A Prominent R Wave in V1 but not in V2 Is a Specific Sign of a Large Lateral Transmural Infarction

Daniele Rovai, Gianluca Di Bella, Giuseppe Rossi, Alessandro Pingitore, and Antonio L’Abbate

Abstract

Introduction and objectives: In the absence of right ventricular hypertrophy or bundle-branch block, a prominent R wave in V1 or V2 is considered to reflect a lateral myocardial infarction. We investigated the differences in infarct location, size and transmural extent between patients with prominent R wave in V1 and those with prominent R wave in V2.

Methods: We studied 50 patients with a previous first infarction involving left ventricular inferior and/or lateral wall at contrast-enhanced magnetic resonance.

Results: A prominent R wave in V1 was present in 8 patients (16%), in V2 in 23 (46%). At magnetic resonance, the infarction involved the inferior wall in 11 patients (22%), the lateral wall in 6 (12%), and both walls in 33 patients (66%). The sensitivity of a prominent R wave in V1 in detecting a lateral infarction was low (17.9%), while the specificity was high (90.9%). The sensitivity and specificity of a prominent R wave in V2 were 46.2% and 54.5%, respectively. In patients with a prominent R wave in V1, infarct size and lateral and transmural extent were greater than in patients without this pattern (P<.005, <.001, and <.05, respectively); conversely, infarct size and transmural extent in the inferior wall and in its basal-posterior segment were not significantly different. In patients with a prominent R wave in V2, infarct size, lateral and transmural extent were not different from patients without this pattern.

Conclusions: Only a prominent R wave in V1 is a specific sign of large and transmural lateral infarction.

Palabras clave: Electrocardiografía, Infarto de miocardio, Resonancia magnética, Exploraciones de imagen, Enfermedad coronaria

Resumen

Introducción y objetivos: Si no hay hipertrofia ventricular derecha o bloqueo de rama del haz, se considera que la presencia de una onda R prominente en V1 o V2 refleja un infarto de miocardio de la pared lateral. Hemos investigado las diferencias existentes en cuanto a localización, tamaño y extensión transmural del infarto entre los pacientes con una onda R prominente en V1 y los que presentan una onda R prominente en V2.

Métodos: Estudiámos a 50 pacientes con un primer infarto previo que había afectado a la pared inferior y/o lateral del ventrículo izquierdo utilizando resonancia magnética con contraste.

Resultados: Se observó la presencia de una onda R prominente en V1 en 8 pacientes (16%) y en V2 en 23 pacientes (46%). En las imágenes de resonancia magnética, el infarto afectaba a la pared inferior en 11 pacientes (22%), la pared lateral en 6 (12%) y ambas en 33 (66%). La sensibilidad de la presencia de una onda R prominente en V1 para la detección de un infarto de cara lateral fue baja (17.9%), mientras que la especificidad fue alta (90.9%). La sensibilidad y la especificidad de una onda R prominente en V2 fueron del 46.2 y el 54.5% respectivamente. En los pacientes con una onda R prominente en V1, el tamaño del infarto y la extensión lateral y transmural fueron mayores que en los pacientes que no mostraban este patrón (p < 0.005, p < 0.001 y p < 0.05 respectivamente); en cambio, el tamaño del infarto y la extensión transmural en la pared inferior y en su segmento posterobasal no mostraron diferencias significativas. En los pacientes con una onda R prominente en V2, el tamaño del infarto y la extensión lateral y transmural no fueron diferentes de lo observado en los pacientes sin ese patrón.

Conclusiones: Tan sólo la presencia de una onda R prominente en V1 constituye un signo específico de un infarto lateral grande y transmural.

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The protocol consisted of cine MRI to evaluate global LV function and contrast-enhanced MRI to determine location, size, and transmural extent of MI. The MRI was performed using a 1.5 T whole-body scanner (GE Medical Systems, Milwaukee, WI). A four-element cardiac phased-array receiver surface coil was used for signal reception. We used a breath-hold, segmented-gradient, fast-imaging echo employing a steady-state acquisition ECG-triggered sequence to evaluate global LV function by standard parameters. In each patient a total of 9 to 12 short-axis views (depending on the LV volume) and 2 long-axis views (one vertical and one horizontal) were acquired, with a minimum of 30 cine frames for each slice. From 10 to 15 min after bolus injection of gadolinium- diethylenetriamine pentaacetic acid (Gadovist, Schering, Berlin, Germany; 0.2 mmol/kg), images were acquired at end-diastole in the same views. A fast-gradient echo inversion recovery sequence was used. The inversion time was optimized until the disappearance of the signal from the viable myocardium.

