in distinct bundles to form muscle bellies. Four muscle bodies proposed, anchoring to the fibrous trigones and aortic and pulmonary roots. (7) Helical ventricular myocardial band (HVMB) (Torrent-Guasp et al. [8,18,62,95]): both RV and LV exist as a single muscle band: the HVMB, organised spatially as a helix and consisting of basal and apical loops. This band is a helical rope-like structure that is unrolled easily from the heart by blunt dissection along natural cleavage planes — that follow the principal fibre direction at any point. A connective tissue sheath enclosing the band segments was not proposed [diagrams for Models 1, 2, 3 and 6, adapted from Jouk et al. [23] with kind permission from Springer Science and Business Media, Model 4 from LeGrice et al. [56] used with permission and Model 5 from Harrington et al. [11] used with permission. Model 7 from Kocica et al. [8] used with permission].
Peskin [41] has carried out an asymptotic analysis of Model 1 and has derived the fibre architecture of the heart from first principles. These models form a conceptual model of cardiac structure from the one macro- and microscopically observed characteristic of principal fibre direction. As such, the nested doughnuts/pretzel surfaces are abstract concepts — no discrete biological equivalents exist. It is possible to create the surfaces by dissection following the observed principal fibre direction, as demonstrated in early studies by Torrent-Guasp [3,42], but in so doing much information is lost. The dissection and histological methods used do not record data of local branching in directions other than the predominant fibre orientation, so by definition no detail of local tissue organisation is modelled. Locally branching fibres are smoothed to a single orientation, which is again smoothed to a global ventricular fibre orientation. With reference to Grant’s principles, these models represent the highest order schema of whole heart fibre orientation in isolation.

It might be assumed that the fibre maps produced by algorithmic analysis and computer visualisation may be more than a conceptual model, but the considerations above apply equally. The DT-MRI fibre-tracing algorithms track the principle fibre orientation from the primary eigenvector only. The automatic computational constructive methods were applied to histological fibre orientation datasets which only record principle fibre orientation. Being limited to tracking the principle fibre orientation alone, these methods cannot reconstruct any detail of myolaminar structure, and as such reproduce idealised fibre-tracing dissections.

Much evidence points to myolaminae as the central feature of the wall motion mechanism [16,43–46]. As such the doughnut and pretzel models represent geometric abstractions of cardiac structure. Their primary uses may be in (i) refining more histologically detailed models for the constraint of fibre orientation and (ii) in modelling the spread of the cardiac action potential, the conduction of which is significantly influenced by principal fibre direction [47].

7. Three-layered myocardium — functional syncytial mesh — Model 3 (Fig. 2, part 3)

This model is similar to the fibre path models (Models 1 and 2). It divides the myocardium into three layers on the basis of fibre orientation. It was put forward by Rushmer et al. [48] as a mechanistic model of cardiac contraction, in which (as first proposed by Krehl [49]) the middle circular layer acted as a constrictor. Further studies endorsed and extended this description [15,50–52]. Paradoxically, these later studies endorse this model whilst also accepting that the separation of layers is artificial as the orientation of fibres changes gradually, and there are fibres which span layers: ‘As a result we can differentiate three layers of fibres: superficial (subepicardial), middle and deep (sub-endocardial). However, it is very important to emphasize that the separation of these three layers is artificial since the change of the direction of the fibres is gradual without cleavage planes of connective tissue. Moreover, a number of fibres change layers in their trajectory around the ventricles’ [15].

As with the previous models, the features described in the three-layer model are largely correct — this is not surprising given the many anatomical dissections upon which the model is based. However, as described, the model is vague or contradictory in its description of the boundary zones between layers, where evidence points to a smooth transition of principal fibre direction through the myocardium. Laminar structure is omitted in early descriptions of the model and, although now accepted by some proponents [4], its higher-level symmetry and regional patterning remain unappreciated.
8. Simple myolaminae model — Model 4 (Fig. 2, part 4)

Sheet structure was first described in detail by Feneis (cited by Hort in [54]) and then Hort [53—55]. Hort described macroscopic sheet structure visible in dissections as "feathering" or "pinnation", and understood these areas to be regions of reduced fibre and connective tissue interconnectivity between sheets of myocardial cells. Grant [2] described parallel fibres, which "by their branching form planes that seem to rotate". Spotnitz et al. [46] not only described open spaces between planes in rat heart histology but also proposed that sliding between these planes accounted for the change in myocardial volume associated with contraction. Greenbaum et al. [50] referred to the grossly visible structures as "layers" and described them as regions of little change in fibre orientation in contrast to regions of rapid change, but disputed different connective tissue morphology between and within layers.

LeGrice et al. conducted the first quantitative analysis of sheet structure and formalised this into a conceptual [34,56] and mathematical [57] laminar model of cardiac structure in which the canine myocardial wall is organised with an ordered laminar structure, separated by extensive cleavage planes and running radially from endocardium to epicardium, integrating whole heart fibre orientation. Examination of the long-axis cut surface of the LV allows myolaminae to be identified without magnification (Fig. 1g). Viewing this image from a distance allows the identification of radial myocardial sheets, but the eye oversimplifies the structure — close-up inspection reveals a highly branched topology with two approximately perpendicular (strictly normal as in three dimensions) sheet populations.

Although the authors of this model recognised regional anisotropy in sheet structure, they underestimated its degree and significance which was recognised by Feneis (cited by Hort in [54]) and has recently been highlighted in several studies [9,11,13,44,45,58]. A mathematical analysis of the co-existing regular helical fibre path and LeGrice's proposed sheet structure (in the manner of Peskin [41] for fibre path alone) has not been carried out, but these features do not seem compatible.

9. The helical ventricular myocardial band — Model 7 (Fig. 2, part 7)

This is a structural and functional hypothesis that the ventricles are a single band of muscle, coiled into two helices with insertions on the pulmonary trunk and aorta. The concept is based on Torrent-Guasp's dissection method, where the predominant fibre direction at given point is followed in hearts boiled in 1% ethanoic acid. Aberrant fibres, going against the plane of dissection, are recognised but are torn in the process of the dissection.

