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Tissue Doppler in Ischemic Heart Disease

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1. Introduction

Tissue Doppler Echocardiography was introduced in the 1960s (Yoshida et al., 1961), and enabled the quantitative assessment of myocardial motion and deformation. The wide use of tissue Doppler as a research tool halted, however, until the early 1990s (Hatle & Sutherland, 2000). Tissue Doppler is now available for high frame rates, wide sector angles and in combination with 2-dimensionlal data acquisition. Although widely used in cardiovascular research, the clinical use is limited, probably due to the time consumption associated with special imaging protocols and tedious post processing. Nevertheless, tissue Doppler echocardiography has contributed to most of the available knowledge on the pathophysiology involved in myocardial contraction deficiency.

The experienced cardiologist can easily identify large myocardial infarcts by visual analysis of echocardiograms, but identification of a small MI may be challenging due to the modest changes in tissue properties. In ischemic tissue, the contractility is reduced and reduced deformation and deformation rate is observed. Due to differences in contractility among ischemic and adjacent healthy myocardium, the ischemic myocardium has a characteristic deformation pattern with stretch in early systole, reduced systolic shortening, and a delayed (postsystolic) shortening when the ventricular pressure decays. This early stretch and post systolic shortening pattern has been described both experimentally (Edvardsen, T. et al., 2002; Skulstad et al., 2002) and clinically (Gjesdal et al., 2008; Jamal et al., 1999; Voigt et al., 2004).

2. Deformation indices

Myocardial function was traditionally assessed by tissue Doppler by the measurement of myocardial velocity and displacement indices in directions parallel to the ultrasound beam direction. These indices allowed quantitative assessment of myocardial function in the longitudinal direction, but could not differentiate velocities caused by deformation in the myocardium from velocities caused by displacement and tethering from adjacent structures.

The deforming heart can be assessed by evaluation of tissue velocities or displacement, deformation (strain) or rate of deformation (strain rate).

The Doppler shift utilized for deformational assessment in tissue Doppler echocardiography is a measure directly correlated to tissue velocity, and strain or strain rate measurements are automatically derived by integration or derivation of the spatial velocity distribution (Figure
During contraction, the apex is relatively stationary and the strain is relatively equally distributed in the different left ventricular compartments. There is therefore a natural gradient of velocity and displacement in the heart, with higher values observed towards the basal part of the heart. Tethering (pulling) by healthy adjacent myocardium, and translational cardiac motion can thus influence the velocities and displacements of diseased myocardium. When these indices are assessed, the position of the region of interest must therefore be evaluated together with the velocity information (Skulstad et al., 2004).

![Diagram of Displacement and Velocity](image)

**Fig. 1. Mathematical relation between deformation indices**

In an attempt to establish a measure that was more specific for myocardial properties the principle of calculating myocardial strain rate from echocardiographic data was introduced (Quinones et al., 1974). This is analogous to the earlier approach to calculate strain rate or “normalized velocities” from apex cardiographic tracings, and the concept of myocardial strain was defined as fractional tissue deformation in response to applied force or stress (Mirsky et al., 1972). In that context, strain represents fractional change of tissue length and is expressed in a dimensionless unit as percent shortening or lengthening (Lagrangian formula). In an effort to improve the ability of TDI to measure regional function, TDI-derived real time strain rate was introduced (Heimdal et al., 1998). Strains are more uniformly distributed within the LV myocardium than tissue velocities, and the assessment of myocardial strain by TDI thus simplifies the analysis of regional contractile function by providing an objective parameter of myocardial deformation. Strain is a relatively new index of myocardial function that describes myocardial deformation as the relative change in myocardial segment length over time (D’hooge, J. et al., 2002), and provides information on segmental or global myocardial deformation (Gjesdal et al., 2007). Strain (deformation) or strain rate (rate of deformation) can be assessed in patients by echocardiography using tissue Doppler or speckle tracking echocardiography (Dandel & Hetzer, 2009) or with tagged MRI (Edvardsen & Rosen, 2005), and in experimental studies by sonomicrometry (Urheim et al., 2000). The indices have proven superior to established indices of myocardial function to assess myocardial infarct size (Becker et al., 2006; Chan et al., 2006; Gjesdal et al., 2008; Vartdal et al., 2007), hypertrophic cardiomyopathy (Serri et al., 2006), and in metabolic syndrome (Gong et al., 2009), adds incremental value in predicting outcome in cardiovascular
disease (Ingul, C. B. et al., 2007), and predicts arrhythmic events after myocardial infarct (Haugaa et al., 2009).

