Current State of Three-Dimensional Myocardial Strain Estimation Using Echocardiography

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With the developments in ultrasound transducer technology and both hardware and software computing, real-time volumetric imaging has become widely available, accompanied by various methods of assessing three-dimensional (3D) myocardial strain, often referred to as 3D speckle-tracking echocardiographic methods. Indeed, these methods should provide cardiologists with a better view of regional myocardial mechanics, which might be important for diagnosis, prognosis, and therapy. However, currently available 3D speckle-tracking echocardiographic methods are based on different algorithms, which introduce substantial differences between them and make them not interchangeable with each other. Therefore, it is critical that each 3D speckle-tracking echocardiographic method is validated individually before being introduced into clinical practice. In this review, the authors discuss differences and similarities of the currently available 3D strain estimation approaches and provide an overview of the current status of their validation. (J Am Soc Echocardiogr 2012; 25: 589-601.)

Keywords: Echocardiography, Myocardial deformation, 3D strain, Validation, Heart

Ultrasonic myocardial deformation imaging offers the possibility of quantifying regional cardiac deformation noninvasively. It is widely used for the detection of myocardial ischemia and has been proposed as a tool to detect heart disease at its preclinical stage to differentiate among various hypertrophy etiologies, and to monitor therapy. Traditionally, cardiac motion and deformation analysis was limited to a visual inspection of the image sequence, in which each myocardial segment was assigned a wall motion score. However, this methodology depended considerably on the expertise of the interpreter and showed relatively low interobserver agreement.

For this purpose, myocardial strain (i.e., the relative lengthening or shortening of the myocardial segment expressed as a percentage of its initial length) and strain rate (i.e., the rate of this lengthening or shortening) imaging was introduced. For further details, the reader is referred to previous literature. At present, the assessment of one-dimensional strain with Doppler tissue imaging (DTI) is a well-established technique to regionally quantify myocardial function. This technique requires high frame rates, which may be difficult to obtain in dilated ventricles. However, it can be overcome by reducing the imaging sector width. Therefore, this should not be considered a true limitation of the technique. On the other hand, just as with other Doppler techniques, this technique is angle dependent, as only the strain component along the ultrasound beam can be measured. Although it allows the measurement of longitudinal strain (LS) in all left ventricular (LV) segments, radial strain (RS) and circumferential strain (CS) can only be measured in a limited number of LV segments.

Non-Doppler-based, two-dimensional (2D) strain estimation methods can partly overcome these limitations. Various echocardiographic techniques have been proposed, which use raw radiofrequency signals or apply block-matching techniques, often termed speckle-tracking (ST) techniques, on B-mode images. Several of these methods based on 2D ST have already been commercialized. However, six acquisitions of different parasternal and apical LV views are still required to obtain all strain components in all LV segments. This can be an issue in certain experimental and clinical situations, particularly when heart rate or LV loading conditions change rapidly.

With developments in ultrasound transducer technology and both hardware and software computing, systems capable of acquiring real-time volumetric LV data are now widely available. Most of the systems use electrocardiographic (ECG) gating to construct three-dimensional (3D) data from smaller subvolumes acquired at subsequent cardiac cycles. As such, reasonable spatial and temporal resolution of 3D data sets can be achieved. The ability to estimate true 3D myocardial motion and deformation using various 3D ST echocardiography (STE) approaches may provide cardiologists with a better view of regional myocardial mechanics, which may be important for diagnosis, prognosis, and therapy. These 3D approaches can measure all strain components in all LV segments from a single acquisition. Furthermore, they are angle independent, do not suffer from strain estimation errors associated with out-of-plane motion, and may in theory allow more precise calculations of LV twist and assessment of shear strain components.

However, tracking in three dimensions naturally poses some challenges. The increased field of view of volumetric images comes at the cost of both spatial and temporal resolution of the data set. In other words, the current volumetric data sets show a coarser speckle pattern and a higher speckle decorrelation between subsequent volumes. The latter effect becomes even more pronounced in data sets acquired at high heart rates (e.g., during stress echocardiography). Moreover,
because of the amount of data available in volumetric data sets, the computational load is also higher compared with 2D data sets. These aspects pose a challenging environment for myocardial motion and deformation imaging. Given the potential of 3D STE techniques and a multitude of publications that have recently appeared in medical journals, we felt that a comprehensive overview that would make this area more accessible to the reader was lacking. In this review, we thus discuss differences and similarities among the currently available 3D STE approaches and provide an overview of their first validation and clinical application studies.

THREE-DIMENSIONAL STRAIN ESTIMATION APPROACHES

General Work Flow
All currently available techniques follow the same subsequent steps to estimate 3D strain. This is illustrated in Figures 1 to 4.