Table: Characteristics of Patients

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Value</th>
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</thead>
<tbody>
<tr>
<td>Number</td>
<td>50</td>
</tr>
<tr>
<td>Age, years</td>
<td>65 (11)</td>
</tr>
<tr>
<td>Male, %</td>
<td>90</td>
</tr>
<tr>
<td>Family history of CAD, %</td>
<td>37</td>
</tr>
<tr>
<td>Diabetes mellitus, %</td>
<td>39</td>
</tr>
<tr>
<td>Hypercholesterolemia, %</td>
<td>53</td>
</tr>
<tr>
<td>Hypertriglyceridemia, %</td>
<td>12</td>
</tr>
<tr>
<td>Arterial hypertension, %</td>
<td>56</td>
</tr>
<tr>
<td>Smoking habit, %</td>
<td>53</td>
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<tr>
<td>Obesity, %</td>
<td>30</td>
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<tr>
<td>No. of stenosed vessels</td>
<td>2 (0.9)</td>
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<tr>
<td>LVEDV, ml/m²</td>
<td>96 (30)</td>
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<tr>
<td>LVESV, ml/m²</td>
<td>54 (27)</td>
</tr>
<tr>
<td>LVEF, %</td>
<td>47 (13)</td>
</tr>
<tr>
<td>LV mass, g/m²</td>
<td>81 (22.5)</td>
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<tr>
<td>DCE Extent, % of entire LV</td>
<td>8 (4.4)</td>
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<tr>
<td>Segments DCE transmural extent 13-25%</td>
<td>1.5 (1.5)</td>
</tr>
<tr>
<td>Segments DCE transmural extent 26-50%</td>
<td>1.6 (1.6)</td>
</tr>
<tr>
<td>Segments DCE transmural extent 51-75%</td>
<td>1 (1.2)</td>
</tr>
<tr>
<td>Segments DCE transmural extent 76-100%</td>
<td>0.6 (1.1)</td>
</tr>
</tbody>
</table>

CAD, coronary artery disease; DCE, delayed contrast enhancement; LV, left ventricular; LVEDV, left ventricular end-diastolic volume; LVEF, left ventricular ejection fraction; LVESV, left ventricular end-systolic volume.
Magnetic Resonance Imaging Data Analysis

The LV endocardial borders were manually drawn on LV short-axis images corresponding to end-diastolic and end-systolic phases to calculate LV volumes and the ejection fraction. The LV myocardium was divided according to a 17-segment model. To assess location, size, and transmural extent of the MI, in each image the boundaries of contrast-enhanced areas were automatically traced (using a signal intensity cut-off of >5 SD over the average of normal remote myocardium), and manually corrected when needed. The reproducibility of this method has been previously validated.

To measure the extent of the MI, in each image the myocardium was automatically divided into equiangular sectors starting from the anterior septal insertion. Each sector was subdivided into 100 radiants, and the extent of contrast enhancement in each radiant was measured automatically (Fig. 1). Three to four contiguous slices were averaged to obtain the extent of the MI in each segment. The transmural extent of the MI in each segment was scored according to a 4-point scale, where 1 corresponds to <25% of LV myocardial thickness, 2 is between 26% and 50%, 3 between 51% and 75%, and 4 corresponds to an extent >75% of LV thickness. The infarct size was calculated as the sum of the scores of all 10 segments of the inferior and lateral walls. Infarct extent in the inferior or lateral wall was calculated as the sum of the scores in the 5 inferior or 5 lateral segments. Infarct transmural extent was calculated as the mean score in the segments with contrast enhanced myocardium.

Electrocardiogram

A 12-lead ECG was recorded at a speed of 25 mm/s and a voltage of 10 mm/mV. Two cardiologists, unaware of the MRI data, analyzed the ECG tracings off-line. In the case of disagreement regarding the ECG interpretation, a consensus was reached by reading the tracing together. The criteria for identification of the ECG patterns of necrosis were defined before the analysis. Q waves were considered pathological if they were >0.04 s in duration. The R wave in V1 or V2 was considered prominent if the R/S ratio was >1 and the R wave duration was >0.04 s. The patients were categorized on the basis of a prominent R wave in V1 or a prominent R wave in V2.

Statistical Analysis

Quantitative data are expressed as mean±1 SD, qualitative data as percentage. The ability of the different ECG patterns of necrosis to detect MI location was evaluated as sensitivity and specificity, according to their definition. The difference between MI size, horizontal extent, and transmural extent in patients with the different ECG patterns of necrosis was tested by the analysis of variance. All statistical tests were 2-tailed; a P-value<.05 was considered significant. Statistical analysis was performed with the JMP 9 software (SAS Institute Inc.).

RESULTS

The 12-lead ECG showed a prominent R wave in V1 in 8 patients (16%); all these patients also had a prominent R wave in V2. The ECG showed a prominent R wave in V2 but not in V1 in 15 patients (30%). Thus, 23 patients (46%) showed a prominent R wave in V2. Inferior Q waves of necrosis were present in 27 patients (54%). These Q waves were associated with a prominent R wave in V1 in 5/8 patients, with a prominent R wave only in V2 in 10/15 patients, and were not associated with a prominent R wave in 12 patients. At cardiac MRI, the infarction scar was located only in the inferior wall in 11 patients (22%), only in the lateral wall in 6 patients (12%), and in both inferior and lateral walls in 33 patients (66%).

