Comparison Study on Different Quantification Methods of Diffuse Myocardial Fibrosis of Dilated Cardiomyopathy Using Myocardial T1 Value

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ABSTRACT

Purpose: The purpose is to compare several quantification methods and clarify which quantification method is reliable to estimate diffuse myocardial fibrosis with cardiac MRI in patients with dilated cardiomyopathy (DCM) using myocardial T1 value.

Methods and Results: Delayed enhancement imaging was performed in 52 patients with DCM and 10 control subjects to identify fibrosis using an inversion time scout sequence. The mean contrast-enhanced myocardial (M) T1 values of the pre and post contrast-enhanced myocardial and left ventricular lumen (L) of control and dilated cardiomyopathy cases were compared. The calculated post M T1 value, pre M T1 value-post M T1 value, and (pre M T1 value-post M T1 value)/(pre L T1 value-post L T1 value) were significantly different between the patient group and the control group (344.5 ± 31.6 vs. 390.4 ± 19.3 msec, 239.9 ± 64.2 msec vs. 134.0 ± 28.9 msec, and 0.50 ± 0.11 vs. 0.30 ± 0.60, respectively). (Pre M T1 value-post M T1 value)/(pre L T1 value-post L T1 value) was significantly the most related to the left ventricular ejection fraction (r = 0.66, p < 0.0001).

Conclusion: (Pre M T1 value-post M T1 value)/(pre L T1 value-post L T1 value) was the most reliable quantification method to estimate the severity of DCM.

Keywords: Inversion Time; Quantification Methods; Myocardial Fibrosis; Dilated Cardiomyopathy

1. Introduction

Recently, the importance of myocardial fibrosis in the development and progression of systolic and diastolic cardiac failure has been highlighted in cardiomyopathy [1-3]. Central to the diagnostic utility of cardiovascular magnetic resonance is its almost unique capacity to reveal myocardial fibrosis through the use of delayed gadolinium enhancement imaging [4].

In patients with cardiac infarction, infarcted regions in the myocardium, having undergone scar formation with collagen deposition, have a much slower washout rate of gadolinium-based contrast material than healthy myocardium, leading to markedly increased signal intensity on T1-weighted imaging [5]. On the other hand, because 60% of dilated cardiomyopathy (DCM) patients have diffuse myocardial fibrosis, the technique of delayed contrast enhancement often shows no regional scarring [5]. Previous study suggested that the mean contrast-enhanced myocardial inversion time at the null point (T1 value) was significantly shorter in a patient group than in a control group [5]. T1 value can be easily obtained, but this technique does not consider the influence of the contrast medium in the blood pool, and thus may not evaluate the enhancement of myocardium precisely.

The purpose of this study was to compare several quantification methods and clarify which quantification method is reliable to evaluate diffuse myocardial fibrosis of DCM using T1 value.

2. Materials and Methods

2.1. Patients

Fifty-two patients (33 men, 19 women; mean age ± SD, 53.3 ± 13.6 y; age range, 18 - 77 y) with DCM and 10 control subjects (5 men, 5 women; mean age ± SD, 54.8 ± 12.4 y; age range, 55 - 74 y) underwent MRI at our institution between June 2008 and July 2011. There was no statistically significant difference in mean age (P = 0.763) between the two groups. The diagnosis of non-ischemic DCM was made according to World Health Organization and International Society and Federation of...
Cardiology criteria [6]. None of the subjects had clinical symptoms or signs of ongoing myocarditis. By the use of coronary angiography, cases of significant coronary artery disease (>50% diameter luminal stenosis in any coronary artery) were excluded from this study. Other exclusion criteria were the presence of any contraindications for cardiac MRI, significant valvular disease, or any evidence of infiltrative heart disease by clinical symptoms or signs and sonographic findings. In addition, all patients with DCM underwent myocardial biopsy. The specimens showed disarray, with varying degrees of interstitial fibrosis of the myocardium, which were consistent with DCM.

On the other hand, all control subjects (n = 10) underwent cardiac MRI because of arrhythmia; however, no subjects had abnormalities according to coronary angiography, laboratory testing, echocardiography, nuclear medicine scan, or myocardial biopsy, which were performed to rule out various myocardial diseases.

