Tissue Doppler imaging: current and potential clinical applications

D J A Price, D R Wallbridge, M J Stewart

Echocardiography when combined with spectral and colour flow Doppler is well established as a safe, non-invasive, and versatile diagnostic modality in cardiology, and is now the predominant technique used for evaluation of left ventricular function and for the assessment and quantification of valvar heart lesions. When combined with physiological or pharmacological stress, echocardiography also enables the identification of reversible myocardial ischaemia and myocardial contractile reserve.1 2 However, assessment of regional cardiac dysfunction at rest and during stress remains subjective and semiquantitative, with high interobserver variability.3–6

Doppler measurement of myocardial motion, using pulsed wave Doppler, was first proposed in 1989 but this technique allowed real time visualisation of only a single myocardial segment and its potential was not realised.7 Only later, with the development of colour flow algorithms to visualise myocardial motion, did the technique begin to gain clinical acceptability.8–10 Subsequent software development has led to improved temporal and spatial resolution and off line processing to enable quantification of multiple segments of myocardium in seconds. These technological advances have been matched by widespread clinical interest in the technique and an explosion of clinical research. We review the current state of knowledge of tissue Doppler imaging, emphasising current clinical applications and likely future roles.

Background

Doppler tissue imaging uses the same principles as colour flow Doppler mapping, applying standard autocorrelation processing but reversing high velocity and low amplitude filters such that the high amplitude/low velocity motion of tissue is displayed in preference to blood flow. As cardiac structures move in a velocity range 0.06 to 0.24 m/s, some 10 times slower than myocardial blood flow, and have an amplitude approximately 40 decibels higher, it is possible to obtain images of tissue Doppler motion of high resolution without significant artefact originating from the blood pool. In such images, each pixel displays one colour representing a mean velocity value. However, comparison with pulsed wave Doppler traces of myocardial motion has indicated that initial Doppler tissue images failed to accurately quantify each phase of the cardiac cycle. The normal cardiac cycle has four distinct peaks in systole and diastole (fig 1).

Figure 1 Normal myocardial velocity profile sampled from basal septum (apical four chamber view), showing typical pattern of two systolic and two diastolic peaks. S1, myocardial velocity associated with isovolumic contraction (3.0 cm/s); S2, peak systolic shortening velocity (9.0 cm/s); E, indicates the peak early diastolic myocardial relaxation velocity (~12.0 cm/s); A, late diastolic myocardial velocity associated with atrial contraction (~8.2 cm/s).
The low frame rate (25 Hz) of initial Doppler tissue imaging systems provided a temporal resolution which was insufficient for accurate depiction of each phase of myocardial motion. Technological advances using digital parallel processing techniques now allow two-dimensional (2D) Doppler tissue imaging with a 60° sector of up to 80 Hz. Combining such images with off line processing allows accurate quantification of regional myocardial motion in both systole and diastole, from multiple sites, with an image dataset being acquired within seconds.

Doppler tissue imaging terminology

In the original description of Doppler tissue imaging, three separate modalities were clearly distinguished.10 Myocardial Doppler velocity imaging uses standard colour coding to depict both velocity and direction of movement, with myocardial motion away from the transducer coded in blue, and towards the transducer coded in red. This bidirectional map allows semi-quantification of myocardial motion. This can be displayed as a standard 2D image or as a Doppler tissue M mode, where temporal resolution is greatly enhanced and differences in myocardial velocity from endocardium to epicardium can usually be appreciated. Recognition of this has led to the concept of the myocardial velocity gradient where the change in velocity from endocardium to epicardium is quantified.11-12 By effectively quantifying the motion within a myocardial segment, this technique helps to eliminate errors arising from the inability to correct for translational motion of the heart within the chest and to reduce angle dependency. Recently, this concept has been further developed with 2D depiction of strain rate imaging.