9.1. HVMB topology

The concept of the ventricular myocardial band is very simple: there are two loops each divided into two segments (basal loop — right and left segments; apical loop — ascending and descending segments). The connectivity and anatomical arrangement of these four segments is shown in Fig. 4a.
9.2. Presentational problems

Early reports of the HVMB describe a 'unique morphologic and physiologic characteristic' of 'unique anatomy' that 'unwrap[s] along a natural cleavage plan' [18]. Although an enclosing connective tissue sheath is not proposed, the band could be considered a muscle belly in all except epimysium. Indeed some [59], after reading this description, have understood Torrent-Guasp to be describing connective tissue sheaths between band segments. A recent description [8] has highlighted the 'infinite anisotropy of the ventricular myocardium', recognised a 'functional syncytial mesh, resembling vascular musculature' and also that 'blunt anatomical dissections following predominant fibre direction, shall reveal unique functional (i.e. vectorial) but not any eclectic (i.e. discrete) anatomical planes in the ventricular mass'.

The continuum concept has always been present in the HVMB literature; in earlier descriptions it was underplayed [18,42,60] but has recently received greater emphasis [8]. While it cannot be expected that an idea will remain static over 30 years, the evolution of the concept is not clearly set out and this has increased confusion in the HVMB debate. This confusion is compounded by the presentation of images and diagrams of the HVMB dissection which create a false impression of the band's discreteness (Fig. 4b). Images of the same dissections in different orientations would emphasise the continuum of structure across band segments.

9.3. Philosophical problems

The HVMB has been dismissed on the basis of histological, optical [23] and DT-MRI [30] evidence of the smooth change in ventricular fibre orientation across the myocardium, as described in the geodesic models and shown in Fig. 3. If fibre angle changes smoothly across the wall with no glitches, how can the band exist? This argument against the band appears powerful, but is flawed. Grant's principles allow for the presence of discrete localisations of branching within a smoothly changing whole ventricle predominant fibre orientation schema. Recent evidence has been found for regionally anisotropic distribution of two sheet populations (discussed at length in Section 10.2). Could this regional sheet anisotropy represent the HVMB?

A more complete consideration of the HVMB requires separate analysis of (i) the merits of the dissection, (ii) the static structural model and (iii) the dynamic concept formulated on the basis of the structure.

A first consideration is that, by Grant's principles, any dissection path followed in the ventricles is a statistical concept. An implication is that if a path is sought from any point to any other within the ventricles it can by definition be found. In fact, infinite paths can be found, and again by definition, one path will also be most statistically favoured. Being the most favoured path alone does not necessarily make the path sensible or informative.

Is the HVMB (i) a bizarre and meaningless pathway, (ii) one of an infinite set of valid schema, (iii) a useful schema of structural features at one level of abstraction of cardiac structure or (iv) a grouping of fibre paths within the syncytium of such general shared fibre direction that it can be considered a unique physiological or anatomical entity (as proposed by Torrent-Guasp et al.)? The answer depends on both the statistical analysis of the fibre branching along the dissection path and on a detailed study of the macro- and microstructural boundary zone between HVMB segments. If the band is one dissection out of infinite possible dissections, each dissection will have a statistically derivable value calculated from the predominance of fibres following the chosen path at each point along its length. The validity of the band can be judged from the significance of the difference between the HVMB and other possible dissections (i.e. the p-value).

As discussed above, the HVMB has been rejected by some on the basis of histological [4,14], optical [23] and DT-MRI [30] evidence of the smooth change in ventricular fibre orientation across the myocardium (Fig. 3) There is, however, convincing and conflicting evidence that the HVMB does follow a favoured path from histological [54], optical [22] and DT-MRI data presented in Fig. 5. This figure synthesises fibre angle change, obtained from DT-MRI, with the dissection diagrams of Hort [54] and optical section reconstructions of McLean and Prothero [22]. Both longitudinal DT-MRI (fibre angle change) sections and the optical reconstruction split the ventricles into zones clearly reflecting the HVMB dissection planes.

DT-MRI-derived cross-sections show V-shaped features correlating to the pinnation features described by Hort [54]. If a dissection plane is made joining these features this reproduces a key feature of the HVMB dissection path. Streeter [3] described the linking of the V-shaped pinnation features in this manner to produce the discrete muscle bundles described by Robb and Robb [61] (Model 6). This evidence does not identify the HVMB as a unique unit but it does show that the dissection of Torrent-Guasp et al. follows, at least in part, a measurable difference within the myocardium. Visualisation of canine myocardial sheet angle (Fig. 8c and d; discussed in detail in Section 10.3) provides yet more convincing evidence that the HVMB follows real anatomical features.

This evidence of a favoured dissection path is not a justification of the HVMB structural model. Examination of high-resolution three-dimensional microscopic images show that (in rat heart at least) the band dissection plane tears across structural features of a complex continuum of merging sheet populations running from epicardium to endocardium (Fig. 6a--c; discussed in detail in Section 11). Laminar sliding and extension have been demonstrated as the most significant component of systolic ventricular thickening [44], and although strain modelling studies are only beginning to extend to consideration of two sheet populations [44,45,58], evidence points to interactions between sheet populations (in three dimensions) being key to LV radial wall thickening in systole.

The HVMB dynamic concept proposes independent band action via longitudinal sliding of HVMB band segments [62,63], and studies have revealed some division of dynamical properties across band regions [63,64]. This is not unexpected as different band areas have differing fibre and laminar structure. This evidence should not be interpreted as justification of the HVMB structural—functional concept, as it is incompatible with microstructural evidence.
At best, the HVMB dissection represents one generation of Grant’s schema of cardiac structure, but as described above, it misleads more than it informs. More useful schema are in the form of the DT-MRI transition data, which demonstrate boundaries of subtly different fibre orientation without overemphasising their importance by tearing and then refolding them as distinct units.