All subjects gave written informed consent, and the protocol was approved by a medical ethics committee.

2.2. Cardiac MRI

All subjects were studied in the supine position using a 1.5-T cardiac MRI system (Avanto, Siemens Healthcare) with an eight-channel body coil. The data acquisition was blinded to diagnosis. The cardiac MRI study included cine steady-state free-precision imaging (TR/TE, 3.4/1.2; in-plane spatial resolution, 1.6 × 2 mm) of left ventricular function. Cine images were acquired with ECG gating and breath-holding. Cine images were obtained in 8-14 short axis (8-mm thickness with 0-mm spacing) from the base to the apex of the left ventricle. Vertical long axis images were also obtained. In all subjects (n = 62), inversion time scout images were obtained before enhancement and 15 minutes from the beginning of bolus injection of gadopentetate dimeglumine (0.2 mmol/kg; Magnevist, Bayer Schering Pharma) to identify regional fibrosis using a 2D inversion recovery gradient-echo technique (inversion time scout sequence: TR/TE, 20.8/1.3; flip angle, 25°; acquisition matrix, 192 × 78; field of view, 34 × 27 cm; slice thickness, 8 mm; inversion time, individually determined to null the myocardial and left ventricular lumen signal). Forty-two images per single midventricular slice were acquired at various inversion times during breath-holding as long as possible (Figures 1(a) and (b)). This single midventricular slice was selected on the basis of long axis scout images of the left ventricle before inversion time scout images were obtained. These inversion time scout images were then processed with a curve-fitting technique to generate T1 maps (Figures 1(c) and (d)). After the inversion time scout sequence was performed, ordinary delayed enhanced images were obtained.

2.3. Data Analysis

All MRI post-processing was performed by a single observer with more than 10 years of experience in cardiac MRI. The left ventricular ejection fraction was derived from cine images using a workstation for analysis (Syngo, Siemens Healthcare). A T1 mapping sequence was used to calculate the pre and post contrast-enhanced myocardial and left ventricular lumen T1 times of a single midventricular slice as an index of diffuse fibrosis. After image acquisition, short-axis images of varying inversion times were transferred to a workstation for analysis (Syngo, Siemens Healthcare). For each image, a region of interest was drawn in the left ventricular lumen and around the entire myocardium to calculate the pre and post contrast-enhanced myocardial and left ventricular lumen T10 values for each subject (Figures 1(a) and (b)). This allowed us to analyze regions of interest to find the average T1 time for that area, as well as a pixel-by-pixel determination of T1, by fitting data acquired at various preparation times to the exponential curve.

We then determined pre and post contrast-enhanced myocardial (M) and left ventricular lumen (L) T1 values (pre contrast-enhanced myocardial T1 values; post M T1 value, pre contrast-enhanced left ventricular lumen T1 value; pre L T1 value, post contrast-enhanced myocardial T1 value; post M T1 value, post contrast-enhanced left ventricular lumen T1 value; post L T1 value) [1,7,8] (Figures 1(c) and (d)). We compared the several quantification methods as follows: 1) post M T1 value, 2) pre M T1 value-post M T1 value, 3) post M T1 value/post L T1 value, and 4) (pre M T1 value-post M T1 value)/(pre L T1 value-post L T1 value).

2.4. Statistical Analysis

All values are expressed as the mean ± SD. Statistical analysis was performed on clinical and morphologic variables with the paired Student’s t test and Mann-Whitney U test for continuous variables. Pearson’s correlation coefficients were used to examine the correlation of left ventricular ejection fraction with all T10 values. Correlation coefficient values of 0.4 - 1.0 were considered to indicate a correlation [9]. In all tests, p < 0.05 was considered significant (SPSS, release 11.5; SPSS).