Doppler acceleration mode imaging refers to a unidirectional map, where the change of velocity (or acceleration) from one frame to another is displayed in a 2D image according to a separate multicolour scheme. This technique has found some application in electrophysiology, enabling recognition of the site of pre-excitation in the Wolf-Parkinson-White syndrome.13-15

Doppler tissue energy mapping, now more frequently referred to as power Doppler imaging, has also been developed as part of this technique. The displayed tissue map represents the intensity of the power spectrum of the received Doppler signal, reflecting the composition of tissue. Hence, this map is both angle and velocity independent. This imaging modality has found a particular role in myocardial contrast echocardiography. Utilisation of autocorrelation processing means that, with the introduction of a left heart contrast agent which displays non-linear oscillation under ultrasound interrogation, there is increased variation in the Doppler energy signal, displayed as a notable increase in colour intensity.16 Use of Doppler energy mapping, combined with a left heart contrast agent, led to one of the first descriptions of contrast within the myocardium following intravenous injection in humans.17 With further technical development, combined with intermittent imaging and pulse inversion technology, this now forms the basis of much ongoing research in myocardial contrast technology.18-19

This review, however, concentrates on Doppler tissue velocity imaging and, to avoid confusion, this is referred to as tissue velocity imaging (TVI) hereafter.

Development and normal values

Initial validation of tissue velocity imaging, using rotating phantoms and a water bath, confirmed accurate representation of motion within the physiological range, but with over estimation and poor spatial resolution of velocity in very low range.10 Further phantom studies also documented the axial and lateral resolution of the 2D velocity map to be 3 × 3 mm, implying resolution sufficient for the documentation of a significant regional wall motion abnormality. While such resolution does not approach that obtained with standard 2D echocardiography, it is important to emphasise and appreciate that these images are entirely dependent on Doppler data, obtained from frequency shift information rather than from the reflected signal amplitude required for a grey scale image. As a result, tissue velocity data can be obtained even from subjects where the traditional echo window is suboptimal, and in particular means that wherever an image is obtained accurate quantification of myocardial motion is still reliable, in contrast to standard B-Mode imaging (fig 2).

As a Doppler based technique angle dependency remains a crucial issue, leading to the potential for error when trying accurately to quantify myocardial motion. As a result the technique as applied at present cannot reliably measure, for example, the velocity of contraction of the lateral wall in the parasternal short
axis view. Contraction of mid-septum and posterior wall, which occurs parallel to the scan line, can be accurately assessed. However, at the much lower velocities encountered with tissue motion, the contribution of overall cardiac translational motion within the thorax becomes important resulting, for example, in underestimation of septal systolic velocity and overestimation of posterior wall systolic velocity in parasternal views, because of movement of the heart towards the transducer. These limitations may explain, in part, the relatively poor reproducibility of this aspect of the technique. While appearing a significant limitation of tissue velocity imaging, this effect may be minimised when measurement of velocity is relative to other myocardial regions, or of different points within the same region, rather than relative to the ultrasound transducer. This theory led to the development of the transmyocardial velocity gradient, for which normal values have been proposed, and more recently to myocardial strain rate imaging. Although still under evaluation, these developments should facilitate the application of TVI technology to imaging patients with ischaemic heart disease.

From the cardiac apex, the impact of translational motion of the heart within the thorax becomes much less important. As myocardial thickening is occurring predominantly at right angles to the direction of Doppler interrogation from the apex, this cannot reliably be measured with tissue velocity imaging. However, longitudinal shortening velocities do appear to be reliably measured from the apical window. Normal myocardial motion is complex, with three separate components: radial contraction, longitudinal shortening, and rotation. At present, no echo technique can simultaneously assess all three components and there is debate about the relative importance of each. There is now little doubt, however, that longitudinal shortening is an integral part of global contractile function, and it is thought to play as important a role as radial thickening in contributing to left ventricular (LV) ejection fraction.

The potential for quantification of longitudinal shortening as a simple estimate of overall systolic function has recently been proposed. Tissue velocity imaging from the apex has the potential to provide equivalent and perhaps more extensive clinical information more readily and more reproducibly.

Longitudinal contraction and relaxation velocities of the LV myocardium are greatest in the basal segments and decrease progressively toward the apex, where in fact velocities may become reversed. This is consistent with previous observations that LV contraction occurs towards a central point situated two thirds along a long axis line from the level of the mitral annulus to the apex. The normal longitudinal myocardial velocity profile when imaged from the apex consists of a systolic phase directed toward the transducer, and a diastolic phase directed away. The systolic phase consists of an early peak coinciding with isovolumic contraction followed by a second, larger peak coinciding with descent of the cardiac base. The normal diastolic phase consists of two distinct peaks, the first early (E) and the second late (A) in diastole. These correspond respectively to early and late LV diastolic relaxation, and thus to early and late ventricular filling, resembling in relative magnitude the E and A waves of the mitral inflow velocity profile measured by pulsed wave Doppler (fig 1).

