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Myocardial SPECT perfusion defect size compared to infarct size by delayed gadolinium-enhanced magnetic resonance imaging in patients with acute or chronic infarction

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Summary

Background: Single photon emission computed tomography (SPECT) perfusion imaging has been considered a reference method for non-invasive estimation of infarct size in man. Recently, delayed gadolinium-enhanced magnetic resonance imaging (DE-MRI) has evolved as an accurate tool to quantify infarct size. Therefore, the present study was designed to compare perfusion defect size by SPECT to hyperenhanced volume by DE-MRI.

Methods: DE-MRI was performed in 30 patients. Fourteen were patients with revascularized first-time acute infarctions, eight revascularized chronic infarctions, and eight clinically referred non-revascularized patients. SPECT was performed in the same patients and analysed by a commercial package.

Results: The hypoperfused volume by SPECT was larger than the hyperenhanced volume by DE-MRI by 8 ± 8 ml (6% ± 5 percentage points), 10 ± 18 ml (6% ± 11 percentage points), and 26 ± 30 ml (12% ± 10 percentage points) in the acute, chronic and clinical populations, respectively. Left ventricle wall volume was smaller by SPECT in all settings.

Conclusion: The SPECT perfusion defect size was comparable with but generally slightly larger than the hyperenhanced volume by DE-MRI in both absolute and relative terms in patients with acute and chronic infarction. The results may be related to systematic differences between modalities but could also be influenced by biological phenomena such as wall thinning or hypoperfused but viable myocardium.

Introduction

The size of irreversible injury following an acute myocardial infarction has been shown to influence function and remodeling of the left ventricle (LV) (Field et al., 1972; Warren et al., 1988) and has also proved to be an important predictor of mortality (Braunwald, 1989; Miller et al., 1995). Therapy, such as thrombolysis or percutaneous transluminal coronary angioplasty (PTCA) that leads to reperfusion, limits infarct size (Braunwald, 1987, 1989; Gibbons et al., 1994). To verify successful treatment in the acute setting, a reliable method for assessing the presence, extent, and location of infarcted myocardium is therefore of clinical importance (Gibbons et al., 2000; Lund et al., 2001). This is also the case in chronic myocardial infarction when determining appropriate treatment and monitoring outcome. Several approaches in the clinical setting can be used to directly or indirectly monitor treatment and infarct size, such as ECG (Hinohora et al., 1988), biochemical markers of injury (Licka et al., 2002), echocardiography (Shen et al., 1991), metabolic activity by positron emission tomography (PET) (Charoenthaitawee et al., 2002), or relative distribution of perfusion by myocardial single photon emission computed tomography (SPECT) perfusion imaging (Gibbons et al., 2000). Scintigraphy using ⁴¹⁰Tl- or ⁹⁹mTc-labelled perfusion agents such as sestamibi and tetrofosmin has often been used as a reference method to estimate infarct size (Miller et al., 1995; Gibbons et al., 2000) and to monitor invasive therapy (van der Wall et al., 1990; Behrenbeck et al., 1991; Gibbons et al., 2000). Delayed gadolinium-enhanced magnetic resonance imaging (DE-MRI) has emerged as a useful method to image myocardial infarction and has in animal studies been shown to correlate well to histologically determined infarct size using triphenyltetrazolium chloride (TTC) staining (Kim et al., 1999; Fieno et al., 2000). Comparison of SPECT and DE-MRI for detection of non-transmural infarcts has shown that SPECT and DE-MRI detect transmural infarcts at similar rates although...
non-transmural infarcts detected by DE-MRI may be missed by SPECT (Wagner et al., 2003).

Therefore, the present study was designed to compare perfusion defect size by SPECT to the hyperenhanced volume by DE-MRI, in revascularized acute and chronic myocardial infarction, as well as in a non-revascularized population with chronic infarction.

**Methods**

All study patients had undergone myocardial infarction based on standard criteria as listed in the ACC/AHA guidelines for the management of patients with ST-elevation myocardial infarction (http://www.acc.org/clinical/guidelines/stemi/). The acute and chronic populations were revascularized and therefore less likely to have hypoperfused but viable myocardium. The clinical population was chosen as a non-revascularized group.