In the initial step, key cardiac events are indicated: end-systole and end-diastole. This can be done automatically (e.g., on the basis of the electrocardiogram) or manually (e.g., by visually inspecting the cardiac cycle and indicating the frames with the smallest and largest volumes, respectively; Figure 1). Second, the region of interest for strain estimation is defined. Depending on the software, the endocardial border, sometimes in combination with the epicardial border, is delineated manually, semiautomatically, or automatically. Typically, this involves contouring the end-diastolic frame, although some software packages may require repeating the process for the end-systolic frame as well (Figure 2). Next, the left ventricle is subdivided into segments for segmental strain analysis, usually ranging from 16 to 18 segments depending on the segmentation model used. An anatomic landmark (e.g., the right ventricular insertion point) is used to indicate the orientation of the segments (Figure 3). Finally, the 3D LV region of interest is tracked throughout the cardiac cycle, and the deformation curves are estimated (Figure 4). The main differences between the currently available 3D STE algorithms are found in this final step. Their underlying principles are briefly discussed in the next sections.

Block Matching
The most common 3D STE approach is based on block matching, which can be considered a direct extension of 2D ST to three dimensions. This technique is based on the assumption of a stable local speckle pattern between subsequent volumes. Local tissue motion can then be extracted from the displacement of those speckle patterns from one frame to another (Figure 5). Because the motion estimates are performed independently from one another, they are usually noisy. An a posteriori regularization step (e.g., by spatially smoothing the initial motion estimates) is therefore often performed. Note that this approach also works on volumetric data acquired through ECG gating, given that no stitching artifacts appear in the volumetric data (otherwise, the assumption of a stable speckle pattern between subsequent frames is violated).

Elastic Registration
Elastic or nonrigid image registration techniques use image-warping techniques to estimate cardiac motion between subsequent volumes. In this method, one image is deformed in multiple steps to look as similar as possible to the next image in the image sequence. Once both images match optimally, interframe motion is known (Figure 6). As opposed to the block-matching approaches, the motion of the whole myocardium is thus estimated simultaneously. However, not all warping solutions are feasible or desired. Therefore, additional conditions are imposed during the motion estimation process (e.g., by enforcing the tissue motion to be smooth in space and/or time). For more details, the reader is referred to previously published articles discussing the technical aspects of elastic image registration. Currently, this method has been used mostly in academic circles and will be made commercially available.

Model-Based Approach
Models incorporating a priori knowledge can also be used to guide the 3D myocardial motion and strain estimation process. Different sources may be used as input to this model; for example, a biomechanical model can describe the deformation of myocardium, incorporating the presence of myocardial fibers, or a statistical model can describe the typical shape of the left ventricle, its typical appearance (i.e., the local grayscale properties of the myocardium), or even its typical motion. Using a statistical model for motion estimation consists of two stages (Figure 7). The first step involves building a large annotated database containing manually segmented volumetric data sets of healthy and pathologic hearts. This database is then used to learn the normal and pathologic geometry and appearance or motion of a ventricle offline. In the second step, myocardial motion in a new data set can be estimated online by combining and comparing the image data with
the database. The expertise of the clinical experts delineating the ventricles is thus captured and used as a priori knowledge.

To improve tracking performance, block-matching results from a selected number of speckles can be included in the motion estimation process. This method has been implemented in the software of one of the manufacturers.37-40

**VALIDATION**

Although all 3D STE methods measure the same deformation of the heart, they do so in different ways. Furthermore, within the same category of 3D STE methods, different implementations may exist (e.g., different regularization choices or postprocessing steps). This was demonstrated in a recent study by Gayat et al.41 highlighting intervendor dependency of strain measurements, although they were all based on the same underlying 3D STE approach (i.e., block matching). Therefore, it is critical that every method is validated individually before being introduced into clinical practice.

The validation of any strain imaging method and its implementation into clinical routine can be regarded as a four-step process, beginning with a validation on simulated models. In the next step, the method is usually validated using in vitro and in vivo experiments. Finally, its implementation in clinical practice can begin. In every step, deformation measurements are compared against a reference measurement (i.e., a ground truth). With every stage, ultrasound images become more realistic (i.e., containing more image artifacts and having an increased level of noise), while obtaining a ground truth deformation measurement becomes more complicated. An overview of the current validation statuses of the available 3D STE approaches is given in Table 1.

**Validation in Simulated Models**

Synthetic ultrasound images can be simulated using dedicated software. The shape and the underlying heart motion are typically simplified (e.g., by representing the left ventricle by the top half of an ellipsoid or by simulating the motion by finite element modeling). Simulated images offer many advantages for testing the performance of a strain estimation method. The displacement (and thus strain) value is known precisely at every image point, and the sensitivity of a method to image acquisition parameters can be investigated in detail (e.g., the signal-to-noise ratio and frame rate). Dysfunctional regions can also be included to test the method’s ability to detect wall motion abnormalities. Although these images have realistic image quality, image artifacts often encountered in clinical images are usually not present, because they are difficult to simulate.