Normal values for longitudinal shortening have been reported as mean (SD) 7.5 (1.3) to 9.5 (1.4) cm/s for the anterior septum, and 10.3 (1.9) cm/s for the basal lateral wall. The highest systolic tissue velocities in the long axis are observed in the mitral annulus and the basal segments of the lateral, anterior, and posterior free walls, followed by those in the basal septum. Peak early diastolic velocities also vary between segments, with highest values in the posterior and lateral walls (16.5 (4.0) to 17.5 (2.9) cm/s) and lowest in the anterior septum (10.9 (2.4) to 12.9 (3.1) cm/s). With the exception of the anterior and posterior septum, diastolic lengthening velocities are significantly higher in the basal segments than the mid wall segments. The ratio of early diastolic to late diastolic velocities (E/A) in normal subjects is similar to the conventional E/A ratio of mitral inflow velocities. Studies aiming to determine normal values and including both male and female subjects have been limited by relatively small numbers. Although there is little evidence thus far of a significant difference in velocities between the sexes, increasing age has been associated with a gradual fall in myocardial velocities both in systole and diastole.

TVI has also demonstrated differences within myocardial segments. M mode studies, obtained from the parasternal window, in a limited number of segments, have shown an intramyocardial velocity gradient with a gradually increasing velocity from the subepicardial to the subendocardial region. When imaging from the apex, there is also a quantifiable velocity gradient between two points separated in the longitudinal axis but lying within the same segment, a feature which may lend itself to assessment of regional contractile function. Recognition of these intramyocardial velocity gradients has led to the suggestion that persistence could represent viability in an otherwise akinetic segment; a hypothesis which requires further confirmatory work.

**Systolic function**

One of the first clinical applications of TVI has been as a method of assessing left ventricular systolic function. Assessment of this parameter remains the commonest indication for echocardiography, but standard methods are often limited by technical difficulties, inaccuracy and poor reproducibility. The advantages of a Doppler based technique as a means of acquiring data make this an attractive alternative to standard 2D imaging, and there is already evidence that TVI applications offer new and easy methods of assessing both global and regional LV systolic function.
GLOBAL LV FUNCTION

Descent of the cardiac base towards the ventricular apex is a feature of normal ventricular systolic function. This has been quantified by measuring the displacement distance of the mitral annulus by M mode echocardiography and correlates with left ventricular ejection fraction.\(^2\)\(^5\)\(^7\) It follows that the rate of mitral annular displacement may also reflect LV systolic function and studies using TVI to measure mitral annular displacement velocity (MADV) confirm this.\(^3\)\(^2\)\(^6\) Measurement is made at several sites—for example, lateral, septal, inferior, and anterior aspects of the annulus—to derive an average figure for MADV, as an adjacent regional wall motion abnormality may affect localised values and a mean velocity is more likely to reflect global LV function. Measurement of MADV is relatively easy as, even in difficult echo subjects, tissue velocity images of the mitral annular region can almost always be obtained from the apical window. On the 2D tissue Doppler map, the sample volume for PWTDs is placed over the mitral annulus and TVI software then allows real time readout of the velocity changes of that region with time. Alternatively, a cine loop can be stored digitally for offline analysis, where computer software allows reconstruction of velocity profiles in graphical form from any point on the tissue Doppler map.

REGIONAL LV FUNCTION

Positioning a sample volume within the myocardium equidistant from endocardial and epicardial borders allows measurement of segmental myocardial velocities, both at rest and with pharmacological stress. Regional velocities can be measured both in real time using PWTDs, or can be analysed offline by storing a digital cine loop. Modern systems capable of high frame rates, and thus high temporal resolution, ensure that velocity profiles derived from a stored cine loop closely reflect those obtained by PWTDs. This allows rapid acquisition of sufficient data for analysis of multiple regions simply by acquiring one cine loop, obviating the need for acquisition of multiple regional samples by PWTDs.

The colour coded 2D tissue velocity map also lends itself to the assessment of regional function, as deviations from the normal pattern of LV motion and colour changes throughout the cardiac cycle are often readily appreciated, providing immediate online appreciation of cardiac asynchrony. Areas of akinesia after myocardial infarction, for example, are shown by darker hues of colour representing low velocity of motion (fig 3), and dyskinetic segments may be seen to exhibit colours opposite to those of adjacent normal segments, particularly during systole.

