Three dimensional echocardiography for the assessment of mitral valve disease

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Whereas conventional two dimensional (2D) echocardiography is crucial to our understanding of the complex anatomy and three dimensional (3D) spatial relationships of cardiac structures, it requires the mental integration of a limited number of 2D imaging planes. This mental 3D reconstruction is inherently variable according to observer experience and expertise, and can only be described to other clinicians (such as surgeons) rather than displayed reproducibly. The display of cardiac anatomy in three dimensions from any perspective would have clear advantages over conventional 2D imaging and provide an insight into the functional and anatomic properties of cardiac structures. Recent advances in ultrasound and computer technology have been combined such that dynamic 3D echocardiographic imaging is now a practical reality.

Three dimensional echocardiography (3DE) has been shown to be more accurate than 2DE in the quantification of cardiac volumes. These studies used either manually contoured, static “wire frame” reconstructions or dynamic “volumetric” automated reconstruction technology that is now commercially available—we concentrate on the latter methodology in this review. The benefits of 3DE are particularly well suited to the study of the mitral valve given its complex morphology and the importance of delineating its anatomy precisely in various pathological states. This was shown by Levine et al who used wire frame reconstruction of the mitral valve to define the 3D morphology of the mitral annulus and its relationship to mitral leaflet position, thereby clarifying the echocardiographic definition of mitral valve prolapse. The assessment of patients with mitral valve disease is one of the most promising clinical applications of this technology.

Dataset acquisition, processing and reconstruction

A 3D dataset is composed of anatomical information from multiple 2D cross sectional images. For reconstruction of the mitral valve in adult patients, transoesophageal echocardiography (TOE) is the preferred approach for 2D image acquisition as it offers a relatively stable site for the imaging probe and superior resolution of the mitral valve apparatus. Images from a commercially available multiplane TOE probe are interfaced with a 3D computer system which incorporates the steering logic for acquisition of a rotational dataset and software for 3D reconstruction and display. With the mitral leaflets in the transverse mid-oesophageal view, the multiplane TOE probe is usually rotated at 2° or 3° increments over 180° to give 90 or 60 sequential 2D cross sections, which are digitised to form a conical dataset. Optimal temporal and spatial registration is achieved by ECG and respiratory gating. Offline processing involves the conversion from polar to cubic Cartesian co-ordinates and interpolation of missing information between 2D slices. From the resultant dataset, novel 2D cut planes in any orientation can be selected (anyplane echo) and multiple parallel cross sectional 2D slices can be generated in any desired plane (paraplane echo). A volume rendered 3D image of the mitral valve can be reconstructed from any perspective (fig 1). Threshold limits are used to separate cardiac structures from blood pool and background. Brightness and shading provide perception of depth. With the added dimension of time we are able to study in detail the motion of the valve during the cardiac cycle.

Mitral stenosis

Three dimensional echocardiography has a role in both the quantitative and qualitative assessment of mitral stenosis. Three dimensional echocardiography overcomes the limitations of image plane positioning inherent in conventional 2DE and offers a more precise approach to measurement of mitral valve area. Paraplane 3DE allows the 2D short axis slice in the optimum plane of the orifice to be selected from the 3D dataset and the smallest complete orifice can be directly measured by planimetry (fig 2). Anterior leaflet calcification often produces acoustic shadows which obscure the mitral orifice in the 2D transthoracic short axis plane. Three dimensional TOE overcomes this problem by visualising the mitral valve from behind, so that acoustic shadows are cast into the left ventricle rather than over the leaflet tips. Chen et al assessed 15 patients with mitral stenosis and showed this...
technique to correlate more closely with the Doppler pressure half time derived area compared with 2D echo planimetry. A second study of 54 patients showed 3D TOE to be superior to 2D transthoracic echo for planimetry of mitral valve area when compared with the area measured at surgery with calibrated dilators. We have reported the first in vitro study to validate the accuracy of 3D TOE for the measurement of mitral valve area. In our study, porcine mitral valves were prepared with the commissures glued to simulate mitral stenosis. These valves were scanned in a water bath using 3D TOE. The reference standard valve area was determined using a digital photographic technique. Paraplane 3D TOE was shown to be more accurate than any previously described technique for quantifying mitral valve area and could become the “gold standard”.

The shape of the mitral valve leaflets proximal to the orifice has an impact on the flow dynamics across a valve. 3DE with stereolithographic modelling has been used to demonstrate that flat shaped valves cause a higher pressure gradient for the same anatomic area and flow rate compared with “funnel” shaped valves. Thus, 3DE provides insights into mitral leaflet geometry which could refine our assessment of mitral stenosis.

3DE also appears to be of value in the assessment of patients undergoing balloon mitral valvotomy (BMV) where the dominant mechanism is splitting of fused mitral commissures. Studies using 2DE have shown that commissural morphology is a powerful predictor of outcome after BMV and at our institution this forms the cornerstone of assessment of patients with mitral stenosis referred for valvotomy. Viewed from the left atrium, 3D reconstruction of mitral stenosis displays the restricted orifice, thickened leaflet margins and prominent left atrial appendage (fig 3A). In our experience, the volume rendered 3D display provides improved visualisation of mitral commissural fusion, particularly when the leaflets are viewed from the perspective of looking upwards from the left ventricle (fig 3B). Following balloon valvotomy, 3DE also defines clearly the extent and site of commissural splitting which may be symmetrical (fig 3C) or eccentric (fig 4). Other investigators have reported improved imaging of the mitral commissures with 3DE compared with 2D TOE. Furthermore, 3DE assessment of commissural splitting following balloon inflation has been shown to relate to increase in mitral valve area.

