Assessment of the relationships between myocardial contractility and infarct tissue revealed by serial magnetic resonance imaging in patients with acute myocardial infarction

Christie McComb1,2 • David Carrick2,3 • John D. McClure3 • Rosemary Woodward2,4 • Aleksandra Radjenovic2 • John E. Foster1,2 • Colin Berry2,3

Received: 25 December 2014 / Accepted: 11 May 2015 / Published online: 6 June 2015
© The Author(s) 2015. This article is published with open access at Springerlink.com

Abstract Imaging changes in left ventricular (LV) volumes during the cardiac cycle and LV ejection fraction do not provide information on regional contractility. Displacement ENcoding with Stimulated Echoes (DENSE) is a strain-encoded cardiac magnetic resonance (CMR) technique that measures strain directly. We investigated the relationships between strain revealed by DENSE and the presence and extent of infarction in patients with recent myocardial infarction (MI). 50 male subjects were invited to undergo serial CMR within 7 days of MI (baseline) and after 6 months (follow-up; n = 47). DENSE and late gadolinium enhancement (LGE) images were acquired to enable localised regional quantification of peak circumferential strain (Ecc) and the extent of infarction, respectively. We assessed: (1) receiver operating characteristic (ROC) analysis for the classification of LGE, (2) strain differences according to LGE status (remote, adjacent, infarcted) and (3) changes in strain revealed between baseline and follow-up. 300 and 258 myocardial segments were available for analysis at baseline and follow-up respectively. LGE was present in 130/300 (43 %) and 97/258 (38 %) segments, respectively. ROC analysis revealed moderately high values for peak Ecc at baseline [threshold 12.8 %; area-under-curve (AUC) 0.88, sensitivity 84 %, specificity 78 %] and at follow-up (threshold 15.8 %; AUC 0.76, sensitivity 85 %, specificity 64 %). Differences were observed between remote, adjacent and infarcted segments. Between baseline and follow-up, increases in peak Ecc were observed in infarcted segments (median difference of 5.6 %) and in adjacent segments (1.5 %). Peak Ecc at baseline was indicative of the change in LGE status between baseline and follow-up. Strain-encoded CMR with DENSE has the potential to provide clinically useful information on contractility and its recovery over time in patients with MI.

Keywords Myocardial infarction • Strain-encoded CMR • Contractility

Background

Regional left ventricular (LV) systolic function has prognostic importance in survivors of myocardial infarction (MI) [1, 2]. Imaging of LV wall motion is an indirect method for assessing contractility, and cardiac magnetic resonance (CMR) can now provide information on LV contractility using strain-specific techniques. Measurement of strain rather than LV motion should theoretically increase the diagnostic accuracy and prognostic value of cardiac imaging in patients with a history of MI [3]. The fact that CMR provides quantitative information on infarct pathology as well as myocardial strain makes this imaging modality attractive for clinical and research purposes in patients with a history of MI.

During the cardiac cycle, the heart moves through and rotates within a given imaging plane. The relationship between the apparent motion of the endocardial and
epicardial borders and the contraction and relaxation of the myocardium is complex, and a true assessment of contractile function requires tracking of discrete points within the myocardium [4]. Three types of magnetic resonance imaging (MRI) pulse sequence which have previously been used for the visualisation and quantification of myocardial deformation are tagging [5–8], velocity-encoded phase contrast imaging [9–12] and strain-encoded (SENC) imaging [13].

The main drawbacks of myocardial tagging and SENC techniques are that the need to apply a saturation grid limits the resolution, and quantitative analysis can be complicated and time consuming. Velocity-encoded phase contrast methods are sensitive to accumulated errors that can result in inaccuracies in position measurements, and displacements can only be measured over short periods of time.

Displacement ENcoding with Stimulated Echoes (DENSE) is an alternative method for the quantification of myocardial strain [14, 15]. Stimulated echoes are used to provide a high spatial density of displacement measurements in the myocardium over long time intervals, which enable large displacements to be encoded. Since displacement is directly encoded in the phase, there is no requirement for tag detection and analysis is less time-consuming than for tagging.