10. Unifying structural model

10.1. Terminology for local aggregations of structure

Primary cardiac structure has been defined as the predominant fibre orientation, secondary as the organisation of the myolaminae through which these fibres course, and tertiary structure as localised aggregations of primary and secondary structure (often referred to as bands) [4,14,65,66].

Some cardiac researchers have dismissed the presence of localised aggregations of cardiac structure in the myocardial wall [4], but there is convincing evidence from DT-MRI data. We present a revisualisation of DT-MRI data and review published DT-MRI results alongside optical, automated...
confocal, strain modelling and histological data. Before introducing this analysis a review of the recent literature on mixed sheet populations and sheet merging is presented.

10.2. The evolving concept of myolaminar sheets: mixed sheet populations, sheet merging, the role of sheets in myocardial contraction and inter-individual variability in sheet structure

There is general agreement based upon histology, optical studies and DT-MRI that myocardial fibre orientation is remarkably uniform between and within species [2,13,16,23—25,27,28,30,35—40,44,50,53,54,67—85] (Fig. 3). Intriguingly, there is recent convincing evidence that, despite this fibre structure uniformity, cardiac sheet structure is considerably variable within species [27,44,45].

In order to review the literature on cardiac sheet structure it is first necessary to clarify what is meant by the term 'sheet angle'. This is not a simple task; there are many distinct definitions of sheet angle in the literature. There is no uniformly accepted system for reporting cardiac geometrical data, but cylindrical [39] or prolate spheroidal [57] coordinate systems have been adopted. In either coordinate system, three standard cardiac axes are defined: the long axis, the radial axis and the tangential axis. These axes in turn are used to define three standard cardiac planes: the radial—longitudinal plane, the transverse plane (also known as the short-axis cross-section plane or the radial—circumferential plane) and the tangential plane [81,86]. Sheets have a separate local orthogonal coordinate system defined by the fibre-orientation axis (primary eigenvector), the sheet normal axis (tertiary eigenvector) and a second axis lying in the sheet plane (secondary eigenvector). In general terms, sheet angles relate the local sheet coordinate system to the standard cardiac coordinate system. It is possible to define many different angles between the sheet and cardiac coordinate systems, depending on the chosen plane or axis from each system, and unfortunately no convention has been established. All the sheet angle definitions used in the literature are geometrically dependent, but they are distinct. An additional complication is created by the differing angular scale adopted in reports, being either $0^\circ \leq \beta \leq 180^\circ$ or the equivalent range $-90^\circ \leq \beta \leq +90^\circ$. These considerations make both qualitative and quantitative comparisons of sheet structure between reports difficult.

The definitions of sheet angle adopted in visualisations in this report are those described in general geometric terms by Costa et al. [86], and are defined in relation to the DT-MRI eigenvectors here:

- $\beta^r$-Sheet angle ($-90^\circ \leq \beta^r \leq +90^\circ$): The angle the sheet makes in the longitudinal—radial plane. This is the angle between the transverse plane and the projection of the secondary eigenvector onto the longitudinal—radial plane. Positive angles rise to the heart base from endocardium to epicardium (or LV endocardium to RV endocardium).
- $\beta^s$-Sheet angle ($-90^\circ \leq \beta^s \leq +90^\circ$): The angle the sheet makes in the transverse plane. This is the angle between the longitudinal—radial plane and the projection of the secondary eigenvector onto the transverse plane. Positive angles rise to the heart base from endocardium to epicardium (or the projection of the secondary eigenvector onto a standard cardiac plane has not been performed). Positive angles rise to the heart base from endocardium to epicardium (or LV endocardium to RV endocardium). This angle is referred to a $\beta$ by Costa et al. [86], but here we denote it with $\beta^s$ as in Chen et al. [16].

The role, importance and the mechanism of action of cardiac sheets in contraction is addressed in many studies (as presented below) but is still an area of controversy. The cited papers should be referred to for the precise definition of the sheet angle reported. To allow qualitative comparison, angles are reported on the $-90^\circ \leq \beta \leq +90^\circ$ scale and positive sheet angle refers to sheets in which $\beta^s$ rises to the heart base from endocardium to epicardium.

Hort [53—55] proposed a model (Fig. 10 in [53] and Fig. 21 in [3]) to explain wall thinning from systole to diastole in which parallel bundles of fibres are arranged in several stacked layers in systole. In diastole, these layers of like-fibre orientation interdigitate, reducing the number of layers, and allowing the wall to become thinner and the cardiac chambers larger. This model could not function in the presence of cardiac sheets, as it requires bundles of myocytes separated by parallel cleavage planes that, due to their separate connective tissue, can interdigitate during the cardiac cycle.

Spotnitz et al. [46] explored the relative contribution of (i) the myofibre diameter and (ii) the sliding between bundles of myocytes, in myocardial wall thickening. They concluded that (i) changes in fibre thickness could not account alone for the changes in myocardial thickness during contraction, (ii) that 'cleavage planes' were present between groups of myocytes, that (iii) the sliding of groups of myofibres (permitted by cleavage planes between them) was an important mechanism in myocardial contraction, and is evidenced by cleavage planes having a more vertical alignment in diastole and a more horizontal alignment in systole, and that (iv) there is no evidence for insinuation or interleaving of fibres as suggested by Hort. They do not address if cleavage planes separate myolaminae or if they separate bundles of myocytes.

As described above, LeGrice et al. [56] carried out the first quantitative histological analysis of laminar structure throughout the ventricles, and formulated this into a mathematical model [57]. Unlike later reports, this model of the canine ventricles described smoothly varying sheets rather than dual sheet populations; however, regions of abrupt change can be recognised in the data (Fig. 8f). Two canine hearts were studied by histological measurement and two by scanning electron microscopy, but direct heart-to-heart comparisons were not performed. The study reported little detail of inter-individual variability in structure but found significant variation in local sheet structure within hearts, which was considered to be branching from a model of regular transmural endocardial to epicardial planes.