Strain and strain rate (SR) were developed as clinical indices of regional myocardial deformation (Edvardsen et al., 2001; Gotte et al., 2001; Mirsky & Parmley, 1973; Rademakers et al., 1994; Zhang et al., 2005) and have been introduced and validated using tagged MRI and sonomicrometry (Derumeaux et al., 2001; Edvardsen, T. et al., 2002; Urheim et al., 2000). Strain is defined as tissue elongation relative to length; usually the length at end diastole (Lagrangian strain) but instantaneous length is also used (Eulerian strain). A positive strain value refers to elongation, whereas a negative strain value describes shortening. Strain and strain rate are related through temporal derivation or integration, respectively. Therefore, negative systolic strain and strain rate values describe a normal contracting myocardial segment. Three main systolic deformation patterns form perpendicular axes in the heart's internal coordinate system (D’hooge et al., 2000); longitudinal shortening, circumferential shortening and radial thickening (Figure 2). In a direct comparison between strain, displacement and ejection velocity, strain by tissue Doppler was found to be superior to describe regional myocardial function, both in an animal model, and in humans (Skulstad et al., 2006).

Fig. 2. The heart’s coordinate system. Longitudinal shortening (Ls), Circumferential shortening (Cs) and Radial Thickening (Rt) is displayed.

Doppler derived measures are angle dependent, and the assessment of global function is limited by angulations of apical segmental myocardium relative to the ultrasound beam direction. To overcome this problem, assessment of mitral plane velocities or displacement has been proposed as an index of global LV function (Alam et al., 1990; Alam, M. et al., 1992; Hoglund et al., 1989; Simonson & Schiller, 1989). Due to tethering from the mid-ventricular and apical segments, the displacement at the base of the left ventricle reflects the average deformation of the LV walls. When opposing walls are evaluated during the same heartbeat, the potential misjudgment caused by apical rocking is reduced to a minimum. Mitral annulus displacement and velocity analyses by TDI has been validated in the clinical setting, is well established and widely available. The method requires dedicated imaging protocols and is angle dependant, but might be more robust compared to speckle tracking imaging.
when the acoustic window is poor. Mitral annulus displacement has proven to be robust and reproducible (Hayashi et al., 2006), and has demonstrated good ability to predict prognosis following myocardial infarct (Brand et al., 2002). Mitral annulus displacement can also be normalized for end diastolic LV length (normalized Mitral annulus Displacement), an index which corresponds to global strain (Gjesdal et al., 2009).

To eliminate the problem of angle-dependency in Doppler-derived deformation analyses, strain measurement based on two-dimensional speckle-tracking echocardiography (2D-STE) has recently been developed (Amundsen et al., 2006; Cho et al., 2006; D’hooge, J. et al., 2002; Helle-Valle et al., 2005; Leitman et al., 2004). Natural acoustic markers (speckles) visualized by gray scale imaging form patterns within myocardial tissue. Dedicated software identifies the speckle patterns, and myocardial deformation assessed based on deformational changes on a frame-to-frame basis. Strain is calculated for each LV-segment as the average relative deformation in circumferential, longitudinal or radial directions (Becker et al., 2006; Chan et al., 2006; Vartdal et al., 2007) and the method furthermore enables assessment of LV-rotation and twist (Helle-Valle et al., 2005). The method is semi-automatic and relatively independent of angle. Spatial resolution is nearly constant with depth in the direction of the ultrasound beam, while the spatial resolution orthogonal to the beam direction is constant with depth with a linear array transducer, and decreases slightly with depth when a sector or phased array transducer is used. Since speckle tracking echocardiography is a developing methodology for assessment of the same indices that are assessed by tissue Doppler echocardiography, this will also be covered briefly. The implementation of TDI in daily clinical work has been relatively slow and most echocardiographic laboratories do not apply TDI as a routine diagnostic method.

3. Infarct assessment

Direct assessment of myocardial infarct by histopathology is the gold standard for infarct sizing, and is an option in post-mortem studies, in animal studies, and in studies on the explanted heart (Kim et al., 1999; Medrano et al., 1996). Biochemical infarct sizing is based on the correlation between the amount of damaged myocardium and release to the blood pool of specific markers of cardiac necrosis. Peak values correlate to infarct size, but the accuracy of the methods depends on correct timing of the blood sampling in relation to the ischemic event (Gibbons et al., 2004). Visualization of MI by imaging techniques is based on differences in tissue properties among normal and infarcted tissue. In CE-MRI, the infarcted myocardium is highlighted due to retention of contrast medium in the infarcted tissue, while positron emission tomography (PET) and single photon emission computed tomography (SPECT), are based on visualization of viable non-infarcted myocardium due to preserved glucose metabolism and retention of radioactive tracers in viable myocytes, respectively (Gibbons et al., 2004). In echocardiography and in cine- and tagged MRI imaging, changes in cardiac motion pattern, deformation, or changes in LV pumping performance form the basis of infarct sizing.