To date, block matching–based22,42,43 and elastic registration–based21,34 3D STE approaches have been shown to perform well in simulated models. These approaches provide reliable deformation curves throughout the cardiac cycle in the longitudinal, circumferential, and radial direction. Some studies have demonstrated that a simulated infarct area could be detected (i.e., an area of low deformation compared with neighboring segments).21,22,42 Moreover, the capability of two methods to capture the underlying torsional motion has also been investigated.21,22

**Validation in an In Vitro Experimental Setting**

In vitro validation involves constructing deformable cardiac phantoms, which are made of a tissue-mimicking material with similar acoustic properties as myocardial tissue.32 Myocardial scar can be mimicked by embedding inclusions of a stiffer material into the phantom wall. The phantom is then mounted into a hydraulic pumping system and submerged in a water tank to allow scanning.44 As an alternative to tissue-mimicking phantoms, some groups have attached
Excised animal hearts to the setup.25,45 These hearts are deformed using a water-filled balloon secured within the LV cavity. At this stage, reference segmental strain values are obtained from the displacements of attached sonomicrometry crystals. In contrast to the simulated models, ground-truth displacement and deformation are known only in the regions where crystals are implanted.

Preliminary results of several studies on the in vitro validation of 3D block-matching methods25,45-47 and 3D speckle-tracking echocardiography on the basis of elastic registration32 are readily available. These studies report good correlations between 3D STE and sonomicrometric regional LS ($r = 0.85–0.99$), CS ($r = 0.80–0.99$),25,32,45,47 and RS ($r = 0.98$).32 The results of these studies are summarized in Table 2.

The capability of a 3D STE method based on elastic registration to detect dysfunctional regions was tested in vitro by Heyde et al.32 In that study, only LS and CS, but not RS, could discriminate regions of simulated myocardial scar.

Validation in an In Vivo Experimental Setting

The third validation step is performed in vivo using open-chest animal models. At this stage, LV deformation is controlled indirectly by pharmacologic inotropic stimulation, and regional myocardial ischemia is induced by coronary artery ligation. Just as in an in vitro setting, sonomicrometric strain curves can be used as reference measurements. Some groups have also described magnetic resonance imaging (MRI) opaque markers that can be implanted in the myocardium to estimate reference strain values.48 The results of the 3D STE validation studies performed in this setting are also summarized in Table 2.

More than a decade ago, Papademetris et al.20,49 were the first to report on the in vivo validation of a model-based 3D STE approach against sonomicrometry20 and opaque MRI markers.49 In more recent years, different implementations of block matching–based approaches,50-54 a new implementation of a model-based approach,40 and an elastic registration 3D STE approach56 have been validated in an in vivo setting. All these studies reported moderate to good correlations ($r = 0.49–0.91$) between regional strain values obtained by 3D STE methods and sonomicrometry40,51,56 (Table 2). However, care should be taken in interpreting and comparing those results, as not all of the studies reported strain values corresponding to a local cardiac coordinate system (i.e., RS, LS, and CS). In some of these studies, principal strain components (i.e., strain values along the three major directions in which no shear strain occurs) were measured instead.20,49,57

In general, the RS component showed the worst correlation and agreement with sonomicrometry for all the tested approaches. To overcome this shortcoming, myocardial incompressibility may be assumed, and RS can then be computed from the other strain components or expressed in terms of the endocardial surface area change during the cardiac cycle (Figure 8). In the latter case, the resulting
compared with MRI tagging–derived strain estimates. The results of
approach, have been compared with 2D ST. CS and RS estimates
some manufacturers, as well as an elastic registration
strain values among different techniques.
performs better than the other, they show only the comparability of
standard method, the results of such comparisons should be
sive deformation imaging techniques can be considered a gold-
ST, MRI tagging). However, because none of the available noninva-
change ratio could detect myocardial ischemia as well as standard
strain components, and it appeared to be advantageous in discrimi-
nating between states of increased and decreased contractility.
Finally, in an in vivo study by Ashraf et al., one of the 3D block-
matching approaches was shown to be able to detect LV twist.

COMPARISON AGAINST OTHER TECHNIQUES AND
CURRENT APPLICATIONS OF 3D SPECKLE-TRACKING
ECHOCARDIOGRAPHY IN A CLINICAL SETTING

Validating methods in a clinical setting is challenging because no ref-
ence measures of myocardial deformation are available. The new
deformation imaging technique can be compared against only one
of the currently available and widely used methods (e.g., DTI, 2D
ST, MRI tagging). However, because none of the available noninva-
sive deformation imaging techniques can be considered a gold-
standard method, the results of such comparisons should be
interpreted with caution. Rather than indicating if one method
performs better than the other, they show only the comparability of
strain values among different techniques.

To date, the block matching–based 3D STE software offered by
some manufacturers, as well as an elastic registration
approach, have been compared with 2D ST. CS and RS estimates
obtained by the software of two different vendors have been compared with MRI tagging–derived strain estimates. The results of
these studies are summarized in Table 2. It should be noted that
most studies report only global strain values, while only one block-matching method and one elastic registration approach have so far been compared with 2D ST at a regional level.