In a mechanics study, LeGrice et al. [43] demonstrated that in dogs >50% of the thickening of the myocardial wall during contraction could be accounted for by slippage along
the cleavage planes between myolaminae. They performed biplane cinemographic radiography on radiopaque beads at two cardiac sites in open-chest dogs. The premise for the study was the observation that myocyte thickening alone could account for only a fifth of ventricular wall thickening. This study utilized small regions of the LV anterior free wall and septum from 10 mongrel dog hearts. Inter-individual variation in sheet structure is not recorded in this report.

This work was extended by Costa et al. [44] using the same methodology, who described generally smoothly varying cleavage planes from epicardium to endocardium in three of six dogs, but with abrupt changes associated with the trabeculata–compacta interface (the zone linking trabeculated endocardium and compact subendocardium) in a further three dogs. They confirmed that the laminar nature of the myocardium is critical for normal ventricular dynamics. They also demonstrated that in addition to the sliding of sheets due to interlaminar shear, the sheets participated in myocardial contraction dynamically by extending during systole. They showed the shear component to contribute ~40% and the extension component ~60% to ventricular transmural thickening.

Arts et al. [45] carried out a strain modelling analysis of the data from the above Dokos et al. [58] study and predicted two distinct sheet populations. Histologically determined sheet angles were pooled for all six hearts and revealed a bimodal distribution by cardiac location, of approximately normal sheet orientations. Theoretical modelling of sheet structure using mechanics supported these measurements. They concluded that sheet orientation is not a unique function of the transmural location but occurs in two distinct populations.

Dokos et al. [58] examined laminar shear in samples of excised ventricular myocardium from six pigs. They demonstrated dual cardiac sheet orientations, describing a predominant orientation and an orientation approximately normal to this. They showed that patterns of sheet intersection were not uniform in the circumferential direction, varying significantly over sub-millimetre dimensions. Inter-individual variation in sheet architecture was not characterised or reported.

Jiang et al. [81] produced high-resolution three-dimensional reconstructions of sheet orientation from DT-MRI of the murine heart. They showed whole heart visualisations of sheet angles extracted from the secondary and tertiary eigenvectors (including β'). These images display regional localisations of structure in the LV, but little qualitative and no quantitative descriptions are provided and inter-individual variations are not characterised.

Harrington et al. [11] also identified two sheet populations from a histological study of tissue blocks from the LV interpapillary muscle region of five sheep. They proposed a model with alternating, approximately normal (in the longitudinal–radial plane) positive and negative sheet orientations across the ventricular wall with sheet orientation dependent on transmural depth; positive epicardial, negative mid-myocardial and positive endocardial orientation (Model 5, Fig. 2, part 5). There was relatively little inter-individual variation in sheet orientation (maximum cluster SD = 9 in the reported sheet angle). This model is a satisfactory explanation of the study findings, but is an attempt to extrapolate the features of a small tissue block to a wider region and is hence only valid in a LV without localised heterogeneous aggregations of structure.

Helm et al. [27,83] describe the two sheet populations in DT-MRI data from seven canine hearts. In some LV regions of individual hearts (including the lateral wall) sheets toggled from negative to positive orientation and sometimes back. Combined analysis of sheet orientation from these seven hearts reveals a bimodal distribution of two alternative angle populations. Twenty-five out of 30 selected anatomical regions have such a bimodal distribution (Fig. 8g), having one dominant sheet orientation cluster present in most hearts and one lesser orientation cluster for the remaining hearts. The two sheet populations are approximately normal. The possibility of eigenvector sorting error producing the bimodal distribution was excluded by a t-test analysis of the secondary and tertiary eigenvector statistical differences.

Cheng et al. [9] identified two sheet populations in the lateral interpapillary LV muscle region of seven sheep and applied a combined strain modelling and histological method. Sheets were found to exhibit a pleated or accordion-like orientation with alternating sign from sub-epicardium to subendocardium in a more pronounced manner in the sheep than has been described in the canine by Helm et al. above [27,83]. Intriguingly, they found that systolic fibre shortening was converted to radial wall thickening by a fundamentally different mix of sheet mechanisms at different wall depths. Discontinuities of sheet structure are described at the 20–50% depth and at 50–80% depth with a transmural pleated configuration.

The rat DT-MRI study of Chen et al. [16] demonstrates by DT-MRI the important role played by laminar shear (sheet sliding) in myocardial contraction. This study is similar in approach to Spotnitz et al. [46], but uses DT-MRI in parallel to histology. The presence of sheets of myolaminae (not specifically mentioned by Spotnitz et al.) as well as cleavage planes is confirmed in this study. Separate populations of sheet orientation are not commented upon despite their presence in their DT-MRI results. The paper instead discusses a simple sheet orientation pattern identical to that described by LeGrice et al. [56] – ‘positive sheet angles at the base to negative sheet angles at the apex’. The data presentation does not allow analysis of inter-individual sheet variation within groups, but it is noted that significant variability of sheet angle measurements was present. This was explained by (i) DT-MRI resolution limitations voxel averaging over ~20 sheets, (ii) sheet branching and (iii) sorting error of the secondary and tertiary eigenvector.

Rohmer et al. [31] carried out a dual algorithmic extraction and visualisation of fibre structure and sheet structure from one human DT-MRI dataset. The fibre-structure model produced is similar to other published algorithmically extracted fibre maps [28–30]. The sheet model is broadly similar to that proposed by LeGrice et al. [34,43,56,57], with individual concave or convex transmural endocardial to epicardial sheets. Overall, the modelled sheet architecture was complex, with sheet orientation depending on myocardial location. Sheets have greater curvature in the apex. The simple laminar structure model (Fig. 2, part 4) was used as a basis for the development of the sheet reconstruction algorithm, and as such single transmural
sheets with convex/concave surfaces are visualised, and the abutments between separate positive and negative sheet populations are not seen. If this alternative conception of dual positive and negative sheet populations had been applied in algorithm development, the visualisation of sheet structure would not be of continuous warped transmural sheets.