Much of the research performed with DENSE so far has been methods development, and its diagnostic value in a clinical setting is uncertain. Recently, Miyagi et al. [16] reported that circumferential strain revealed by DENSE is associated with the presence of late gadolinium enhancement (LGE). However, the relationships between regional LV contractility and infarction are not uniform. Regional contractility may be influenced by the spatial proximity to the infarct tissue, which itself may change in composition and size according to the time interval from the acute MI event [17]. Therefore, we aimed to assess the relationships between regional myocardial contractility revealed by DENSE and LGE in patients with acute MI undergoing serial imaging.

Methods

Study population

Fifty male patients (age 56 ± 10 years) underwent CMR within 7 days of ST-elevation myocardial infarction (STEMI), and were invited to return for a follow-up scan after 6 months. All patients underwent revascularization with primary percutaneous coronary intervention (PCI) in the Golden Jubilee National Hospital, according to contemporary standards of care.

The research protocol was approved by the regional Research Ethics Committee and written informed consent was obtained from each subject.

Cardiac MRI

All images were obtained using a 1.5 T Siemens Avanto MRI scanner (Siemens, Erlangen, Germany) with a 6-channel phased-array body coil (anterior) and an 8-channel phased-array spine coil (posterior). The CMR protocol included DENSE imaging for the assessment of strain and LGE imaging in a co-registered mid-ventricular short axis slice, positioned at mid-papillary level. Cine imaging was also acquired to enable quantification of LV dimensions and function.

DENSE imaging parameters were as follows: echo time 8 ms; repetition time 16.3 ms; flip angle 20°; slice thickness 8 mm; field of view 360 mm × 270 mm; matrix size 112 × 84; displacement encoding of 0.2 π/mm; EPI factor of 8.

Early gadolinium enhancement (EGE) imaging was acquired 1, 3, 5 and 7 min post-contrast injection using a TrueFISP readout and fixed inversion time (TI) of 440 ms. Late gadolinium enhancement images covering the entire LV were acquired 10–15 min after IV injection of 0.15 mmol/kg of gadoterate meglumine (Gd²⁺-DOTA, Dotarem, Guebert S.A., Villepinte, France) using segmented phase-sensitive inversion recovery (PSIR) turbo fast low-angle shot sequence in all cases [18]. LGE imaging parameters were as follows: echo time 3.4 ms; repetition time 8.7 ms; flip angle 20°; slice thickness, 8 mm; field of view 340 mm × 270 mm; matrix size 256 × 156. The voxel size was 1.8 × 1.3 × 8 mm³. A Look-Locker TI scout scan was undertaken to determine the inversion times associated with optimal nulling of the myocardial signal. The inversion times were in the range of 260–350 ms.

Cine images were acquired using a b-SSFP sequence with the following parameters: echo time 1.2 ms; repetition time 3.3 ms; flip angle 70°; slice thickness 7 mm; field of view 340 mm × 270 mm; matrix size 256 × 180.

The imaging protocol was the same for all patients for the baseline and follow-up CMR scans.

Image analysis

The images were analysed on a Siemens work-station by observers with at least 3 years CMR experience (C.M., D.C.). The LGE and DENSE analysis were performed independently by two trained operators (DC, CM) who were blinded to the results of each other’s findings.
Analysis of left ventricular volumes and ejection fraction

Left ventricular dimensions, volumes and ejection fraction were quantified using computer-assisted planimetry (syngo MR®, Siemens Healthcare, Erlangen, Germany). Endo- and epicardial borders were manually delineated on the cine images, and LV dimensions and systolic function (ejection fraction) were measured with the automated analysis software. The LV was segmented using the anterior right ventricular-LV insertion point as the reference point.

Analysis of strain-encoded MRI with DENSE

Each of the short axis DENSE images were divided into six segments according to the American Heart Association (AHA) model in order to standardise the approach to regional analysis of LV strain [19]. Endocardial and epicardial borders were contoured and strain values were measured on a per-segment basis.