Myocardial contraction is severely reduced within scars, and systolic deformation of the left ventricle therefore decreases with increasing myocardial scar load (Chan et al., 2006; Hoglund et al., 1989; Skulstad et al., 2006; Vartdal et al., 2007). Assessment of myocardial deformation at a global or segmental level thus represents alternatives to the direct assessment of myocardial scar load by CE-MRI (Figure 3). Echocardiographic scanners are
less expensive, widely distributed, and are mastered by most cardiologists, and enables myocardial deformation assessment by a variety of methods. Evaluation of LV-deformation has traditionally been performed by assessment of the relative volume reduction during systole (Left ventricular ejection fraction, LVEF), by visual assessment of the wall thickening in individual LV-segments during systole (Wall motion score, WMS) (Lang et al., 2005), or by assessment of the longitudinal LV-shortening or rate of shortening (mitral annulus displacement or velocity) (Simonson & Schiller, 1989).

Echocardiographic assessment of LVEF is easily available and feasible, but does not provide information on segmental LV-function. Evaluation of regional function by analyses of endocardial motion or local wall thinning and thickening characteristics is user dependent and requires well-trained personnel. Measurement of longitudinal LV-deformation by mitral annulus (MA) displacement or velocity can be performed by tissue Doppler imaging (TDI), pulsed TDI or M-mode echocardiography (Hayashi et al., 2006). Assessment is relatively easy, but the reference values differ among the methods. Furthermore, the methods do not provide information on segmental deformation, and regional deformation may be influenced by tethering or apical rocking. MA-displacement, however, predicts future events in patients with myocardial infarction, heart failure or hypertensive heart disease (Ballo et al., 2008; Hillenbrand et al., 2000; Willenheimer, R. et al., 1997).

Fig. 3. Bulls eye plot of peak systolic longitudinal strain by speckle tracking echocardiography (left) and gadolinium contrast MRI (right). The center of the plot represents apex, and the rim represents the basal LV segments.

Low systolic myocardial velocities in ventricles with myocardial damage, hypertrophy and cardiomyopathy have been demonstrated by several authors (Bach et al., 1996; Gorcsan, III et al., 1996; Miyatake et al., 1995; Uematsu et al., 1995). Mitral annular and myocardial longitudinal velocity measurements have been used in several studies to characterize systole in normal individuals and in different heart diseases (Fukuda et al., 1998; Gulati et al., 1996; Oki et al., 1999; Pai & Gill, 1998; Wilkenshoff et al., 1998). Generally, systolic longitudinal velocity was reduced in most heart diseases. This longitudinal approach has also been useful to describe the diastolic function in normal subjects and patients with LV
hypertrophy showing a decrease in VE with age and in LV hypertrophy (Rodriguez et al., 1996). Several conditions are thus responsible for decreased myocardial systolic and diastolic velocities and myocardial single finding of reduced velocity. Based on the available literature, it is likely that focus on the highest systolic velocity or deformation is not sufficient to characterize ischemia from other diseases affecting the contraction of the heart.

4. Validation studies

Tissue Doppler derived strain and strain rate have been validated using tagged MRI and sonomicrometry in humans and in animal studies (Derumeaux et al., 2001; Edvardsen et al., 2002; Urheim et al., 2000). Myocardial longitudinal strain was assessed as the time integral of regional Doppler velocity gradients in a dog model, and compared with strain derived from sonomicrometry crystals placed near the LV apex and base, respectively (Derumeaux et al., 2001; Edvardsen, T. et al., 2002; Urheim et al., 2000). Comparisons during baseline, apical ischemia and preload alterations demonstrated good correlations (r = 0.92, p<0.01). In human studies longitudinal myocardial Doppler velocities have been shown to decrease progressively from base to apex, while myocardial strain rates and strains are uniformly distributed among all segments (Edvardsen, T. et al., 2002). Comparisons between myocardial longitudinal strains by SDE and tagged-MRI showed excellent correlations (r=0.89 and r=0.96, for longitudinal and radial strains respectively (p<0.001), in healthy individuals, infarct patients, and during stress echocardiography.