Three-Dimensional Speckle-Tracking Echocardiography for the Estimation of Global LV Function

Several recent publications indicate that 3D speckle-tracking echocardiography is a trustworthy technique for the evaluation of global LV function. All demonstrated good correlations (r = 0.81–0.91) between 3D STE global strain and conventional parameters of global LV systolic function such as ejection fraction and Doppler-derived cardiac output. Moreover, 3D global LS values have been shown to be significantly lower in patients with known ischemic heart disease and patients with hypertension compared with controls. Reduced global 3D CS has been reported only in patients with ischemic heart disease.

Besides that, all of the published comparative studies have reported good correlations between global LS values obtained with 3D STE methods and with 2D ST (r = 0.72–0.91; Table 2). The reported limits of agreement (in absolute strain) between these two techniques were ± 5% with respect to the bias. It is also worth noting that in general, all of the 3D STE methods tend to measure less deformation compared with 2D ST. The block-
matching approach offered by one manufacturer, and an elastic reg-
istration approach, reported approximately 3% lower absolute strain values compared with 2D ST, whereas for the block-
matching software offered by another manufacturer, this difference was only 0.5% lower absolute strain values. Interestingly, global CS extracted by 3D speckle-tracking echocardiography seemed to be 10% higher (i.e., more negative) in comparison with tagged MRI.

Three-Dimensional Speckle-Tracking Echocardiography for the Estimation of Regional LV Function

To date, myocardial deformation imaging is most widely applied in the setting of ischemic heart disease, for which detecting areas of regional myocardial dysfunction is as important as estimating global LV function. However, it still remains unclear whether 3D STE regional strain values are as reliable as those obtained by DTI or 2D ST.

In the clinical setting, delayed enhancement MRI is commonly used as a gold-standard method to locate and quantify the extent of myocardial scar. As such, measured segmental strain values can be compared against those segments identified as being scar tissue. It is important to note that this is not a direct comparison of two techni-
ques, as strain is a parameter of myocardial function, whereas de-
layed enhancement MRI characterizes the ventricle morphologically.

To date, delayed enhancement MRI has been used as a reference method by only one 3D STE study, in which Hayat et al. found significantly decreased segmental LS, CS, and RS values obtained by both 3D block matching and 2D ST in the segments identified as transmural myocardial scar.

Occasionally, strain values are compared with the wall motion score estimated by eyeballing. However, the meaningfulness of such comparisons is questionable, as wall motion scoring is known to be rather subjective and to require considerable expertise of the observer.

Despite 3D STE methods being capable of detecting regional LV dysfunction, both regional LS and CS values extracted with a block-matching method or an elastic registration approach...
have correlated only moderately with those derived by 2D ST ($r = 0.41–0.63$). The limits of agreement between these techniques were rather large as well. For the block-matching approach, they were $6.17\%$ (LS) and $6.24\%$ (CS) with respect to the bias, while for the elastic registration approach, they were $6.9\%$ (LS) and $13\%$ (CS).

In fact, the poor observed agreement between 2D and 3D techniques at the regional level might partially be due to the difficulty in matching corresponding LV segments. This may not be an important issue in a healthy heart, in which deformation is expected to be fairly homogeneous. However, in the presence of regional myocardial dysfunction, differences in strain values measured by two techniques might be induced by inaccurately matching segments. Moreover, the agreement between CS values measured with 2D and 3D strain estimation techniques can also be influenced by the discrepancies of endocardial border definition in the 2D and 3D data sets, in combination with a relatively large intramural gradient of this strain component. For the block-matching approach, they were $6.17\%$ (LS) and $6.24\%$ (CS) with respect to the bias, while for the elastic registration approach, they were $6.9\%$ (LS) and $13\%$ (CS).

Table 1 Overview of the current validation status of available 3D STE approaches

<table>
<thead>
<tr>
<th>3D STE approach</th>
<th>Simulated models</th>
<th>In vitro experimental setting</th>
<th>In vivo experimental setting</th>
<th>Comparison with other methods in clinical setting</th>
</tr>
</thead>
<tbody>
<tr>
<td>Elastic registration</td>
<td>Elen et al. (2008)$^{21}$ De Craene et al. (2012)$^{34}$</td>
<td>Heyde et al. (2012)$^{32}$ Heyde et al. (2011)$^{56}$</td>
<td>—</td>
<td>Jasaityte et al. (2012)$^{40}$</td>
</tr>
<tr>
<td>Model based</td>
<td>—</td>
<td>—</td>
<td>Papademetriz et al. (2001)$^{20}$ Papademetriz et al. (2002)$^{49}$ Bouchez et al. (2012)$^{90,\ast}$</td>
<td>—</td>
</tr>
</tbody>
</table>

*Abstracts presented at scientific congresses.
†Only qualitative results are presented.
Table 2  Validation studies of the currently available 3D STE approaches