**Acute study population**

Fourteen patients (13 men, mean age 61 years, range 42–78 years) with no earlier evidence of myocardial infarction, presenting with acute ischaemia due to an occluded coronary artery, were included prospectively in the study. They underwent primary PTCA resulting in TIMI 3 flow in the opened artery. All patients received GPIIb/IIIa inhibitor during PTCA and were transferred to the coronary care unit for conventional therapy.

**Chronic study population**

Eight patients (eight men, mean age 65 years, range 57–76 years) with LV ejection fraction of <50% by echocardiography who underwent first-time elective coronary artery bypass grafting (CABG) without simultaneous valve repair were included prospectively.

**Clinical study population**

Eight clinically referred patients (seven men, mean age 68 years, range 57–74 years) who had underwent SPECT perfusion imaging were included and scheduled for DE-MRI.

**Study protocol**

The protocol and procedures were approved by the research ethics committee at Lund University Hospital, Sweden. Subjects were included after written informed consent. SPECT and DE-MRI were performed to determine the perfusion defect size and hyperenhanced volume, respectively.

The MRI was performed 6–9 days (median 7·5) after PTCA, and 28–38 days (median 32·5) after CABG in the acute and chronic population, respectively. SPECT was performed 0–5 days (median 0) and 0–1 day (median 0) after MRI in the acute and chronic population, respectively. MRI of the clinical population was performed 0–63 days (median 3·5) after SPECT. All patients were considered to be adequately medicated and were free from chest pain at time of imaging. The size of the perfusion defect measured by SPECT was compared with the hyperenhanced volume measured by DE-MRI.

**SPECT imaging**

Depending on body weight, 500–700 MBq $^{99m}$Tc-labelled tetrofosmin (Amersham Health, Buckinghamshire, UK) was injected at least 30 min before SPECT. The acquisition was performed within 3 h from isotope injection according to the standard clinical protocol at rest using a dual-head camera (ADAC, Milpitas, CA, USA). The subject was placed in supine position and imaged in steps of 5·6 degrees using a $64 \times 64$ matrix with a pixel size of 5·02 mm. Image acquisition time was approximately 25 min.

**SPECT image analysis**

Short- and long-axis images gated to ECG were reconstructed, covering the LV (Fig. 1a). This was done using a commercial application (AutoSPECT+Instill™ 6·0, ADAC) with an iterative method (maximum-likelihood expectation-maximization), using 12 iterations and a Butterworth filter with a cut-off value of 0·6 and an order of 5·0. Analysis of SPECT perfusion size is subject to a variety of technical and interpretation issues (Kojima et al., 1989; King et al., 1991; Ceriani et al., 1996; Issner & Patterson, 1999). We therefore chose to evaluate SPECT data by a validated (Germano et al., 2000; Sharir et al., 2000) and widely used commercial package (AutoQUANT™ 4·3·1 and a standard database; ADAC), in line with clinical practice, rather than to elaborate on the analysis details. LV wall volume (LVWV) and perfusion defect size were quantified in millilitre.