5. Acute ischemic heart disease

5.1 Deformation characteristics

During acute ischemia, the contractility of the ischemic myocardium is reduced. Postsystolic shortening was identified as a sign of viability in a dog model of ischemic heart disease (Takayama et al., 1996), and tissue Doppler enabled the characterization of regional wall motion disturbances during ischemia and reperfusion (Derumeaux et al., 1998). Also in humans, abnormal postsystolic contraction was observed during ischemia (Jamal et al., 1999). Deformation assessment by strain and strain rate demonstrated better ability to evaluate ischemic myocardium compared to velocity based indices in animal models (Hashimoto et al., 2003) and human studies (Edvardsen, T. et al., 2002; Jamal et al., 2002; Zhang et al., 2005), and provide information on small changes in function over time (Ingul et al., 2005).

In a study during and after angioplasty in humans, three major characteristics were identified for the systolic velocity pattern of ischemic myocardium (Edvardsen et al., 2000; Edvardsen et al., 2001). When comparing the ischemic regions of LV with the non-ischemic regions, the ischemic region was recognized by early systolic stretch and reduced peak systolic shortening, followed by a postsystolic shortening when the LV pressure decays during isovolumetric relaxation and early diastole (Figure 4). When the ischemia becomes more severe, the segment gradually becomes passive, and the deformation depends on the passive elastic properties of the myocardial tissue (Skulstad et al., 2002).
Fig. 4. Strain rate and strain from typical normal (grey) and ischemic (black) myocardium. Ischemic myocardium is characterized by an early peak positive strain (PPS), and peak systolic strain (PSS) is typically lower than end systolic strain (ESS). A post systolic shortening is often seen, and peak strain (PS) therefore occurs after end systole.

During ischemia, adjacent myocardium also display altered deformation characteristics. This might be caused by stunning or transient ischemia, but also by alterations in local loading conditions. Remote myocardium generally has been thought to increase the contraction in a compensatory manner to accomplish the demands to cardiac output. This mechanism could be caused by neuro-hormonal activation, but also but the Frank-Starling mechanism secondary to the increased preload of the ischemic heart. Recently, strain by Doppler and speckle tracking was demonstrated to display similar ability to separate among levels of transmurality in acute ST elevation myocardial infarction, but that reproducibility was somewhat better for speckle tracking derived strain (Sjoli et al., 2009). Post-systolic shortening has also been demonstrated in approximately one third of segments in healthy individuals, and indexation of post-systolic shortening relative to peak systolic shortening has therefore been suggested (Voigt, J. U. et al., 2003). Systolic strain by vector velocity imaging is also found to correlate with infarct in the acute phase, and peak systolic strain was a better predictor of infarct than the post systolic shortening index (Jurcut et al., 2008).

5.2 Identification of area at risk

Strain imaging by tissue Doppler echocardiography shortly after PCI for acute myocardial infarct correlated strongly with global infarct size assessed by delayed enhancement MRI 9 months later (Vartdal et al., 2007). Similar results were also found for speckle tracking echocardiography (Sjoli et al., 2009). Moreover, the strain value correlated with infarct transmurality level, suggesting that this index might be useful for prognostication.
5.3 Non ST elevation infarction
Global strain by speckle tracking deteriorates in non ST-elevation patients while waiting for angiography, and the patients with occlusion did not recover by revascularization, thus suggesting a potential benefit from a more aggressive strategy in these patients (Grenne et al., 2010). Strain measured immediately before revascularization was found to correlate significantly with infarct size assessed by delayed enhancement MRI also in non ST-elevation infarcts (Eek et al., 2010), and presence of reduced function in 4 or more adjacent segments was shown to predict coronary occlusion with high accuracy (Eek et al., 2010).

6. Chronic ischemic heart disease
6.1 Deformation characteristics
In chronic ischemic heart disease, the infarcted area has transformed to scar tissue with increased resistance to passive stretch. As a consequence, early systolic stretch is a less common finding. There is often, however, ischemic areas and areas with subendocardial infarct present adjacent to the infarcted areas, and early stretch can be identified in these areas. In the MESA study, reduced myocardial deformation was found in healthy individuals with increased coronary artery calcification (Edvardsen et al., 2006).