Prominent R Wave in V1 or V2 and Location of Myocardial Infarction

Of the 8 patients with a prominent R wave in V1, 7 showed areas of delayed contrast enhancement in the lateral wall; specifically, the infarction scar was located only in the lateral wall in 2 patients, and in both inferior and lateral walls in 5. In the remaining patient with a prominent R wave in V1, a small subendocardial area of delayed contrast enhancement was located in the mid-inferior segment. Of the 42 patients without prominent R wave in V1, 32 showed areas of necrosis in the lateral wall, and 10 did not. Thus, the sensitivity of a prominent R wave in V1 in detecting a lateral MI was low (17.9%), while the specificity was high (90.9%). The positive and negative predictive values were 87.5% and 23.8%, respectively.

Of the 23 patients with prominent R wave in V2, 18 showed lateral infarction scars, and 5 did not. Of the 27 patients without this pattern, 21 showed evidence of lateral necrosis at MRI, and 6 did not. Thus, the sensitivity and the specificity of a prominent R wave in V2 in detecting lateral infarctions were 46.2% and 54.5%, respectively.

Prominent R Wave in V1, Infarct Size, and Transmural Extent

In patients who presented a prominent R wave in V1, the overall size of the infarction at contrast-enhanced MRI, the extent of the MI in the lateral wall, and infarct transmural extent were higher than in patients without this electrocardiographic pattern (Fig. 2).

Figure 1. Contrast-enhanced, short-axis image of the heart. The lines in the right panel correspond to left ventricular endocardial (yellow) and epicardial boundaries (red), and to the edges of contrast-enhanced area (green).
Conversely, infarct size in the inferior wall and in its basal segment (formerly called posterior) were not significantly different between patients with and without a prominent R wave in V1 (Fig. 3).

Prominent R Wave in V2, Infarct Size, and Transmural Extent

In patients with prominent R wave in V2, the overall infarct size and its lateral and transmural extent were not significantly different from those of patients without such a pattern (Fig. 4), nor were the size of the MI in the inferior wall (P=.348) or in its inferobasal segment (P=.187).

In the 20 patients with inferior Q waves, infarct size in the lateral wall was not significantly different (P=.419) from that of patients without Q waves, while MI extent in the inferior wall was significantly higher (P=.017).

DISCUSSION

This study shows that a prominent R wave in V1 and in V2 are markers of a lateral MI. However, a prominent R wave in V1 is a specific marker but not very sensitive, while a prominent R wave in V2 is a more sensitive but less specific marker. In practical terms, a prominent R wave in V1—in the absence of right ventricular hypertrophy or bundle branch block—is a diagnostic sign of a lateral infarction, while a prominent R wave in V2 may lead to misdiagnosis. Although these conclusions can be envisaged by previous studies, this is the first investigation that compares the different meanings of a prominent R wave in V1 and V2.

The relationship between an R wave in V1 and V2 and infarct location has been investigated in several studies. In a previous study we evaluated patients with Q-wave MI or equivalents. Thus, we selected patients based on electrocardiographic evidence of necrosis, while in this study we selected patients based on the MRI evidence of infarction. Furthermore, we previously included all possible infarct locations, while in the present study we selected only patients with inferior, lateral, or inferolateral MI. This study is also different from the previous ones performed by Bayes de Luna et al.,3–6 where the MRI data were categorized in a binary way (as presence or absence of a previous MI) while we have included a quantitative data analysis on infarct size and transmural extent.

According to the results of various studies, including the present one, the commonly utilized nomenclature of MI location should be reconsidered. Specifically, a prominent R wave in right precordial leads should be considered a sign of lateral MI, not of posterior MI. This is not a matter of semantics, since the LV lateral wall is generally perfused by the left circumflex coronary artery, often by the obtuse marginal branch, while LV inferobasal wall is usually perfused by the right coronary artery. This consideration should be taken into account when patients undergo coronary arteriography and myocardial revascularization. Finally, although this was not the end-point of the study, our results confirm that the Q-waves of necrosis in the inferior leads not only have no relationship with lateral MI, but correspond to a scar located in the inferior LV wall.

This study revealed the innovative information that a prominent R wave in V1 carries quantitative information on MI size and transmural extent. In fact, a prominent R wave was associated with a larger infarct size, while in the absence of this sigh the infarction was smaller and more confined to the subendocardial layers. In a study of 100 patients with previous MI, Moon et al. found the Q waves to be indicative of a large infarct; in addition, a strong relationship was found between classification as a Q-wave MI and quintiles of transmural extent of MI measured by contrast-enhanced MRI.13 In another study, although several parameters were predictors of Q waves at univariate analysis, multivariate analysis showed that quantified scar tissue extent was the single
**CONCLUSIONS**

A pathologic R wave in V1 is a specific marker of lateral infarction, and unmasks a large infarction scar, transmurally well extended. A prominent R wave in V2 is a more sensitive but less specific marker of lateral MI, and can induce a reasonable suspicion.

**FUNDING**

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**REFERENCES**