<table>
<thead>
<tr>
<th>3D STE approach</th>
<th>Validation study</th>
<th>Study setting</th>
<th>Reference method</th>
<th>GLS</th>
<th>GCS</th>
<th>GRS</th>
<th>SLS</th>
<th>SCS</th>
<th>SRS</th>
<th>Correlation coefficient (r) (LOA [mean bias ± 1.96 SD])^*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Block matching</td>
<td>Duan et al. (2007)^60</td>
<td>Clinical</td>
<td>MRI tagging</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>0.91 (2.18% ± 3.47%)</td>
</tr>
<tr>
<td></td>
<td>Duan et al. (2009)^67</td>
<td>In vivo</td>
<td>Sono</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>0.86 (1.8% ± 3.56%)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Seo et al. (2009)^51</td>
<td>In vivo</td>
<td>Sono</td>
<td>0.89</td>
<td></td>
<td></td>
<td></td>
<td>0.9</td>
<td>0.84</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Maffessanti et al. (2009)^55</td>
<td>Clinical</td>
<td>2D ST</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>0.49</td>
<td>(0.4 ± 16.8%)</td>
<td>0.43 (−2.8 ± 24%)</td>
<td>0.24 (−0.8 ± 52.2%)</td>
</tr>
<tr>
<td></td>
<td>Ashraf et al. (2010)^65,†</td>
<td>In vitro</td>
<td>Sono</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>0.85</td>
<td>(5.0 ± 6.86%)</td>
<td>—</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Sahn et al. (2011)^25,†</td>
<td>In vitro</td>
<td>Sono</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>0.85</td>
<td>(6.5 ± 6.86%)</td>
<td>—</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Hjertaas et al. (2011)^67,†</td>
<td>In vitro</td>
<td>Sono</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>0.95</td>
<td>(−0.02 ± 2.45%)</td>
<td>0.80 (−1.95 ± 3.04%)</td>
<td>to (2.92 ± 3.86%)</td>
</tr>
<tr>
<td></td>
<td>Seo et al. (2011)^54</td>
<td>In vivo</td>
<td>Sono</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>0.87</td>
<td>(0.45 ± 17.35%)</td>
<td>—</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Negishi et al. (2011)^59</td>
<td>Clinical</td>
<td>2D ST</td>
<td>0.72</td>
<td>(3.73 ± 6.86%)</td>
<td>—</td>
<td>—</td>
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<tr>
<td></td>
<td>Reant et al. (2012)^24</td>
<td>Clinical</td>
<td>2D ST</td>
<td>0.91</td>
<td>(1.3 ± 4.4%)</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Hayat et al. (2012)^58</td>
<td>Clinical</td>
<td>2D ST</td>
<td>0.86</td>
<td>(0.5 ± 4.29%)</td>
<td>—</td>
<td>—</td>
<td>—</td>
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<td></td>
</tr>
<tr>
<td></td>
<td>Kleijn et al. (2012)^61</td>
<td>Clinical</td>
<td>MRI tagging</td>
<td>0.8</td>
<td>(10 ± 3.32%)</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td></td>
</tr>
<tr>
<td>Elastic registration</td>
<td>Heyde et al. (2011)^56</td>
<td>In vivo</td>
<td>Sono</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>0.63</td>
<td>0.60</td>
<td>—</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Jasaityte et al. (2012)^60</td>
<td>Clinical</td>
<td>2D ST</td>
<td>0.93</td>
<td>(2.94 ± 5.19%)</td>
<td>0.86</td>
<td>(2.48 ± 4.51%)</td>
<td>—</td>
<td>0.63</td>
<td>(3.28 ± 9.56%)</td>
</tr>
<tr>
<td></td>
<td>Heyde et al. (2012)^32</td>
<td>In vitro</td>
<td>Sono</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>0.96</td>
<td>0.92</td>
<td>0.98</td>
<td>—</td>
</tr>
<tr>
<td>Model based</td>
<td>Papademetris et al. (2001)^20</td>
<td>In vivo</td>
<td>Sono</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>0.89</td>
<td>—</td>
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<td></td>
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<tr>
<td></td>
<td>Papademetris et al. (2002)^49</td>
<td>In vivo</td>
<td>MRI markers</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>0.84</td>
<td>0.46 (0.73)</td>
<td>0.85 (0.91)</td>
<td>—</td>
</tr>
<tr>
<td></td>
<td>Bouchez et al. (2012)^50,†</td>
<td>In vivo</td>
<td>Sono</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>0.74</td>
<td>0.91</td>
<td>0.49</td>
<td>—</td>
</tr>
</tbody>
</table>

ACR, Endocardial area change ratio; GCS, global CS; GLS, global LS; GRS, global RS; LOA, limits of agreement; SCS, segmental CS; SLS, segmental LS; Sono, sonomicrometry; SRS, segmental RS.

Studies are subdivided into those reporting data in an in vitro experimental setup, in an in vivo experimental setup, or in a clinical setting. When available, correlation coefficients and limits of agreement are reported for either global or segmental strain.