3. Results

The results are summarized in Table 1. In patients in the DCM group, the mean pre M T1 value, post M T1 value, pre L T1 value, and post L T1 value were 581.3 ± 58.3 msec, 344.5 ± 31.6 msec, 741.2 ± 92.5 msec, and 257.5 ± 46.1 msec, respectively. In the control group, the mean pre M T1 value, post M T1 value, pre L T1 value, and post L T1 value were 519.7 ± 30.4 msec, 390.4 ± 19.3 msec, 731.4 ± 64.4 msec, and 299.3 ± 35.5 msec.
respectively (Table 2). There were statistically significant differences between the patient and control groups in pre M T1 values, post M T1 values, and post L T1 values ($P = 0.0052$, $P < 0.0001$, $P = 0.002$, respectively). However, there was no significant difference between the patient and control groups in pre L T1 value ($P = 0.559$).

In patients in the DCM group, post M T1 value, pre M T1 value-post M T1 value, post M T1 value/post L T1 value, and (pre M T1 value-post M T1 value)/(pre L T1 value-post L T1 value) were $344.5 \pm 31.6$ msec, $239.9 \pm 64.2$ msec, $1.37 \pm 0.21$, and $0.50 \pm 0.11$, respectively. In the control group, post M T1 value, pre M T1 value-post M T1 value, post M T1 value/post T1 value, and (pre M T1 value-post M T1 value)/(pre L T1 value-post L T1 value) were $390.4 \pm 19.3$ msec, $134.0 \pm 28.9$ msec, $1.31 \pm 0.13$, and $0.30 \pm 0.60$, respectively (Table 3). There were statistically significant differences between the patient and control groups in post M T1 value, pre M T1 value-post M T1 value, and (pre M T1 value-post M T1 value)/(pre L T1 value-post L T1 value) ($P < 0.0001$, $P < 0.0001$, $P < 0.0001$, respectively). However, there was no significant difference between the patient and control
Table 1. Summary of results in patient and control groups.

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Control Subjects (n = 10)</th>
<th>Patients (n = 52)</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (Y)</td>
<td>54.8 ± 12.4</td>
<td>53.3 ± 13.6</td>
<td>NS</td>
</tr>
<tr>
<td>Sex (no.)</td>
<td></td>
<td></td>
<td>NS</td>
</tr>
<tr>
<td>Male</td>
<td>5</td>
<td>33</td>
<td></td>
</tr>
<tr>
<td>Female</td>
<td>5</td>
<td>19</td>
<td></td>
</tr>
<tr>
<td>LVEF (%)</td>
<td>62.5 ± 7.0</td>
<td>37.2 ± 14.1</td>
<td>&lt;0.0001</td>
</tr>
</tbody>
</table>

Note: LVEF = Left ventricular ejection fraction, NS = not significant.

Table 2. Comparison of pre and post contrast-enhanced myocardial and left ventricular lumen T1 values between control and DCM groups.

<table>
<thead>
<tr>
<th></th>
<th>Control</th>
<th>DCM</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pre M T1 value (msec)</td>
<td>519.7 ± 30.4</td>
<td>581.3 ± 58.3</td>
<td>0.005</td>
</tr>
<tr>
<td>Post M T1 value (msec)</td>
<td>390.4 ± 19.3</td>
<td>344.5 ± 31.6</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Pre L T1 value (msec)</td>
<td>731.4 ± 64.4</td>
<td>741.2 ± 92.5</td>
<td>NS</td>
</tr>
<tr>
<td>Post L T1 value (msec)</td>
<td>299.3 ± 35.5</td>
<td>257.5 ± 46.1</td>
<td>0.002</td>
</tr>
</tbody>
</table>

Note: NS = not significant.

Table 3. Comparison of four quantification methods of T1 value between control and DCM groups.

<table>
<thead>
<tr>
<th></th>
<th>Control</th>
<th>DCM</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Post M T1 value (msec)</td>
<td>390.4 ± 19.3</td>
<td>344.5 ± 31.6</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Pre M T1 value - post M T1 value (msec)</td>
<td>134.0 ± 28.9</td>
<td>239.9 ± 64.2</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Post M T1 value/post L T1 value</td>
<td>1.31 ± 0.13</td>
<td>1.37 ± 0.21</td>
<td>0.3422</td>
</tr>
<tr>
<td>(pre M T1 value - post M T1 value)/(pre L T1 value - post L T1 value)</td>
<td>0.30 ± 0.60</td>
<td>0.50 ± 0.11</td>
<td>&lt;0.0001</td>
</tr>
</tbody>
</table>

Note: M = myocardial, L = left ventricular lumen, DCM = dilated cardiomyopathy.