In summary, the literature reveals (i) the two mechanisms by which sheet structure acts as the central mechanism of wall deformation (laminar extension and laminar sliding) and an older, now rejected model of fibre interdigitation, (ii) the existence of two sheet populations with a complex distribution throughout the left ventricle that is somewhat variable between species and (iii) the inter-individual variation in the canine in the distribution of sheet orientation throughout the heart. The inter-individual variation is highly constrained with sheet orientation belonging to clusters. The nature of the boundaries between these sheet populations is poorly understood. Three-dimensional maps of sheet structure throughout the entire ventricle have been published but their features have not been described, nor have they been formalised into models and only limited statistical analysis has been performed. Those studies which have described topological detail of sheet structure have only examined small slabs of the lateral left ventricle, with emphasis on sheet changes between systole and diastole [16] and in cardiac disease [13,24].

10.3. A novel description of canine left ventricular tertiary structure

The data acquired by Helm et al. and Beg et al. [13,26,27,83] have been extended into a comprehensive resource of cardiac DT-MRI data available from the Centre for Cardiovascular Biomechanics and Modelling (CCBM; http://www.ccbm.jhu.edu/index.php). We revisualised these data alongside DT-MRI data from three mongrel dog hearts, acquired and pre-processed by Hsu et al. [35], then at the Center for In Vivo Microscopy, Duke University Medical Center.

As discussed, Helm et al. [83] provide a description of bimodal populations (Fig. 8g) of sheet orientation when ventricular regions are compared between dogs by cardiac location. Although visualisations are presented in these reviewed studies, these are not comprehensive and do not explore the variable distribution of cardiac structure in different canine hearts, or how this bimodal angle distribution is distributed throughout individual hearts in three dimensions. A hypothesis was formed on the basis of the findings of Helm et al. [83] that (i) bimodal distributions of sheet orientation, (ii) other observations on sheet structure in the literature and (iii) consistently varying fibre helix angles in all studied hearts: cardiac sheet structure can belong to a range of structural forms, with the smooth separation of bimodal peaks indicating a distribution of form, rather than two or more distinct structural models. The range of forms adopted by sheet structure is constrained by the myocyte fibre orientation (as these cells course through the sheets) which has a uniform helical organisation in the left ventricle.

In order to investigate this hypothesis we have revisualised the sheet orientation from selected basal, equatorial and apical slices from 12 normal canine hearts (Fig. 7a). Established DT-MRI visualisation methods [83] and tools were applied to this public domain data and anatomical views were selected to provide a comprehensive presentation of sheet architecture in three dimensions in individual hearts. The visualisations presented are of the $\beta'$-sheet angle and the angular scale $90^\circ - \beta' < 90^\circ$ is adopted. The sheet structures in visualised short-axis slices are described.

The structural continuity within individual hearts underly the bimodal sheet orientation described above can be seen. Some hearts (6, 16, 12, 11) have predominant $\beta'$-sheet angles greater than 22.5° or less than 22.5° ($-22.5 < \beta' > +22.5$) with relatively little in short-axis plane sheet orientation (henceforth referred to as SA-planar). Other hearts have larger areas of SA-planar ($-22.5 < \beta' < +22.5$) $\beta'$-sheet angle (9, 15, 13, 10, A, C and B).

Eleven hearts (9, 15, 13, 10, 8, 6, 16, 11, 12, A and C) have predominantly negative $\beta'$-sheet angle in the LV equatorial posterior region, with one heart (B) having predominantly positive $\beta'$-sheet angle in this zone. The equatorial antero-lateral LV free wall is predominantly planar in four (9, 15, 10 and B), negative in four (13, 8, 6, 11) and positive in four (16, 12, A and C) hearts.

$\beta'$ in the basal anterior LV free wall is positive in ten (9, 15, 13, 10, 8, 16, 12, A and C), mixed in one (10), negative in one (B) and planar in one (6) heart.

The three-dimensional cut out slab (Fig. 8e) shows that described regions of confluent positive (red) or negative (blue) sheet angulation are continuous stacks in three dimensions. The green SA-planar areas also form contiguous regions merging more gradually with red and blue stacks. It is inappropriate to describe these features as bands — they are best interpreted as stacks of similar sheet orientation, reminiscent of a stack of angled books.

This variability in sheet structure is far from random. As noted above and in Helm et al. [83], sheet angles belong to two populations following a bimodal distribution with average peaks at 45.5° and 117.6° (equivalent to $-44.5^\circ$ and $27.6^\circ$ on $\beta'$-sheet angle scale adopted here, see Fig. 8g). Sheet structure is present as circumferentially confluent zones of near-continuous orientation, often having sharp terminations. Regions of positive and negative $\beta'$-sheet angle often run adjacent in the circumferential direction with sudden angular change at their abutments. In other cardiac regions sheets more gently merge into each other.

As such, positive and negative sheet stacks are not features of a regularly undulating continuum of sheet structure. This is demonstrated by the sheet orientation 10° contour map in Fig. 8b, which shows that abutting red and blue confluent sheet zones have very steep angular change between them. It should also be noted that, although this angular change is steep, it is not discontinuous; they represent regions of rapid orientation change with distance (Fig. 8b, iii).

Three hearts (15, 16, 11), representative of the principal sheet features of the continuum of structure, are visualised in three dimensions in Fig. 8a. The apex of heart 15 has an inner ring of positive sheets encircled by an outer ring of negative $\beta'$-sheet angle. At the apical origin of the LV
papillary muscles the negative sheet population has dominated the positive population. Basal to this, at the basal extent of the papillary muscles, positive sheets dominate the anterior LV, while negative sheets dominate the posterior LV. In the basal LV, overall positive sheet predominates.