**Magnetic resonance imaging**

Either a 1·5 T system (Magnetom Vision, Siemens, Erlangen, Germany) with a CP body array coil or a 1·5 T system (Philips Intera CV, Philips, Best, the Netherlands) with a cardiac synergy coil was used. The subject was placed in supine position. Short- and long-axis images were acquired, covering the LV. The gradient-recalled echo (GRE) cine sequence and the segmented inversion-recovery (IR) GRE sequence were triggered by ECG and images were acquired during breath hold of approximately 15 s. The short-axis GRE cine images (slice thickness 8 mm, slice gap 2 mm) were used to measure the volume of the LV wall. The delayed gadolinium-enhanced IR-GRE images (slice thickness 8 mm, slice gap 2 mm) were used to measure the hyperenhanced volume. DE-MRI using the IR-GRE sequence utilizes an increase in regional fractional distribution of extracellular MRI contrast media within the injured area (Tong et al., 1993; Saeed et al., 1994; Arheden et al., 1999; Flacke et al., 2001) to achieve contrast between viable and non-viable...
myocardium (Simonetti et al., 2001). A commercially available gadolinium-based contrast agent (acute and clinical populations: gadoteric acid, Gd-DOTA, Guerbet, Gothia Medical AB, Bildal, Sweden; chronic and clinical populations: gadopentetate dimeglumine, Gd-DTPA, Schering Nordiska AB, Järfalla, Sweden) was administered at a dose of 0.2 mmol per kilogram of body weight, and delayed gadolinium-enhanced images were acquired in the same views as those used for the GRE cine protocol. Images were acquired during end diastole 26 ± 8 (SD) min after gadolinium administration. The inversion time (TI) was adjusted to give null signal from the myocardium, typically giving a TI of 170 ms (range 150–210) or 270 ms (range 260–280) for the Siemens or Philips system, respectively, because of different imaging sequence construction between systems. Image acquisition time was approximately 40 min.

Figure 1 Images from a patient with an acute infarction in the inferior wall, showing short axis slices from apex to base by (a) SPECT. Corresponding images by (b) cine MRI, and (c) DE-MRI. Arrowheads indicate infarcted area (white). LV, left ventricle; RV, right ventricle.

MR image analysis

Analysis of MR images was performed blinded to other results and patient information. All data were analysed using the program ImageJ ver 1.29x (http://rsb.info.nih.gov/ij/). In order to estimate interobserver variability, a subset of six patients was analysed by two observers. LVWV and hyperenhanced volumes were determined in end-diastolic (ED) images. Endocardial and epicardial borders were outlined manually on each ED and end-systolic frame in each short axis stack (Fig. 1b). Papillary muscles were included as LVWV. In the base of the LV where the left atrium was seen, only the portion of the slice that could be identified as LV was included. Hyperenhanced volume was outlined on the delayed contrast enhanced IR-GRE images throughout the LV (Fig. 1c). The long-axis images were used to verify the distribution of the hyperenhanced region. Total
myocardial and hyperenhanced volume were calculated as the planimetric measurements of each slice multiplied by slice thickness and slice gap. Relative hyperenhanced volume was expressed as percentage of the MRI LVWV.

**Statistical analysis**

Values were expressed as mean ± SD. Bland–Altman plots (Bland & Altman, 1986) were used to show differences between SPECT and MRI. A Wilcoxon signed-rank test was performed and P<0.05 was considered to show significant difference.

**Results**

**Acute study population**

Hyperenhanced volume by DE-MRI ranged from 0 to 100 ml involving up to 39% of the LVWV. LVWV was smaller by SPECT compared with MRI (23 ± 22 ml; P = 0.001; Fig. 2a). The hypoperfused volume by SPECT was in absolute and relative terms larger than the hyperenhanced volume by DE-MRI (8 ± 8 ml; P = 0.002; Fig. 2b; 6% ± 5 percentage points; P = 0.001; Fig. 2c).

**Chronic study population**

Hyperenhanced volume by DE-MRI ranged from 3 to 40 ml involving up to 17% of the LVWV. LVWV was smaller by SPECT compared with MRI (36 ± 26 ml; P = 0.016; Fig. 2a). The hypoperfused volume by SPECT was in absolute and relative terms larger than the hyperenhanced volume by DE-MRI (10 ± 18 ml; P = 0.344; Fig. 2b; 6% ± 11 percentage points; P = 0.344; Fig. 2c).

**Clinical study population**

Hyperenhanced volume by DE-MRI ranged from 14 to 53 ml involving up to 29% of the LVWV. Mean LVWV did not differ between SPECT and MRI (0 ± 46 ml; P = 0.438; Fig. 2a). The hypoperfused volume by SPECT was in absolute and relative terms larger than the hyperenhanced volume by DE-MRI (26 ± 30 ml; P = 0.047; Fig. 2b; 12% ± 10 percentage points; P = 0.023; Fig. 2c).