Analysis of the intramyocardial velocity gradients also aids detection of regional dysfunction. M mode studies reveal reduction or loss of the subendocardial–subepicardial gradient during ischaemia or infarction, though the restriction of this technique to only a few myocardial segments reduces its utility as a practical tool.\(^4\)\(^2\) In contrast, the ability to measure

![Figure 3](http://heart.bmj.com/)

**Figure 3** Tissue velocity map superimposed on a 2D parasternal short axis view from a patient with recent antero-septal myocardial infarction. Myocardial velocity profiles are shown from the mid-septum (green marker and line) and mid-posterior wall (yellow marker and line). The peak systolic contraction velocity of the posterior wall is within the normal range (yellow S2 = 2.4 cm/s) whereas the corresponding velocity from the septum is minimal (green S2 = −0.6 cm/s), indicating akinesia. Diastolic velocities in the akinetic septal segment are similarly reduced. SI, myocardial velocity during isovolumic contraction; S2, peak systolic myocardial velocity; E\(_m\), peak early diastolic myocardial velocity; A\(_m\), late diastolic myocardial velocity associated with atrial contraction.
Figure 4  Myocardial velocity profiles obtained from the basal inferior segment (apical 2 chamber view) recorded at rest (panel A) and after full dose dobutamine stress (panel B). The systolic profile shows fusion of S1 and S2 (which often occurs, especially with rapid heart rates) and demonstrates an increase in peak systolic velocity, S2, from 8.0 cm/s to 13.0 cm/s in a subject with normal response. In this example the diastolic velocities E, and A, are also seen to increase with dobutamine stress (from \(-5.5\) to \(-7.1\) and from \(-12.8\) to \(-15.8\) cm/s respectively), the significance of which remains to be established. S1, myocardial velocity during isovolumic contraction; S2, peak systolic myocardial velocity; E, peak early diastolic myocardial velocity; A, late diastolic myocardial velocity associated with atrial contraction.

longitudinal velocity gradients between two points within the same segment, using the new datasets known as strain and strain rate imaging, appears to overcome this limitation and has the advantage of being quantitative rather than qualitative.22

LIMITATIONS OF ASSESSING SYSTOLIC FUNCTION BY TVI
The systolic descent of the cardiac base is a consequence of shortening of the ventricular long axis and predominantly reflects contraction of the longitudinally orientated subendocardial muscle fibres.23 Conversely, the radially orientated fibres are predominantly responsible for radial contraction and endocardial motion and may not be adequately assessed by measures of base descent.45 Hence MADV may only partly reflect global function, potentially limiting its value as a measure of overall LV systolic function.

When assessing regional function, the measured velocity of an individual myocardial segment may be influenced by the motion of adjacent muscle—for example, an akinetic basal segment interrogated from the apical window may yield near normal values if influenced by a hyperkinetic mid-segment, even if not contributing actively to longitudinal shortening itself. Furthermore, the velocity of a segment derived using PWTDS represents only that component occurring towards the transducer, which may include translational and rotational components. This is particularly problematic when using the parasternal window due to anterior cardiac motion in systole, and can be minimised by taking measurements from the apex, which is relatively fixed, from where contraction and relaxation in the axial plane can be measured without the need for angle correction. Thus, velocities measured relative to the transducer must be interpreted with caution when trying to assess the systolic function of individual segments.

Stress echocardiography
Normal subjects show an increase in peak systolic velocity with exercise or dobutamine stress (fig 4), with an attenuated response in patients with impaired coronary flow reserve.46–48 TVI provides an alternative to conventional semiquantitative methods for the diagnosis and assessment of ischaemic heart disease by stress echocardiography. Although limited by many of the drawbacks which apply to its use at rest (angle dependency, heart translational motion, and the influence of adjacent myocardial segments on velocities measured in the segment of interest) it does appear to introduce greater objectivity in regional quantification and thus improve reproducibility, overcoming some of the limitations of conventional stress echocardiography.49 50

Initial work involved use of PWTDS, requiring real time data acquisition from multiple segments during a stress echo study, thus limiting its practical utility.46–47 However, the development of new ultrasound systems with the ability to acquire real time colour Doppler images for subsequent offline analysis, with sufficiently high temporal resolution to ensure accurate measurement of peak velocity, has made the application of TVI methods to stress echocardiography a practical reality.44 51