DENSE images were analysed using CIM_DENSE2D software (University of Auckland, Auckland, New Zealand) [20], and for each myocardial segment, values were obtained for peak circumferential strain (Ecc) (%).

Analysis of late gadolinium enhancement MRI

The presence of acute infarction was established based on abnormalities in cine wall motion, rest first-pass myocardial perfusion, and delayed-enhancement imaging in two imaging planes. LGE image analysis was performed using Argus software (Siemens, Erlangen, Germany). Each of the short axis LGE images were divided into six segments according to the AHA model [19]. A region of interest (ROI) containing at least 100 pixels was drawn in an area of remote myocardium, and a threshold level was set at the mean value of the ROI plus 5 standard deviations (SD) [21]. The territory (area) of infarction was delineated manually using computer-assisted planimetry and based on the 5 SD threshold above the remote reference region ROI. The myocardial mass of late gadolinium (grams) was quantified and expressed as a percentage of total LV mass.

Tissue categorisation: designation of zones of tissue based on pathology

The terms ‘remote’, ‘infarct’ and ‘adjacent’ were used to reflect areas of myocardial tissue that were defined by the presence (‘infarct’ and ‘adjacent’) or absence (‘remote’) of MI pathology, as revealed by LGE on MRI. The remote myocardium was defined as myocardium 180° from the affected zone with no visible evidence of infarction or wall motion abnormalities (assessed by inspecting corresponding contrast enhanced LGE and cine images, respectively).

Statistical analysis

All analyses were performed using Minitab 16 (Minitab Inc, PA, USA), with the exception of ROC analysis, which was performed using SPSS 19 (IBM, New York, USA).

Prior to performing statistical analysis, all data were checked for normality using Anderson–Darling tests. Peak Ecc was found to be non-normally distributed, and analysis was therefore performed using non-parametric tests.

ROC analysis, sensitivity and specificity

ROC analysis was performed based on the presence or absence of LGE within each segment, at both baseline and follow-up. Thresholds for peak Ecc were established by finding the values which corresponded to the maximum average sensitivity and specificity.

A score was allocated to each segment according to the transmural extent of infarction (percentage of LGE) within the segment (0: 0 %, 1: 1–25 %, 2: 26–50 %, 3: 51–75 %, 4: 76–100 %). For each score, the percentage of segments which were correctly identified as containing LGE (scores 1–4) or not containing LGE (score 0) was calculated.

Comparison with LGE Status

At both baseline and follow-up, segments were classified into three groups depending on LGE status: infarcted, adjacent (infarction in one or more adjacent segments) and remote (no infarction in adjacent segments). Strain values in the three groups were compared using a Kruskal–Wallis test along with individual Mann–Whitney tests.

Assessment of longitudinal changes

For segments categorised as remote, adjacent and infarcted at the baseline scan, Wilcoxon signed rank tests were used to compare the baseline and follow-up results for each category separately. A Kruskal–Wallis test along with individual Mann–Whitney tests was then used to compare the strain differences (follow-up—baseline) between the three categories.

The data were then further categorised according to the change in LGE status between baseline and follow-up e.g. remote (baseline) → remote (follow-up), remote (baseline) → adjacent (follow-up), adjacent (baseline) → infarcted (follow-up) etc. Segments which were categorised as remote, adjacent and infarcted at baseline were considered separately, and the changes in strain (follow-up—baseline) between sub-categories [e.g. remote (baseline) → remote (follow-up) vs remote (baseline) → adjacent (follow-up)]
were compared using Kruskal–Wallis with individual Mann–Whitney tests. A similar comparison was then performed to assess if differences in strain values obtained at baseline could be detected between sub-categories.

A Bonferroni correction for multiple testing was used with non-parametric tests including the Kruskal–Wallis and Mann–Whitney tests. A p value of 0.05 was adopted to reject the null hypothesis of no difference.

**Results**

Table 1 summarises the clinical characteristics of the STEMI patients and LV function and volumes as determined by cine CMR. The images obtained from a 70-year-old male with an acute sub-endocardial scar in the lateral left ventricular wall are shown in Fig. 1.