*Bias is reported as absolute strain value (3D speckle-tracking echocardiography – reference method).
†Abstracts presented at scientific congresses.
‡Correlations obtained for LS and CS values together.
§All principal strain components.
{k} Each principal strain component individually.
{i} Range over different frame rate settings.
true, as reflected by their good correlations with sonomicrometry in an in vivo experimental setting.\(^{20,40,49,51,56,57}\)

**RS Estimates with 3D Speckle-Tracking Echocardiography**

As can be noted from Table 2, all current 3D strain estimation approaches can estimate LS and CS with acceptable accuracy, whereas the estimation of RS remains difficult.\(^{50,55}\) Either the correlation was moderate to poor or no correlations were found or reported. This suboptimal performance could be related to the fact that the spatial motion gradient must be calculated over a relatively small region because of the limited wall thickness, in combination with limited spatial resolution. Again, because of the orientation of the left ventricle in the volumetric data, beam density and consequently spatial resolution are lower in the radial direction than, for example, in the longitudinal direction. It is also worth mentioning that until the present day, RS estimation has been more difficult and prone to errors, even with 2D strain estimation techniques.\(^{65}\) Currently, the radial component may thus better be estimated by assuming volume conservation over the myocardium, as mentioned earlier (also see the Appendix).

**Reproducibility of 3D STE Measurements**

An equally important aspect for the clinical applicability of 3D speckle-tracking echocardiography is the reproducibility of the strain measurements. The reported intraindividual and interobserver variability for the different 3D STE approaches, listed for each strain component separately, is shown in Table 3. Note that in the text below as well as in Table 3, the variability of the strain measurements is reported as the ratio between the absolute difference and the mean of two repeated measurements expressed as a percentage.

To summarize, intraindividual variability of global 3D STE strain measurements obtained with different methods varied from 1% to 13%, whereas interobserver variability varied from 2% to 14%. The intraindividual and interobserver variability for regional 3D STE strain measurements ranged from 3% to 18% and from 5% to 25%, respectively. In general, RS values showed higher variability than values of LS and CS. By means of comparison, the interobserver variability of DTI-derived segmental strain values ranged from 16% to 18%.\(^{66}\) The reported 2D ST intraindividual and interobserver variability was 11%\(^{67}\) and 14%,\(^{68}\) respectively. In contrast, a widely recognized parameter such as LV ejection fraction has intraindividual variability of >10% when no contrast enhancement is used.\(^{69,70}\) The repeatability of current 3D strain measurements thus has the same order of variability, if not lower, than the other established techniques for the quantification of cardiac function.

Interexamination variability or test-retest repeatability of 3D speckle-tracking echocardiography has been assessed in only three studies.\(^{24,30,62}\) Interexamination variability of the different global strain components ranged from 8% to 11% in one study.\(^{24}\) In two other studies performed in the same patient group, test-retest repeatability expressed as an intraclass correlation coefficient ranged from 0.41 to 0.77 for the different segmental strain components.\(^{30,62}\)

It is worth noting that better reproducibility of strain measurements can be obtained for techniques with lower degrees of user interaction. Indeed, 3D STE methods using fully automatic or semiautomatic segmentation to indicate the region of interest tend to have better reproducibility than DTI, 2D ST, or other 3D STE methods in which contouring of the myocardial borders and segmentation must be done manually.

**Clinical Potential of 3D Speckle-Tracking Echocardiography**

Several recently published studies have shown the potential of 3D speckle-tracking echocardiography to replace 2D ST and DTI in a clinical setting in which the value of deformation imaging is already proven (e.g., the detection of myocardial ischemia).\(^{1-3}\) Evidently, full-volume data sets can be acquired in a more time efficient manner compared with multiple 2D cross-sections from several standard 2D planes. It also seems advantageous over 2D ST and DTI when significant and fast changes in loading conditions of the heart or heart rate can be expected, because a single acquisition is sufficient to obtain the deformation of all LV and right ventricular segments at once. Being able to provide a physician with a fast overview of the deformation properties of the heart further adds to this technique’s appeal and makes it attractive in routine clinical practice.

Furthermore, 3D speckle-tracking echocardiography can provide a more robust measure of torsion, because it overcomes the major shortcomings of 2D ST by enabling the easy acquisition of parallel LV short-axis planes and precise measurement of the distance between them. Moreover, because 3D STE methods calculate the complex deformation of the entire myocardium, the extraction of shear strain components, which combine deformation in two different directions, will become feasible.

One of the biggest expectations in the future for 3D speckle-tracking echocardiography is its ability to provide better insight into cardiac physiology and disease processes. For example, instead of using a standardized 17-segment approach for LV segmentation (as proposed by the American Heart Association),\(^{71}\) more pathology-specific segments (e.g., infarct core, adjacent or remote myocardium, coronary artery territories)\(^{72}\) could be used to examine myocardial deformation properties with 3D speckle-tracking echocardiography. Furthermore, measuring regional curvatures of the ventricle in a robust manner might soon become possible from volumetric data sets. As such, it would help estimate segmental LV load, which
combined with strain measures would give more information about regional LV function.