Heart 16 (Fig. 8a) has a similar overall LV architecture but has significantly greater positive $\beta$-sheet angle in the anterior LV throughout the equatorial zone.

In heart C (Fig. 8a) positive $\beta$-sheet angle extends from the equatorial zone to the apical zone, where positive and negative sheets are both present.

Fig. 8c and d is a visualisation of posterior view of heart 15, showing $\beta$-sheet angle. It should be noted that the underlying sheet structure is similar to that described for the HVMB in Fig. 4.

Fig. 8f shows a top view of an equatorial LV slice of histologically derived sheet orientation. The data are derived from LeGrice et al. [27]. The $\beta$-sheet angle orientation shown is not identical to any of the 12 canine hearts in Fig. 7a but fits into the continuum of structure being intermediate between hearts 12 and B.

10.4. Discussion of sheet stacks

How do the described stacks of laminar structure fit into the general cardiac sheet literature? All the sheet structural features described in the reviewed literature — two populations [9,11,13,27,44,45,58,83], toggling positive/negative accordion pattern [11,27], patches [45], smoothly varying and abrupt changes [44,58], inter-individual variation of sheet orientation with a bimodal distribution [27,45,83] — can be accommodated by the restrained structural continuum model proposed here.

The sheet stacks described in this study are often clearly defined. Sheet reconstructions showing the same general pattern stacks have been published for the dog (Figs. 7 and 8) [27], mouse [81], rabbit [27] and rat [16], but without explicit descriptive identification of structures analogous to our stacks. Crescent-shaped stacks are visible in the murine DT-MRI sheet reconstruction of Jiang et al. [81], similar in morphology to those visualised here.

The rat study of Chen et al. [16] reveals prominent abutting positive and negative sheet stacks; however, these have differences in conformation to those described here for the dog. There are no SA-planar regions reported, rather overlapping crescents of opposite polarity (Fig. 6e). The boundary zones between positive and negative stacks are very sharp, suggesting a sudden change of orientation.

Some of the sheet features described in this review are present in the registered canine DT-MRI visualisations of Helm et al. [13], but certain features are more pronounced.

RV endocardium. (a) $\beta$-sheet angle maps from three-dimensional reconstruction of DT-MRI for 12 normal canine hearts. Numbered hearts are from datasets, courtesy of Patrick Helm and Raimond Winslow et al. at The Center for Cardiovascular Bioinformatics & Modeling and Elliot McVeigh at the National Institute of Health. Data for hearts denoted by capital letters were acquired from Hsu et al. [35], then at the Center for In Vivo Microscopy, Duke University Medical Center. A top view of basal, equatorial and apical slices (left to right) is shown alongside an approximately anterior view of a mid-heart slice. The hearts are ordered by eye so as to demonstrate a distribution of changing $\beta$-sheet angle properties from heart 9 (top) to heart 11 (bottom). (b) $\beta$-sheet angle scale bar from $-90^\circ$ red to $-90^\circ$ blue.

Fig. 7. Reconstruction of canine ventricular $\beta$-sheet angle. $\beta$-sheet angle has been defined as the angle between the transverse plane and the projection of the secondary eigenvector onto the radial plane [81]. Positive $\beta$-sheet angles rise to the heart base from endocardium to epicardium (or LV endocardium to RV endocardium).
Fig. 8. (a) Anterior view of three-dimensional detailed representation of $\beta$-sheet angle from hearts 15 (top), 16 (middle) and C (bottom) showing detail of sheet morphology in three dimensions. Basal, equatorial and apical slices are shown from these three hearts, which represent major forms within the continuum of cardiac structure. $\beta$-sheet angle is coloured as shown in the scale bar (h) from $+90^\circ$ red to $-90^\circ$ blue. (b) Top view of short axis cross-section of lateral LV free wall of heart 16. In the top panel $\beta$-sheet angle is coloured as shown in the scale bar (h) from $+90^\circ$ red to $-90^\circ$ blue, in the bottom panel a cardiac outline is shown in black with no
while others are more subtle. This is likely to be a consequence of the data registration and image averaging methodology utilised. This algorithmic methodology combines data from several heart DT-MRI scans into a single spatially registered and combined average visualisation of sheet structure. This emphasises features with fixed presence between hearts, and smooths out those that have a more variable presence between individuals. Given the described bimodal angle population present between individuals [27, 83] and the visualisation presented here the application of such methods to cardiac sheet structure data is likely to be of limited value.

10.5. Cardiac sheet structure features, pinnation, fibre paths and the HVMB

The described sheet structures are related in location, although not coincident, with the DT-MRI primary eigenvector transitions (Fig. 5a–d). The precise relationship between the DT-MRI determined sheet features and the histologically described (i) pinnation features of Hort [54] (Fig. 5e and f) or (ii) fibre orientation regions of Greenbaum et al. [50] and Anderson et al. [4] (Fig. 5) cannot be determined without further studies, as the histological features are from hearts of unknown sheet structure. As they all form part of the same complex continuum of structure, these features must be related.

11. Sheet stack boundaries

11.1. The nature of sheet stack boundaries

The DT-MRI data provide a picture of global sheet behaviour throughout the ventricles. Due to resolution limitations (discussed in detail below) little information is provided about the nature of sheet interactions at stack boundaries. The nature of these boundaries is best studied by high-resolution three-dimensional confocal microscopy. Two such studies [20, 21] have analysed lateral LV transmural tissue blocks from rats (producing the data shown in Fig. 6a–c). These reports have focused on technique validation for use in electrical modelling [87], rather than the anatomical structure revealed. Detailed study of the microstructural architecture in Fig. 6 and the video from which these images were extracted reveals intriguing and unreported detail of myolaminar architecture.