From the 50 STEMI patients who underwent baseline scans, 47 attended for follow-up scans, and DENSE images from 4 of these patients were considered non-diagnostic due to breathing artefacts. This gave totals of 300 and 258 segments which were available for analysis at baseline and follow-up respectively. LGE was present in 130/300 (43 %) and 97/258 (38 %) segments, respectively.

**ROC analysis, sensitivity and specificity**

The ROC curves for strain-encoded CMR with DENSE and the classification of LGE (present/absent) on a per-segment basis at baseline and at follow-up are shown in Fig. 2.

The peak Ecc strain threshold, area-under-the-curve (AUC), sensitivity and specificity for the classification of contractile abnormalities associated with the presence of LGE at baseline and follow-up are shown in Table 2. The AUC for detection of infarct tissue (LGE) by DENSE was 0.88 on baseline scans and 0.76 at follow-up. The percentage of segments which were correctly identified as containing no LGE (score 0) and containing LGE (scores 1–4) are shown in Table 3. The percentage of segments that were correctly identified at baseline and at follow-up was greatest for scores LGE scores of 0 (no LGE) and 4 (100 % transmural extent of LGE). Further investigation of the non-infarcted segments (score 0) which were incorrectly classified by DENSE as containing LGE showed that 61 and 52 % were located adjacent to segments which contained LGE at baseline and follow-up respectively.

**Comparison of myocardial strain with the presence and transmural extent of LGE**

Differences were observed for peak Ecc measured in infarcted segments versus peak Ecc in both remote and adjacent segments, and between peak Ecc in remote and adjacent segments, at both baseline and follow-up. The results are illustrated in Fig. 3.

**Relationships between myocardial strain and LGE revealed by serial imaging after acute MI**

From the 43 patients who attended for both baseline and follow-up scans, 122 segments had LGE at baseline and/or at follow-up.

A comparison of the peak Ecc strain values obtained at baseline and follow-up found differences in segments classified as adjacent and infarcted, but not in remote segments. An increase in peak Ecc was observed in both adjacent and infarcted segments, with median differences (follow-up—baseline) of 1.5 and 5.6 % respectively.

The results of the Kruskal–Wallis test with individual Mann–Whitney tests comparing the three categories of tissue pathology (remote, adjacent, infarcted) are illustrated in Fig. 4a. Differences were observed between the change in strain in remote and infarcted segments, and between adjacent and infarcted segments.

### Table 1 Characteristics of STEMI patients (n = 50)

| Characteristic                      | Value       |
|------------------------------------|-------------|
| Age (years)                        | 56 ± 10 (33–80) |
| BMI                                | 28 ± 4 (20–38) |
| Smoker                             | 38 (63 %)   |
| Hypertension                       | 11 (18 %)   |
| Hypercholesterolaemia              | 10 (17 %)   |
| Prior myocardial infarction        | 1 (2 %)     |
| Culprit coronary artery            |             |
| Left anterior descending           | 19 (38 %)   |
| Left circumflex                    | 9 (18 %)    |
| Right                              | 22 (44 %)   |
| Electrocardiogram                  |             |
| Q-waves                            | 25 (50 %)   |
| Troponin T (ng/l)                  | 3353 (2005, 6625) |
| LVEF (%)                           |             |
| Baseline                           | 54 ± 9 (35–70) |
| Follow-up                          | 62 ± 8 (40–76) |
| LVEDV (ml)                         |             |
| Baseline                           | 153 ± 33 (85–231) |
| Follow-up                          | 163 ± 69 (92–251) |
| LVESV (ml)                         |             |
| Baseline                           | 72 ± 23 (35–119) |
| Follow-up                          | 64 ± 26 (23–150) |

Age, BMI, LVEF, LVEDV and LVESV are expressed as mean ± SD (range). Troponin T is expressed as median (interquartile range). The electrocardiogram (ECG) was obtained during the index hospitalisation.