**CURRENT LIMITATIONS OF THREE-DIMENSIONAL SPECKLE-TRACKING ECHOCARDIOGRAPHY**

General advantages and limitations of 3D STE approaches in comparison with DTI and 2D ST are summarized in Table 4. Indeed, good reproducibility of strain values and time efficiency make 3D STE methods very attractive for clinical users. However, multiple factors can affect the performance and reliability of 3D strain estimation in certain clinical scenarios, such as stress echocardiography or the detection of LV dyssynchrony. Some of these factors are intrinsic to the technology, while others can be avoided.

**Feasibility and Image Quality**

Most of the current ultrasound systems use ECG gating to produce volumetric data sets of sufficient temporal and spatial resolution. As such, the acquisition of volumetric data sets is limited to patients with no heart rhythm irregularities. Additionally, patients must be capable of holding their breath for at least four cardiac cycles to allow ECG gating. These technical factors may dramatically limit the number of patients in whom 3D speckle-tracking echocardiography is feasible.

High-quality volumetric data sets are crucial for reliable 3D strain estimates. In clinical practice, this can be challenging, because shadowing artifacts and signal dropouts often occur. It also requires a sufficient amount of training and skill to acquire high-quality data sets for strain quantification.\(^7^3\)

The currently reported feasibility of 3D STE methods ranges from 63% to 83%\(^2^4,5^8-6^0\) and is somewhat lower than that of 2D ST techniques (80%–97%).\(^1^8,7^4\) Moreover, it should be noted that the feasibility of 3D speckle-tracking echocardiography was calculated after the exclusion of patients with arrhythmic disorders or patients who were not able to perform a sufficient breath hold.

**Intervendor Dependency of Strain Values**

As already mentioned, similar strain estimation methods can be implemented in many different ways (e.g., different regularization or postprocessing steps). Furthermore, nomenclature can differ among different vendors (e.g., instead of RS, area strain may be reported) and may mislead inexperienced users. It remains difficult to compare normal strain values among different commercial systems, largely because these platforms are closed (i.e., they act like black boxes). The associated high intervendor dependency of strain values is a known issue but is related not only to 3D STE techniques\(^4^1\) but also to 2D ST\(^7^2\) and DTI.\(^7^5-7^7\)

However, it is worth mentioning that a working group of experts has recently been initiated by the European Association of Cardiovascular Imaging and the American Society of Echocardiography to work on the standardization of 2D ST software. Hopefully, similar measures will be undertaken in the future to standardize 3D STE software as well.

**Temporal Resolution**

Three-dimensional echocardiography comes at the expense of temporal resolution, which may limit its applicability in patients with high heart rates (e.g., during stress echocardiography and in patients with tachycardia). Moreover, because of the intrinsically low frame
rate, the precise timing of peak systolic strain values is difficult. As such, one might question how reliably 3D speckle-tracking echocardiography can currently detect LV dyssynchrony (which may be defined by the standard deviation of time to peak systolic strain in 16 LV segments). However, some studies have shown that this might potentially provide interesting insights.78,79 The frame rate of 3D acquisitions can be increased by ECG gating, but stitching artifacts caused by breathing or probe motion may limit its use for functional quantification. Recently, advances in ultrasound beam forming have led to systems capable of acquiring real-time 3D volumes with reasonable temporal resolution. However, to date, no validation studies of those systems have been published.

Interestingly, Byram et al.80 investigated the influence of frame rate on the tracking performance of a block matching–based 3D ST method. For this purpose, an experimental setup with a deformable phantom was built, and images were acquired at a volume rate between 50 and 1,000 Hz. Motion estimates showed improvements up to a frame rate of 200 Hz. Moreover, in a clinical setting, the influence of frame rate on the global strain values was evaluated recently by Negishi et al.59 They demonstrated that one of the block matching–based 3D STE approaches gives global LS values comparable with those obtained using 2D techniques at frame rates of 34 to 50 Hz. Strain values extracted from acquisitions obtained at lower or higher frame rates did not correlate well. Similar results were reported in a phantom study by Hjertaas et al.47 The frame rate was modified in the range of 22 to 52 Hz. The highest correlations against sonomicrometry for LS and CS were obtained in the middle of this range (37 Hz). This is not surprising, because too low frame rate results in higher speckle decorrelation and thus less influenced by increased speckle decorrelation between frames.

**Strain Rate Estimation**

It has been elegantly shown that even though both strain and strain rate increase with dobutamine and decrease with esmolol infusion, they are not equal in estimating contractile function.81,82 In fact, strain is very much dependent on LV loading conditions, structure, and heart rate, thus making this parameter mainly a reflection of LV stroke volume.81,82 Strain rate, on the other hand, is less influenced by these factors.81,82 As such, strain rate can yield important insights into myocardial contractility and give a more complete view of regional myocardial mechanics.