**Interobserver variability**

In the present MRI material interobserver variability was found to be 1 ± 4 ml (0 ± 3%) for LVWV and 0 ± 1 ml (1 ± 3%) for infarct size.

**Discussion**

The results show that the hypoperfused volume by myocardial SPECT perfusion imaging was generally slightly larger than the hyperenhanced volume by delayed contrast enhanced MRI. In addition, LVWV by SPECT was smaller compared with MRI in man in both the acute and chronic setting.

The results of the present study in patients are consistent with previous results comparing SPECT and explanted hearts showing that the relative hypoperfused volume measured by \textit{ex vivo} SPECT is slightly larger than \textit{ex vivo} measurements of relative infarct size by TTC (Medrano et al., 1996). SPECT has also been shown to underestimate the amount of viable myocardium when compared with PET (Arrighi et al., 1997). As SPECT depicts a perfusion defect, it is possible that not only the infarcted area but also a border zone of hypoperfused but viable myocardium is included. This border zone, previously described experimentally (Reimer et al., 1977), may explain the discrepancy between perfusion defect size and hypoperfused volume by DE-MRI. In addition, wall thinning by itself, in absence as well as in presence of infarction, may be detected as a perfusion defect by SPECT (Smith et al., 1997; Eisner & Patterson, 1999), giving rise to a larger perfusion defect size compared with hypoperfused volume by DE-MRI. In the acute study population, no thinning but however stunning (Braunwald & Kloner, 1982; Kloner et al., 2001) and thereby normally perfused but hypofunctioning myocardium may be present which possibly may be depicted as a perfusion defect because of absence of thickening-related count increase. Furthermore, systematic differences between SPECT and MRI may depend on technical issues such as temporal and spatial resolution.

In cases where the perfusion defect size by SPECT was smaller than the hypoperfused volume by DE-MRI (Fig. 2b), this could for example be because of the presence of non-transmural infarctions with otherwise well perfused epicardial regions, resulting in possible underdetection and underestimation of the hypoperfused volume by SPECT (Elkamhawy & Chandna, 2001; Wagner et al., 2003). However, in the current study not only non-transmural but also transmural infarctions were in some cases found smaller by SPECT compared with DE-MRI.

It has been proposed that the accuracy of the algorithms used for determination of the SPECT perfusion defect size may depend on myocardial wall thickness which may be of specific importance in cases of larger perfusion defects giving rise to inconsistencies in perfusion defect detection (Eisner & Patterson, 1999).

A smaller LVWV by SPECT than by MRI is likely because of an underestimation on behalf of SPECT rather than overestimation by MRI, since previous findings show that MRI of the LVWV correlates well with autopsy findings (Caputo et al., 1987; Lund et al., 2000).

**Limitations of the study**

Patients in the clinical population who have not undergone revascularization treatment may have hypoperfused but viable
Figure 2  (a) Agreement (left) of LVWV between SPECT and MRI in the acute (●), chronic (○), and clinical (×) populations. Bland–Altman plot (right) showing differences between SPECT and MRI. (b) Corresponding data and analysis for perfusion defect size by SPECT and hyperenhanced volume by DE-MRI. (c) Corresponding data and analysis for relative perfusion defect by SPECT and relative hyperenhanced volume by DE-MRI. LVWV, left ventricle wall volume. Dashed line indicates line of equality.
myocardium present, so called hibernating myocardium. This may give rise to a much larger perfusion defect size by SPECT compared with hyperenhanced volume by DE-MRI. However, to address this issue, we studied revascularized patients in the acute and chronic populations.

Conclusion

The present study has shown a better agreement between SPECT and DE-MRI to estimate infarct size in revascularized myocardial infarction compared with non-revascularized infarction. The perfusion defect size by SPECT is generally slightly larger than the hyperenhanced volume by DE-MRI in both absolute and relative terms. The results are likely related to systematic differences between modalities but could also be influenced by biological phenomena such as wall thinning or hyperperfused but viable myocardium.

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