BMI body mass index, LVEF LV ejection fraction, LVEDV LV end-diastolic volume, LVESV LV end-systolic volume.
Within each designated pathological tissue category (remote, adjacent, infarcted), comparison of tissue categories showed no statistically significant differences when...
considering the change in strain between baseline and follow-up. However, differences were observed when comparing baseline strain between sub-categories, as illustrated in Fig. 4b.

On a per-segment basis at the mid-ventricular level, peak Ecc (Fig. 5a) and change in peak Ecc at follow-up from baseline (Fig. 5b) correlated moderately well with the transmural extent of infarction at baseline, and these relationships were consistently similar for each of the anatomical regions.
Discussion

The results of this investigation have shown that, firstly, in line with previous studies [17], the number of segments with evidence of LGE is less at 6 months compared with 2 days post-MI. Secondly, peak myocardial circumferential strain as measured by DENSE has high discriminative value between healthy myocardium and infarct tissue as revealed by LGE. Thirdly, this relationship was observed within the first week after STEMI and again at a follow-up scan 6 months later, and it was influenced by the time from the index event, the proximity of the segment to the infarct territory, and the transmural extent of MI. Finally, the change in peak circumferential strain over time i.e. strain recovery is associated with the transmural extent of infarction at baseline.

The standard CMR assessment of LV function in patients with a history of MI involves using cine images to measure global parameters such as LV end-systolic volume, LV end-diastolic volume and LV ejection fraction. However, most pathologies do not affect the heart uniformly, and regional dysfunction may be masked by seemingly “normal” values of global function [22]. Assessment of regional myocardial function therefore has an important role to play in the diagnosis and management of cardiac disorders, and in determining the long-term prognosis for these patients, including in patients with a history of MI.

In a study which used strain measurements obtained with SENC to compare myocardial segments with non-transmural and transmural infarction defined as >75 % hyperenhancement in LGE imaging, an optimal threshold for peak Ecc of 10 % was identified, with a corresponding sensitivity of 97 %, specificity of 94 % and AUC of 0.96 [23]. A similar study which used tagging to compare non-infarcted and infarcted segments identified a threshold for peak Ecc of 20 %, with a corresponding sensitivity of 92 % and a specificity of 99 % [3]. More recently, Miyagi et al used ROC analysis to compare non-infarcted and infarcted segments, and obtained an AUC value of 0.92 for peak Ecc [16]. In this study, thresholds for peak Ecc of 13 and 16 % were identified at baseline and follow-up respectively. The corresponding sensitivity, specificity and AUC values shown in Table 2 were lower than those obtained in other studies, but were still considered to be high, particularly at baseline.

The higher threshold value identified at follow-up compared to baseline indicates that the presence of LGE at follow-up may be associated with a lower degree of contractile dysfunction, however the relationship between peak Ecc and LGE may be complicated by the complex remodelling process. In a patient with recent MI, the injured territory revealed by LGE may also contain oedema and viable myocardium amenable to salvage [17]. The potential for salvage is influenced by clinical factors, such as the timing and success of coronary revascularization [24]. Therefore, the presence of LGE in a patient with recent MI may be associated with functional recovery. Alternatively, the persistence of chronic LGE at follow-up is typical of permanent infarction although the extent of infarction typically falls over time reflecting infarct remodelling and salvage of acutely jeopardised myocardium in some segments, but not in other segments. Given the complexities of infarct remodelling and salvage, direct measurement of regional LV contractility could provide further information on the potential for recovery or not, for clinical and research purposes.

Differences in peak circumferential strain between remote and infarcted segments can be detected using tagging [25]. The results of this study have shown that measurements of peak Ecc obtained using DENSE can additionally reveal differences between remote and adjacent segments, not only at baseline but also at 6 months post-MI.

We observed that when segments were sub-categorised according to the change in LGE status between baseline and follow-up, there were differences in baseline strain between sub-categories for adjacent and infarcted segments. Higher strain values were obtained in adjacent segments which would subsequently be re-classified as remote, and lower strain values were obtained in segments which would be re-classified as infarcted. Infarcted segments which would subsequently be re-classified as adjacent had higher strain than those which remained infarcted. Strain measurements in adjacent and infarcted segments at baseline may have prognostic value relating to the progression or recovery of contractile abnormalities in the chronic setting.