Early attention focused on measurement of peak systolic velocity as the parameter likely to be most useful in assessing regional LV systolic function with stress, but more recently other parameters such as the time to peak systolic velocity and the velocity time integral have been proposed as equally important measures.49 The multicentre, multinational MYDISE study (MYocardial Doppler In Stress Echocardiography) is an ongoing study assessing the feasibility and reproducibility of these TVI methods. Early results indicate that TVI does enhance both objectivity and reproducibility of stress echocardiography.52

Furthermore, the susceptibility of subendocardial longitudinal muscle fibres to early ischaemia suggests that longitudinal shortening velocity measured from the apical window should be a sensitive marker of impaired contractile function—perhaps more so than recognition of reduced endocardial excursion, on which conventional assessment by stress echo traditionally depends.52 Longitudinal shortening velocities measured from the apex are reported to show greater response to dobutamine and to be more reproducible than radial contraction velocities measured from the parasternal window.46 This enhancement of stress echocardiographic methods by use of TVI applications may be improved still further if developments in strain rate imaging allow incorporation of this technique in future.
Diastolic function

TVI holds great promise for the assessment of LV diastolic function. Whereas conventional Doppler measures of LV filling derived from mitral inflow velocities reflect only global diastolic function, TVI offers the ability to measure regional diastolic function by echocardiography for the first time. Spectral PWTDs allows quantification of diastolic myocardial velocities from a small sample volume placed in a region of interest in the same way as for systolic velocities, and this has enabled characterisation of the normal pattern of early and late diastolic velocity peaks ($E_1$ and $A_2$, respectively), with highest values at the mitral annulus, decreasing progressively towards the apex.$^8$

Relaxation velocity of the mitral annulus, averaged from four different sites, reflects global LV diastolic function and correlates (in normal subjects) with conventional measures derived from LV filling patterns and the time constant of isovolumic relaxation ($\tau$).$^{53,54}$ Regional myocardial diastolic velocities, measured as segmental longitudinal relaxation velocities from the apical window, are relatively uninfluenced by translational motion although—as for regional systolic function—care must be taken when interpreting localised values in view of the possible influence of adjacent segments. A conventional measure of global diastolic function, E/A ratio, has been shown to fall with an increase in the number of myocardial segments exhibiting a reduction in the tissue velocity $E_1/A_2$ ratio, lending support to the hypothesis that TVI offers a useful measure of regional diastolic function.$^9$

Moreover, there is evidence that diastolic tissue velocities are less influenced by the changes in preload which commonly compensate for diastolic dysfunction and which confound assessment by standard measures based on LV filling patterns.$^{55,56}$

With normal LV filling, the peak early diastolic myocardial velocity ($E_1$) measured in the long axis plane is > 10 cm/s in the young, falling to > 8 cm/s with aging. Both early and intermediate stages of diastolic dysfunction (the delayed relaxation and pseudonormal phases respectively), are characterised by a reduction in $E_1$ velocity to < 8 cm/s, falling further still in the most severe, or restrictive, stage.$^{54}$ Hence myocardial velocities are persistently reduced even in those stages of diastolic dysfunction characterised by increased preload compensation and thus normal or high mitral inflow velocities. This lack of concordance between the two parameters in the pseudonormal and restrictive stages means that myocardial velocity measurement can allow better recognition of these patterns, which are otherwise often difficult to distinguish from normal.

This has practical value in differentiating restrictive cardiomyopathy from pure constrictive physiology, a difficult distinction when relying on standard 2D and Doppler criteria alone. Garcia et al have reported that a tissue Doppler $E_1$ velocity of 8 cm/s reliably separates restrictive cardiomyopathy (characterised by a low $E_1$ value in the context of “pseudonormalised” LV filling with high $E$ velocity and a normal $E/A$ ratio) from constrictive physiology (normal/high $E_1$ velocity if systolic function remains normal).$^{36}$

Mitrail inflow velocity patterns also differ from diastolic myocardial velocity patterns in some groups with ventricular hypertrophy, enabling differentiation between the physiological hypertrophy of athletes and pathological hypertrophy of hypertrophic cardiomyopathy (HCM), for example. Palka et al reported that myocardial velocity gradient (MVG), measured across the posterior wall of the LV in the parasternal short axis view, discriminated between patients with HCM and those with hypertensive LV hypertrophy (LVH) or athletic hearts.$^{57}$ There was no significant difference in the conventional Doppler indices of LV filling to differentiate the HCM and the athletic LVH groups, suggesting that myocardial $E_1$ velocities are less influenced by alterations in preload, which can confound assessment of diastolic function by conventional LV filling indices. Further evidence of the relative preload independence of $E_1$ velocity was presented by Oki et al, who reported that the time constant of isovolumic LV relaxation ($\tau$) correlated well with $E_1$ velocities rather than filling pressure.$^{58}$