Two sheet populations can be seen to co-exist in the same slab of tissue. A highly intricate but ordered pattern of sheet merging is observed, with \( \beta \approx 45^\circ \) and \( \beta \approx 45^\circ \) myolaminae coalescing on myolaminae of rhomboid cross-section in the mid-myocardium. These rhomboid structures appear to carry the circumferential fibres [4, 48] (as described in Model 3 [48]). Merging is subtle, and working back from the rhomboid central zone, two sheet populations can be seen to co-exist simultaneously in the same region (Fig. 6b and c, blue circles). On moving towards the epicardium or endocardium, the positive or negative sheet orientation dominates.

The detail of the sheet interactions is best considered in the light of the rat global sheet architecture as revealed from DT-MRI (Fig. 6e). These DT-MRI data are from a different rat (and a different research group) but the same sheet features are revealed (negative endocardial sheets meeting positive epicardial sheets). In the transition zone, yellow sheet orientation (almost SA-planar, \( \beta \approx 15^\circ \)) is recorded.

No detailed confocal data are currently available for the canine, so the nature of the boundaries between the \( \beta \approx \pm 45^\circ \) sheet stacks and the transverse \( \beta \approx 0^\circ \) sheet stacks is unknown. There may be a similar pattern of merging with co-dominance at the stack boundaries, or this may be limited to the intersection regions between positive and negative \( \beta \approx \pm 45^\circ \) sheet stacks, with a more gradual change of a single sheet population in the rest of the myocardium.

11.2. Role of stack interaction boundaries (positive to negative sheet stack abutments)

No observations of the role sheet merging across stack boundaries have been undertaken. Given the complex merging of laminae across these boundaries it is possible that sheet sliding and extension is continuous across this zone but with the changing of the direction of strain. It is possible that the loose structure of the merging zone between positive and negative sheets, with the rhomboid myolaminae which resemble bundles of myocytes, may allow the interdigitation proposed by Hort [53–55] as a mechanism of myocardial thickness change between systole and diastole. This inter-leaving would not be possible in regions of confluent sheets (predominant positive or negative orientation, towards the epicardium and endocardium of Fig. 6a–c). Inter-leaving was not observed by Spotnitz et al. [46], but their study concentrated on regions of confluent cleavage planes, and they did not discuss dual populations of cleavage planes (despite their presence in their data), and as such they could have different characteristics.
have missed this finding. Indeed, such thickening of myocyte bundles and obliteration of cleavage planes may be observed in the positive to negative cleavage plane abutment to the left of Plate 2 in [46] (note that the myocardial images in this plate are aligned images from different fixed rat hearts).

The junctional region may act as region of meeting of forces with the sheet populations acting antagonistically and/or to stabilise each other, as explored by Sengupta et al. [88]. During isovolumic contraction, shortening of subendocardial myofibres is accompanied by stretching (and hence stiffening) of subepicardial fibres. These authors hypothesise that this could form a functional analogue of the ‘rigid external cylinder’ proposed by Torrent-Guasp et al.

12. Technical limitations of techniques

12.1. Technique focus

It is tempting to view the data produced from novel imaging techniques as unbiased, absolute and definitive – not as artefact-prone and subject to investigator bias as with traditional techniques. However, equal care is needed in the interpretation of these data – its weakness can be that it seems to produce all the answers. In fact, each imaging technique has a focus – a level of structure which it reveals – and in so doing it can obscure higher- and lower-order structure. Confocal imaging of tissue blocks has revealed detailed laminar structure and cross-meshing, but obscures overall sheet patterns and predominant fibre orientation. Primary eigenvector fibre angles and optical polarisation maps reveal primary fibre trajectory but obscure local branching and hide sheet structure. Indeed, DT-MRI reports presenting only the primary eigenvector (principal fibre orientation) have been used to argue against regional structural differences within the ventricles [30].

12.2. DT-MRI: resolution

There are technical reasons for the focus of each technique. The resolution of DT-MRI applied to cardiac studies has generally been approximately 300 µm within slices and 1000 µm between slices [30,35—40]. This is not sufficient to depict the finer structure of myocardial laminae [16] (of dimension ~50 µm in dog [56]). The measured β-sheet angle reported in each DT-MRI voxel will be an averaging of ~6—20 sheets. As discussed, this has implications for the power of DT-MRI to resolve sheet behaviour at stack boundaries. At current resolution it produces a schema of locally averaged sheet direction – so results should be interpreted in this light.

12.3. DT-MRI: eigenvector sorting

In DT-MRI processing, sheet angle is calculated from the secondary eigenvector and sheet normal from the tertiary eigenvector. These values are sorted by magnitude, the secondary always being greater than the tertiary. It has been hypothesised that an eigenvector-sorting problem could occur where secondary and tertiary are similar in magnitude [25,83], and that this could invalidate determined sheet angles. A statistical analysis by Helm et al. [83] addressed this question and concluded that sorting was robust to this error, but the study did not consider sheet co-dominance in regions of sheet merging. Analysis of the microstructural detail of sheet features in confocal data reveals that the true structure consists of complex sheet merging with co-dominance of positive and negative sheets, while in DT-MRI data this is represented by ~SA-planar sheet conformation. An implication is that histological confirmation of detail of sheet structure is required, and that this should be incorporated with the statistical analysis of DT-MRI data to allow identification of co-dominant sheet areas. The canine histological sheet orientation data (Fig. 8f) provide some validation and show that the gross patterning, if not the local microstructural detail, of sheet features is robust to sorting artefact.

12.4. Confocal microscopy: fixation artefact

Tseng et al. [25] described considerable distortion of myolaminae orientation between ink prints from un-fixed tissue and histological measurements after fixation and sectioning. It is likely that the cleavage planes in Fig. 6a–c have opened up due to shrinkage and dehydration in the fixation process.