Limitations

The results of this study show that DENSE has the potential to provide clinically useful information in patients with recent and chronic MI. However, the limitations in the design should be noted. Only one mid-ventricular slice was obtained for each participant, and the analysis method does not take adjacent segments and/or slices into account. The implementation of DENSE available at our institution does not account for through-plane motion of the myocardium during image acquisition; however 3D versions of the DENSE pulse sequence have been developed [26–28] and could be assessed in future studies.

Conclusions

DENSE has the potential to provide clinically useful information relating to contractile abnormalities associated with the presence of LGE, and the recovery of contractile
function over time. Continuing developments in the technique, including 3D versions which account for through-plane motion, will increase the applicability in the clinical setting.

Acknowledgments This work was supported by Health Science Scotland, Medical Research Scotland and British Heart Foundation Project Grant PG/11/2/28474, and through a research agreement with Siemens Healthcare. The authors thank Peter Weale, cardiac applications specialist at Siemens, for his assistance during this study.

Conflict of interest None.

Open Access This article is distributed under the terms of the Creative Commons Attribution 4.0 International License (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made.

References

1. Heikkila J, Tabakin BS, Hugenholzt PG (1972) Quantification of function in normal and infarcted regions of the left ventricle. Cardiovasc Res 6:516–531
2. Neizel M, Korosoglou G, Lossnitzer D, Kuhl H, Hoffman R, Ocklenburg C, Giannitissi E, Osman NF, Katus HA, Steen H (2010) Impact of systolic and diastolic deformation indexes by strain-encoded imaging to predict persistent severe myocardial dysfunction in patients after acute myocardial infarction at follow-up. J Am Coll Cardiol 56(13):1056–1062
3. Gotte MJ, Van Rossum AC, Twisk JWR, Kuijer JPA, Marcus JT, Visser CA (2001) Quantification of regional contracture function after infarction: strain analysis superior to wall thickening analysis in discriminating infarct from remote myocardium. J Am Coll Cardiol 37(3):808–817
4. Reichek N (1999) MRI myocardial tagging. J Magn Reson Imaging 10:609–616
5. Zerhouni EA, Parish DM, Rogers WJ, Yang A, Shapiro EP (1988) Human heart: tagging with MR imaging—a method for noninvasive assessment of myocardial motion. Radiology 169:59–63
6. Axel L, Dougherty L (1989) MR imaging of motion with spatial modulation of magnetization. Radiology 171:841–845
7. Fischer SE, McKinnon GC, Maier SE, Boesiger P (1993) Improved myocardial tagging contrast. Magn Reson Med 30:191–200
8. Ibrahim ESH (2011) Myocardial tagging by cardiovascular magnetic resonance: evolution of techniques—pulse sequences, analysis algorithms and applications. J Cardiovasc Magn Reson 13:36
9. Pelc NR, Sayre J, Yun K, Castro LJ, Herfkens RJ, Miller DC, Pelc NJ (1994) Evaluation of myocardial motion tracking with cine-phase contrast magnetic imaging. Invest Radiol 29:1038–1042
10. Pecl NJ, Drangova M, Pelc NL, Zhu Y, Noll DC, Bowman BS, Herfkens RF (1995) Tracking of cyclic motion with phase-contrast cine MR velocity data. J Magn Reson Imaging 5:339–345
11. Constable RT, Rath KM, Sinusas AS, Gore JC (1994) Development and evaluation of tracking algorithms for cardiac wall motion analysis using phase velocity MR imaging. Magn Reson Med 32:33–42