6.2 Estimation of infarct size and level of transmurality
In chronic myocardial infarct, strain values assessed by tissue Doppler or speckle tracking echocardiography are reduced with increasing level of transmural infarct distribution (Becker et al., 2006; Chan et al., 2006; Gjesdal et al., 2007; Sachdev et al., 2006; Weidemann et al., 2003), both at the segmental, at the global, and at a territorial level (Gjesdal et al., 2008). This is interesting, since segments with transmural infarct defined as more than 50% of the segment mass is less likely to benefit from revascularization.

6.3 Risk stratification for arrhythmias
Recently, post myocardial infarct patients with indication for implantable cardioverter-defibrillator (ICD) were followed for arrhythmias requiring appropriate ICD therapy (Haugaa et al., 2009). In the patients with arrhythmia the dispersion of peak strain, calculated as standard deviation of time to maximal strain for all segments, was increased compared to patients who did not require therapy. Moreover, the global strain value was better in the patients without arrhythmia compared to the ones who received ICD therapy.

6.4 Stress testing
Tissue Doppler echocardiography has been used to quantify segmental function during exercise testing by velocity imaging (Pasquet et al., 1999), and has been demonstrated to predict functional recovery from revascularization (Schneider et al., 2005). When comparing
deformation imaging and velocity imaging for the detection of regional inducible ischemia during dobutamine stress echocardiography, deformation indices demonstrated superior ability to detect ischemia compared to velocity based indices (Voigt et al., 2004). Moreover, in another study, postsystolic shortening demonstrated the best ability to identify stress induced ischemia after normalization to peak deformation, since postsystolic shortening also occurred to some extent in healthy segments (Voigt, J. U. et al., 2003). Strain and strain rate had been found superior to wall motion score to identify angiographic significant stenoses (Ingul, C. B. et al., 2007), and also to predict outcomes in patients admitted to stress echocardiography for suspected coronary heart disease (Ingul, C. B. et al., 2007).

7. Prognosis
Assessment of strain early following myocardial infarction predicts remodeling defined as more than 15% increase in end diastolic volume (Park et al., 2008). Global LV function assessed by tissue Doppler measurements of the mitral annulus displacement predicts events in chronic infarction (Brand et al., 2002) and congestive heart failure (Willenheimer, R. B. et al., 1997). The addition of strain rate during stress echocardiography added prognostic value over traditional risk factors and echocardiographic parameters (Stanton, T. et al., 2009), and an association between resting deformation indices and events was also demonstrated in people referred to echocardiography (Stanton, T. et al., 2009). Global strain by speckle tracking echocardiography was assessed within 48 hours of myocardial infarct in 659 patients, and found to be superior to LVEF and WMSI to predict mortality and clinical Events (Antoni et al., 2010).

8. Conclusions
All indices of myocardial function demonstrate reduced systolic deformation in infarcted myocardium. The deformation gradually reduces with increasing infarct size and transmural distribution. Ischemic tissue is characterized not only by reduced peak deformation, but also by an altered deformation pattern characterized by early systolic stretch and post systolic shortening.

9. References
Amundsen BH, Helle-Valle T, Edvardsen T, Torp H, Crosby J, Lyseggen E, Stoylen A, Ihlen H, Lima JA, Smiseth OA, Slordahl SA. Noninvasive myocardial strain measurement by speckle tracking echocardiography: validation against...
sonomicrometry and tagged magnetic resonance imaging. *J Am Coll Cardiol* 2006;47:789-93.


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Hashimoto I, Li X, Hejmadi BA, Jones M, Zetts AD, Sahn DJ. Myocardial strain rate is a superior method for evaluation of left ventricular subendocardial function compared with tissue Doppler imaging. *J Am Coll Cardiol* 2003;42:1574-83.


Lang RM, Bierig M, Devereux RB, Flachskampf FA, Foster E, Pellikka PA, Picard MH, Roman MJ, Seward J, Shanewise JS, Spencer KT, Sutton MS, Stewart WJ. Recommendations for chamber quantification: a report from the American Society of Echocardiography’s Guidelines and Standards Committee and the Chamber Quantification Writing Group, developed in conjunction with the European Association of Echocardiography, a branch of the European Society of Cardiology. J Am Soc Echocardiogr 2005;18:1440-63.


Since the introduction of Doppler Echocardiography, Nuclear Cardiology and Coronary CT imaging, clinicians and researchers have been searching for ways to improve their use of these important tools in both the diagnosis and treatment of heart disease. To keep up with cutting edge improvements in these fields, experts from around the world have come together in this book to provide the reader with the most up to date information to explain how, why and when these different non-invasive imaging tools should be used. This book will not only serve its reader well today but well into the future.

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