In the DCM groups, (pre M T1 value-post M T1 value)/(pre L T1 value-post L T1 value) was the most significantly related to the left ventricular ejection fraction ($r = 0.66; P < 0.0001$) (Figure 2). The other quantification methods (post M T1 value, pre M T1 value-post M T1 value, and post M T1 value/post L T1 value) had smaller positive correlations than (pre M T1 value-post M T1 value)/(pre L T1 value-post L T1 value) with the left ventricular ejection fraction (post M T1 value; $r = 0.53; P < 0.0001$) (Figure 3), pre M T1 value-post M T1 value ($r = -0.47; P < 0.0001$) (Figure 4), and post M T1 value/post L T1 value ($r = 0.10; P = 0.43$) (Figure 5).

Figure 2. Scatterplot shows correlation between left ventricular ejection fraction and (pre M T1 value-post M T1 value)/(pre L T1 value-post L T1 value) in patients with DCM. (Pre M T1 value-post M T1 value)/(pre L T1 value-post L T1 value) was most significantly related to left ventricular ejection fraction ($r = 0.66; P < 0.0001$).

Figure 3. Scatterplot shows correlation between left ventricular ejection fraction and post M T1 value in patients with DCM. Post M T1 value was significantly related to left ventricular ejection fraction ($r = 0.53; P < 0.0001$).

Figure 4. Scatterplot shows correlation between left ventricular ejection fraction and pre M T1 value-post M T1 value in patients with DCM. Pre M T1 value-post M T1 value was significantly related to left ventricular ejection fraction ($r = -0.47; P < 0.0001$).
4. Discussion

In patients with DCM, an important mechanism for the occurrence of arrhythmias and failure to respond to treatment is the presence of myocardial fibrosis [10-13]. Therefore, in patients with DCM, fibrosis is one of the most important risk factors for mortality.

Delayed enhanced MRI has enabled depiction of myocardial damage with high spatial resolution in various myocardial diseases. Delayed enhanced cardiac MRI has been used to evaluate focal myocardial fibrosis, but it is difficult to evaluate diffuse fibrosis of the myocardium. Prior studies have identified pathologic findings of myocardial fibrosis and disarray can show enhancement on delayed enhanced images, but delayed enhancement related to disarray is usually faint. Measurement of contrast-enhanced myocardial T1 value can provide an estimate of diffuse fibrosis of the myocardium [1,5,7]. Therefore, the prior studies used contrast-enhanced myocardial T1 value, which is easily calculated on a workstation [14-16]. In addition, T1 value was used as the optimal inversion time when delayed enhanced MR images were acquired.

According to previous study, the T1 values were 273 - 420 msec, 15 minutes after administration of contrast material in patients with DCM [5]. This quantification method has been postulated to be useful as this contrast-enhanced myocardial T1 value was significantly shorter in the patient group than in the control group. However, this quantification method does not consider the influence of the contrast medium of the blood pool and thus this quantification method might not evaluate the enhancement of myocardium. Another prior study identified quantification methods such as post M T1 value/post L T1 value, which considered the influence of the contrast medium of the blood pool [17], but this quantification method did not measure the difference between post M T1 value and pre M T1 value, or between post L T1 value and pre L T1 value. Therefore, this quantification method might not evaluate the enhancement of myocardium.

To our knowledge, this is the first study to determine pre and post contrast-enhanced myocardial and left ventricular lumen T1 values. This study compared several quantification methods.

In this study, post M T1 value, pre M T1 value-post M T1 value, and (pre M T1 value) showed statistically significant differences between patient and control groups. These methods could be useful diagnostic tools for DCM. However, because the quantification method of post M T1 value did not measure a difference between post M T1 value and pre M T1 value, this quantification method may not evaluate the enhancement of myocardium precisely. Moreover, because this quantification method may not measure the difference between post L T1 value and pre L T1 value, this quantification method did not consider the influence of the contrast medium of the blood pool.