12. Arai AE, Gaither GC, Epstein FH, Balaban RS, Wolff SD (1999) Myocardial velocity gradient imaging by phase contrast MRI with application to regional function in myocardial ischaemia. Magn Reson Med 42:98–109
13. Osman NF, Sampath S, Atalar E, Prince JL (2001) Imaging longitudinal cardiac strain on short-axis imaging using strain-encoded MRI. Magn Reson Med 46:324–334
14. Aletras AH, Ding S, Balaban RS, Wen H (1999) DENSE: displacement encoding with stimulated echoes in cardiac functional MRI. J Magn Reson 137:247–252
15. Aletras AH, Balaban RS, Wen H (1999) High-resolution strain analysis of the human heart with fast-DENSE. J Magn Reson 140:41–57
16. Miyagi H, Nagata M, Kitagawa K, Kato S, Takase S, Sigfridsson A, Ishida M, Dohi K, Ito M, Sakuma H (2013) Quantitative assessment of myocardial strain with displacement encoding with stimulated echoes MRI in patients with coronary artery disease. Int J Cardiovasc Imaging 29:1779–1786
17. Dall’Armellina E, Karia N, Lindsay AC, Karamitsos TD, Ferreira V, Robson MD, Kellman P, Francis JM, Forfar C, Prendergast BD, Banning AP, Channon KM, Kharbanda RK, Neubauer S, Choudhury RP (2011) Dynamic changes of edema and late gadolinium enhancement after acute myocardial infarction and their relationship to functional recovery and salvage index. Circ Cardiovasc Imaging 4:228–236
18. Simonetti OP, Kim RJ, Fieno DS, Hillenbrand HB, Wu E, Bundy JM, Finn JP, Judd RM (2001) An improved MR imaging technique for the visualization of myocardial infarction. Radiology 218:215–223
19. Cerqueira MD, Weissman NJ, Dilsizian V et al (2002) Standardized myocardial segmentation and nomenclature for tomographic imaging of the heart. Circulation 105:539–542
20. Young AA, Kirton RS, Cowan BR (2012) Generalized spatiotemporal myocardial strain analysis for DENSE and SPAMM imaging. Magn Reson Med 67:1590–1599
21. Flett AS, Hasleton J, Cook C, Hausenloy D, Quarta G, Arti C, Muthurangu V, Moon JC (2011) Evaluation of techniques for the quantification of myocardial scar for differing etiology using cardiac magnetic resonance. JACC Cardiovascular Imaging 4(2):150–156
22. Simpson RM, Keegan J, Firmin DN (2013) MR assessment of regional myocardial mechanics. J Magn Reson Imaging 37:576–599
23. Neizel M, Lossnitzer D, Korosoglou G, Shaufele T, Peykarjou H, Kuhl H, Hoffman R, Ocklenburg C, Giannitissi E, Osman NF (2009) Strain-encoded MRI for evaluation of left ventricular function and transmurality in acute myocardial infarction. Circ Cardiovascular Imaging 2:116–122
24. Berry C, Kellman P, Mancini C, Chen MY, Bandettini WP, Lowrey T, Hsu LY, Aletras AH, Arai AE (2010) Magnetic resonance imaging delineates the ischemic area at risk and myocardial salvage in patients with acute myocardial infarction. Circ Cardiovascular Imaging 3:527–535
25. Inoue Y, Yang X, Nagao M, Higashino H, Hosokawa K, Kido T, Kurata A, Okayama H, Mochizuki T, Murase K (2010) Peri-infarct dysfunction in post-myocardial infarction: assessment of 3-T tagged and late enhancement MRI. Eur Radiol 20:1139–1148
26. Gilson WD, Yang Z, French BA, Epstein FH (2004) Measurement of myocardial mechanics in mice before and after infarction using multislice displacement-encoded MRI with 3D motion encoding. Am J Physiol Heart Circ Physiol 288:H1491–H1497
27. Spottiswoode BS, Zhong X, Lorenz CH, Mayosi BM, Meinjes EM, Epstein FH (2008) 3D myocardial tissue tracking with slice followed cine DENSE MRI. J Magn Reson Imaging 27:1019–1027
28. Zhong X, Spottiswoode BS, Meyer CH, Kramer CM, Epstein FH (2010) Imaging three-dimensional myocardial mechanics using navigator-gated volumetric spiral cine DENSE MRI. Magn Reson Med 64:1089–1097