Sohn et al also demonstrated that manipulation of LV filling dynamics produced alterations in the LV Doppler inflow patterns but with no significant changes in the $E_1$ velocities.$^{59}$ $E_1$ velocity thus has a unimodal distribution, Doppler myocardial velocities showing a steady decline with increasing diastolic dysfunction, in contrast to the bimodal distribution of mitral inflow velocity $E$ which occurs due to preload compensation in the more severe stages. Again, these observations are consistent with relative independence of $E_1$, in relation to compensatory alterations in loading conditions.$^{55,56}$

Assessment of diastolic function by conventional Doppler echocardiography relies heavily on study of the mitral inflow velocity profile, but this is of limited value in the presence of atrial fibrillation or other arrhythmia. As TVI enables the study of diastolic function by direct measurement of myocardial motion, it does not suffer this limitation of conventional methodology.$^{56}$

Reversible myocardial ischaemia affects diastolic function earlier than systolic function suggesting that regional diastolic function measurement could aid its assessment.$^{50}$ PWTDs has enabled rapid and accurate measurement of both myocardial diastolic velocities and regional isovolumic relaxation times, which have been used to differentiate between normally perfused and ischaemic segments.$^{7,56,61}$ Myocardial ischaemia is characterised by a reduction in $E_1$ velocity and consequently also in $E_1/A_2$ ratio.$^{49}$ The $A_2$ velocity however has been reported to remain unchanged, which may be because whereas early diastolic changes reflect an active, energy requiring process, late diastolic filling merely reflects passive stretch after atrial contraction.$^{38,61}$

Full clinical utility of these
observations requires further work but, particularly when added to a stress study, could extend this technique still further.

**Future applications—myocardial strain and strain rate imaging**

The recent development of myocardial strain rate imaging also promises to improve the quantification of regional myocardial ischaemia, overcoming some of the limitations of conventional TVI. Doppler measurement of myocardial strain has resulted from the further development of TVI principles and allows quantification of myocardial shortening within an individual segment (fig 5). This represents a more logical method of assessing regional contractile function as these datasets are not influenced by the function of adjacent myocardial segments and are less dependent upon the direction of shortening in relation to the transducer. Strain rate is a measure of the velocity of deformation of myocardium, and is given by the formula \((V_2-V_1)/d\), where \(V_1\) and \(V_2\) are velocities of myocardial shortening at two points separated by distance \(d\); this gives the difference in tissue velocity per unit length.\(^{22}\)

Integration of this parameter with respect to time gives myocardial strain, a measure of the per cent compression of myocardium during systole.\(^{22}\) This is also given by the formula \((L_1-L_2)/L_1\), where \(L_1\) is the distance between two points at end diastole and \(L_2\) is the distance between the same two points at end systole; this parameter cannot otherwise be measured directly by 2D echo without placement of myocardial markers. This approach will clearly allow identification of akinetic segments, where strain rate and strain will be zero, and may also allow identification of inducible ischaemia and viable myocardium.\(^{52-65}\)

The lower part of the panel shows a graphical representation of changes in strain rate during the cardiac cycle; this can be seen to be approximately a mirror image of the changes in myocardial velocity. The hatched area during ventricular systole represents the integral of myocardial strain rate which gives the myocardial strain.

**Summary**

Tissue velocity imaging is another important development in the field of cardiac ultrasound and promises a number of applications to further extend the unique diagnostic role of echocardiography. Accurate quantification of global and particularly regional systolic LV function is closer than ever before, and the promise to improve the reproducibility and diagnostic value of stress echocardiography looks set to be realised, though full results from MYDISE are eagerly awaited.

Tissue velocity imaging has enabled echocardiographic assessment of regional diastolic function which appears to be less limited by the compensatory changes in loading conditions that confound measurement of diastolic function by conventional echocardiographic methods. Tissue velocity imaging thus offers a practical clinical tool to differentiate physiological from pathological LV hypertrophy, and restrictive from constrictive physiology by refreshingly simple means.

The development of the new datasets strain and strain rate promise to further enhance the value of echocardiography by overcoming some of the limitations of early TVI techniques; this should allow not only superior assessment of regional contractile function at rest and with stress, but also reliable identification of viable myocardium.

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