Unfortunately, because of the low temporal resolution of 3D speckle-tracking echocardiography, none of the currently available 3D STE methods is suitable for strain rate measurements. Indeed, frame rates of 20 to 40 Hz might be sufficient for strain estimation, but frame rates $\geq 160$ Hz are required to measure strain rate reliably.83

**Spatial Resolution**

Finally, all current 3D STE approaches seem to have difficulties estimating deformation in the basal parts of the left ventricle. The lowest correlation between the block matching-based 3D STE approach and 2D ST or triplane imaging was seen in the basal LV segments.59 The elastic registration–based 3D STE approach had the largest displacement estimation errors in the basal segments in simulated models.21 Consistently, in human data sets, the biggest differences in 2D and 3D LS values were seen in the basal segments as well.60 This is not entirely unexpected, as these segments move at the highest velocities during the cardiac cycle, which may be more difficult to measure accurately, especially given the already mentioned low frame rate of the current 3D ultrasound data sets. Moreover, it seems plausible that this is related to the spatial resolution as well. Because of the orientation of the left ventricle in the image volume, ultrasound beams diverge at the base, resulting in a lower spatial resolution and making tracking intrinsically harder.

**FUTURE DEVELOPMENTS AND CONCLUSIONS**

To date, different 3D STE methods have been proposed, all differing primarily in their underlying tracking mechanisms. However, to the best of our knowledge, no direct comparison studies have been reported so far. As such, there is currently no preferred method, and they all remain useful.

Initial experiences with 3D strain estimation are promising, although there is still room for improvement. The techniques can already successfully be applied for the estimation of global LV strain, but regional strain estimation remains challenging. In the future, more validation studies will be required to investigate their capability in detecting dysfunctional regions. Their biggest shortcoming is their inability to measure strain rate and the timing of peak strain values precisely, while the measurements of RS might be replaced by its calculation from LS and CS components.

Major research efforts are currently made to develop new advanced ultrasound transducers and beam-forming techniques that...
will improve spatial and temporal resolution. This will improve the performance of the current 3D STE techniques. In parallel, industry and academia are working on merging different tracking methods, combining their advantages while counteracting their individual disadvantages. Using more automatic segmentation strategies in the future should also have a positive effect on the reproducibility of strain measurements.

Furthermore, 3D STE software packages offered by different vendors should be standardized, and population-based studies should be initiated to define normal reference values for 3D STE global and regional strain values. Finally, large-scale clinical studies will be required to assess whether 3D STE strain values have any diagnostic or prognostic value in different clinical settings before these methods can effectively translate into clinical use.

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APPENDIX

Radial Strain Estimation

Because a direct estimation of the RS component remains difficult, some authors and vendors have proposed computing it from the other strain components. This appendix highlights how these calculations can be done and what assumptions they are based on.

Consider a small myocardial element, as shown in Figure 8. In reality, the surfaces of this element are curved, but locally this element may be considered a cube with initial dimensions $R$, $L$, and $C$ in the radial, longitudinal, and circumferential directions, respectively. The initial volume ($V_0$) of this cube is simply the product of its sides:

$$V_0 = RLC.$$  
(A1)

If the volume is conserved during deformation (i.e., by assuming myocardial incompressibility), $\Delta V$ must equal zero, implying from the above equation that

$$(R + \Delta R)(L + \Delta L)(C + \Delta C) - RLC = 0.$$  
(A2)

Noting that $\Delta R/R$ is the change in length of the radial side of the cube with respect to its original length ($R$), this is the RS. With similar reasoning for the longitudinal and circumferential directions, we thus obtain

$$(1 + RS)(1 + LS)(1 + CS) = 1.$$  
(A3)

When expanding the above product, we find

$$1 + RS + LS + CS + RS \cdot CS + RS \cdot LS + LS \cdot CS + RS \cdot CS \cdot LS = 1.$$  
(A5)

If the strain values are small (i.e., RS, CS, and LS are much smaller than 1), their products become even smaller and can, to a first order of approximation, be neglected. As such, in the case of small deformations, equation A5 further simplifies to

$$RS + LS + CS = 0.$$  
(A6)

or

$$RS = - (LS + CS).$$  
(A7)

stating that RS is the negative sum of LS and CS.

Using the above notations, the endocardial area ($A$) of the cube can simply be written as $LC$. Area strain (AS) is defined as the relative change in surface area over the cardiac cycle and can thus be written as

$$AS = \frac{1}{2} \left[ (A + \Delta A) - A \right] = \frac{1}{2} \left[ (L + \Delta L)(C + \Delta C) - LC \right] / LC.$$  
(A8)

which reduces to

$$AS = (1 + \Delta L/L)(1 + \Delta C/C) - 1.$$  
(A9)

or

$$AS = (1 + LS)(1 + CS) - 1.$$  
(A10)

Note that this deformation parameter is also referred to as area change ratio by some vendors. Again, if a small deformation can be assumed, then equation A10 further simplifies to

$$AS = (LS + CS).$$  
(A11)

In other words, $AS$ is the sum of LS and CS. From equation A7, it is then obvious that RS is simply the negative of AS (when assuming volume conservation):

$$RS = - AS.$$  
(A12)