12.5. Confocal microscopy: real myolaminae and artefact myolamine

There has been some debate about the presence of myolaminae and their structure. Harrington et al. [11] divided planes present in sectioned fixed tissue into true laminae (the predominant sheet orientation visible on the sectioned frozen tissue surface and on un-sectioned tissue) and artefact. On placing the tissue on a slide, gaps opened up dividing the tissue into laminae along the visible sheet orientation over a period of 3 min. On observing the sections for longer periods, additional gaps opened up, which they described as cracks and dehydration artefact. Are any of the sheet-merging features described in Fig. 6 artefact? This is unlikely – the rat hearts were embedded in epoxy resin after fixation, fixation is not known to produce crack artefacts, and cracking of the epoxy resin block on sectioning would be grossly visible, would produce optical distortion and would not occur along planes of equal thickness matching known myolaminae width (four to six fibres). It is also unlikely that perpendicular planes would be created, meeting so regularly on rhomboid myolamine. It is possible that some of the cracking observed by Harrington et al. [11] were real cleavage planes opening up, or that much greater care is needed with frozen tissue than fixed tissue.

These technical problems highlight the need for an intelligent integration of results from the different techniques, building together of schema of heart structure to form a united concept of cardiac architecture.

13. Validity of the proposed sheet stack structure by Grant’s principles

In drawing together a novel description of cardiac structure one risks clouding the debate rather than clarifying
it. To avoid this it is necessary to place the model in a clear context of the known structural facts and the principles as outlined by Grant. An important component in this is to consider the technical limitations and possible artefact due to experimental method.

As evidenced by the contour and microstructural description of boundary zones between sheet stacks, the new model fits into the principle of a syncytial myocardial community. It is also appreciated that the sheet structure obtained from DT-MRI secondary eigenvector data is only one level on which the myocardium can be approached — and that other valid approaches reveal no discernable detail of sheet structure (see the primary eigenvector fibre orientation data of Fig. 3). In this sense the sheet structure is best viewed as a schema of one level of cardiac structure — whole heart sheet behaviour, underlying the higher-order schema of predominant fibre direction and overlying the schema of local myolaminar meshing. The DT-MRI technique is inherently statistical and of uniform bias across the entire structure. The $\beta'$-sheet angle maps (Fig. 8) clearly demonstrate the described sheet features. A complete statistical analysis of the stack structures from grouped hearts and across species is required, but this preliminary analysis demonstrates that Grants’ criterion of statistical uniqueness is likely to be satisfied. Debate on whether the sheet stacks described are a schema of cardiac structure or distinct anatomical bundles (as described by Grant) is largely semantic. Addressing whether the stacks constitute unique physiological bundles is also a naive approach. It is more useful to address the role played by interaction of these ubiquitous, intricately intermeshed structures in cardiac excitation and contraction.

14. Implications

The distribution and merging pattern of the sheet populations lead to many questions to be answered by future directed research. What are the sheet sliding (shear) and sheet extension implications for the sheet stack merging pattern? Is the heterogeneity of myocardial action potential morphology [89] related to sheet populations rather than simple transmural or base–apical ventricular location?

Yet, more questions are raised by the heterogeneity of cardiac sheet structure between dogs. Reviewed evidence points to the critical role of sheets in myocardial contraction, so how do such different structures act to produce the same function? Are some sheet conformations more efficient, more resistant to myocardial failure or of different electrical conduction and arrhythmogenesis potential? Do inbred populations such as laboratory rats and mice have more uniform cardiac sheet structure, as suggested by the data in Spotnitz et al. [46]? Do dogs having perhaps the greatest body form heterogeneity of any domesticated species (due to selective breeding and paedomorphism [90]) also have much greater cardiac heterogeneity than man or pigs? Does the maintenance of such a regular fibre angle distribution despite $\beta'$-sheet angle heterogeneity reflect a greater importance of fibre angle than sheet orientation in myocardial contractile function, contrary to evidence in the reviewed literature?

The recent demonstration of sheets in echocardiography [88] may provide a methodology to study the dynamics of sheet interaction throughout the cardiac cycle, particularly if sheets can be visualised in three-dimensional ultrasound images.

The cardiac topology as described by Torrent-Guasp et al., although reflecting some genuine features of heart structure, is a schema of minor importance and is unlikely to have physiological consequences. It does not apply equally to all canine individuals and may not apply equally in other species.

Fibre orientation and sheet architecture are known to be important in cardiac contraction [16,44,91] and are remodelled in cardiac disease [13,24,85,92]. The importance of sheet stacks described in this review has not been studied, nor has their alteration in cardiomyopathy. Given the known importance of sheets in cardiac contraction it is likely that sheet stacks will also be functionally important, particularly the sharp orientation changes at boundaries between positive and negative sheets.

15. Conclusions

The novel description of cardiac sheet stacks in this review is compatible with the published histological and DT-MRI reports of dual sheet populations and inter-individual variability of sheet structure.

Greenbalm et al. [50] concluded that there was a complex regional variation in left ventricular structure, that modelling based on uniform wall structure could not explain wall movement and that traditional dissection techniques and histology would be inadequate to fully characterise possible regional differences in LV arrangement.

New imaging technologies allow greater characterisation of regional differences, but the application of this knowledge to modelling wall movement and guiding surgery is in its infancy. This report of sheet structure describes the overall pattern in the left ventricle as revealed by the DT-MRI secondary eigenvector and confocal microscopy. It does not constitute a finalised complete model of cardiac structure, but features described here must be incorporated into any future such model.

Note on images: All short-axis (cross-section) cardiac views are presented as top views looking towards the cardiac apex.

Acknowledgements

The canine ventricular DT-MRI datasets presented in Figs. 3, 5 and 7 were acquired and pre-processed by Hsu et al. [35], then at the Center for In Vivo Microscopy, Duke University Medical Center, and the datasets presented in Fig. 8 were acquired and pre-processed by Patrick Helm and Raimond Winslow et al. at the Center for Cardiovascular Bioinformatics & Modeling and Elliot McVeigh at the National Institute of Health.

References