The quantification method of post M T1 value/post L T1 value did not show a significant difference between the patient and control groups, although this quantification method considers the influence of the contrast medium of the blood pool, this quantification method may not evaluate the enhancement of myocardium precisely.

The quantification method of post M T1 value/post L T1 value did not show a significant difference between the patient and control groups, although this quantification method considers the influence of the contrast medium of the blood pool, this method may evaluate the enhancement of myocardium most precisely among the four methods.

In prior studies, left ventricular ejection fraction was suggested to be associated with serious clinical symptoms [18-20]. According to our results, (pre M T1 value-post M T1 value)/(pre L T1 value-post L T1 value) was the most significantly related to the left ventricular ejection fraction \((r = 0.66; P < 0.0001)\) in the DCM group. Our study shows that the quantification method of (pre M T1 value-post M T1 value)/(pre L T1 value-post L T1 value) considers the influence of the contrast medium of the blood pool and evaluates the enhancement of myocardium. Therefore, this method may be the most reliable diagnostic method to evaluate the extent of myocardial fibrosis.

4. Discussion

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To our knowledge, this is the first study to determine pre and post contrast-enhanced myocardial and left ventricular lumen T1 values. This study compared several quantification methods.

In this study, post M T1 value, pre M T1 value-post M T1 value, and (pre M T1 value-post M T1 value) showed statistically significant differences between patient and control groups. These methods could be useful diagnostic tools for DCM. However, because the quantification method of post M T1 value did not measure a difference between post M T1 value and pre M T1 value, this quantification method may not evaluate the enhancement of myocardium precisely. Moreover, because this quantification method may not measure the difference between post L T1 value and pre L T1 value, this quantification method did not consider the influence of the contrast medium of the blood pool.

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The quantification method of (pre M T1 value-post M T1 value)/(pre L T1 value-post L T1 value) measured the difference between post M T1 value and pre M T1 value, and between post L T1 value and pre L T1 value. Because this quantification method did not consider the influence of the contrast medium of the blood pool, this method may evaluate the enhancement of myocardium most precisely among the four methods.

In prior studies, left ventricular ejection fraction was suggested to be associated with serious clinical symptoms [18-20]. According to our results, (pre M T1 value-post M T1 value)/(pre L T1 value-post L T1 value) was the most significantly related to the left ventricular ejection fraction \((r = 0.66; P < 0.0001)\) in the DCM group. Our study shows that the quantification method of (pre M T1 value-post M T1 value)/(pre L T1 value-post L T1 value) considers the influence of the contrast medium of the blood pool and evaluates the enhancement of myocardium. Therefore, this method may be the most reliable diagnostic method to evaluate the extent of myocardial fibrosis. 

Figure 5. Scatterplot shows correlation between left ventricular ejection fraction and post M T1 value/post L T1 value in patients with DCM. Post M T1 value/post L T1 value was not significantly related to left ventricular ejection fraction \((r = 0.10; P = 0.43)\).
cardiac fibrosis non-invasively.

There were several limitations to this study. First, we used inversion times to measure a single slice. Ideally, measurement of the whole myocardium is needed to evaluate diffuse fibrosis. Therefore, further studies regarding markers of fibrosis are needed. In addition, it is not easy to draw a myocardial boundary. This process may potentially bias the results.

Second, this study lacked a comparison of severity of myocardial fibrosis, histologically. However, it may be impossible to correlate the areas of fibrosis on biopsy with the areas seen on MRI.

Third, we evaluated only four quantification methods and we must carry out further examination to find a more useful quantification method.

5. Conclusion

In conclusion, (pre M T1 value-post M T1 value)/(pre L T1 value-post L T1 value), which considered the influence of the contrast medium of the blood pool, evaluated the enhancement of myocardium precisely. This method was the most significantly related to the left ventricular ejection fraction. These data suggest that (pre M T1 value-post M T1 value)/(pre L T1 value-post L T1 value) is the most reliable quantification method to estimate the severity of DCM.

REFERENCES