Mitrval valve prolapse
Mitrval valve prolapse is a frequently encountered problem in clinical cardiology and the most common cause of isolated mitral regurgitation requiring surgical treatment. Mitral repair is associated with better outcome than valve replacement but demands a detailed understanding of valve morphology. The complex anatomy of the mitral valve makes interpretation of conventional 2D images difficult and is occasionally misleading. Dynamic volume rendered 3DE of the mitral valve displays both leaflets in their entirety from any desired perspective. This allows clear visualisation of the site and extent of leaflet prolapse during systole (figs 5 and 6). A number of studies have demonstrated the feasibility of 3DE in optimally displaying mitral valve prolapse. The accuracy of 3DE in localising the involved leaflets has been confirmed at surgery. By elucidating the non-planar shape of the mitral annulus and its spacial relation to the mitral leaflets, 3DE might also improve the

Figure 2 3D paraplane echo for measurement of mitral valve area. A series of parallel 2D short axis cross sections are generated in the optimal plane of the mitral valve orifice. The 2D slice defining the smallest complete orifice is selected and the valve area measured by planimetry.

Figure 3 3DE of rheumatic mitral stenosis. 3D images reconstructed from the left atrial (A) and left ventricular (B) perspectives. The left ventricular view clearly shows thickened leaflets with a restricted orifice due to symmetrical commissural fusion. (C) shows the same valve after successful balloon mitral valvotomy demonstrating increase in orifice area due to bilateral commissural splitting.
diagnostic sensitivity and specificity in mitral valve prolapse.\textsuperscript{15} Reconstruction of the mitral valve en face from the left atrial perspective provides a view of the valve similar to that seen intraoperatively by the cardiac surgeon, with the major advantage of displaying the dynamic motion of the valve within the beating heart.\textsuperscript{12} Clearly, 3DE has great potential to facilitate the preoperative planning of mitral valve repair.

With expected advances in computer technology and interactive software, we anticipate accurate simulation of surgery prior to repair operations in the form of “virtual repair”.

**Mitrail regurgitation**

As yet there is no widely accepted, non-invasive technique for accurate quantitation of mitral regurgitation. The regurgitant orifice in severe mitral regurgitation cannot be defined by 2DE, and calculations based on Doppler flow are technically difficult and not widely applied. 3DE allows direct visualisation and planimetry of the regurgitant orifice in the optimal 2D plane, and measurements obtained by this method have been shown to correlate well with flow convergence derived measurements.\textsuperscript{15} However, this is only applicable in severe regurgitation. Recently, the ability to reconstruct Doppler colour flow has provided an insight into the mechanism and shape of mitral regurgitant jets.\textsuperscript{17–18} Three dimensional colour jet volume has been shown to correlate significantly with regurgitant volumes and may improve the assessment of eccentric MR because the full extent of these jets is better appreciated.

An important development in echocardiography has been the use of colour Doppler to evaluate the regurgitant flow convergence zone and calculate its proximal isovelocity surface area (PISA). Accurate measurement of the PISA would allow precise quantitation of mitral regurgitant volume, flow rate and regurgitant orifice. However, this method relies on assumptions regarding the geometry of the flow convergence zone—for example, that it is hemispheric. Cape \textit{et al} have shown that the flow convergence zone is a complex 3D shape dependent on the shape of the regurgitant orifice\textsuperscript{19}; this can be reconstructed precisely using 3DE in vitro\textsuperscript{20} thereby measuring PISA directly, without geometric assumptions, upon which accurate calculations of flow rate can be based. The quantification of mitral regurgitant volume and effective regurgitant orifice area remain important yet elusive goals; we await clinical evidence to support these theoretical advantages of 3DE.

**Limitations of three dimensional echocardiography**

The standard of the 3D reconstructed display depends critically on the quality of the original 2D cross sectional images. Until recently, in adult patients, this necessitated TOE. However, the development of harmonic imaging has made it feasible to reconstruct from a transthoracic rotational dataset. Minor movements of either patient or operator will distort the images and result in dropout which may be misinterpreted. Atrial fibrillation or a variable respiratory pattern prolongs the acquisition time and impairs the dataset resulting in artefact. Operator dependent changes in threshold settings, which define the tissue-blood interface on the 3D rendered display, can affect the apparent mitral orifice. Therefore, measurements on reconstructed images should be made with caution.
contrast in mitral stenosis hinders reconstruction of the valve from the left atrial perspective; this can be minimised by reducing the probe frequency. Highly mobile structures such as a ball valve thrombus, vegetations, and ruptured chords are not easily seen. In our opinion, 3DE has not improved visualisation of the mitral subvalvar apparatus and areas of calcification are not apparent in the volume rendered display. At present this technology provides information which complements that gained from a comprehensive 2D and Doppler echocardiographic study.

Conclusions
In the past, the major practical limitation of 3DE was the long time required for raw data processing and image reconstruction and suboptimal 2D image quality. With improved ultrasound technology, particularly harmonic imaging and faster digital processing, these problems are being overcome, and 3D software is being integrated into modern echo machines. A rotational 3D TOE dataset can be acquired, processed, and displayed within 10 minutes and has been shown to be feasible and useful in the intraoperative setting. “Real time” 3D transthoracic probes have been developed and are already commercially available. These factors will enhance the clinical applicability of 3D echocardiography in the future.