Non-Invasive Magnetocardiography for the Early Diagnosis of Coronary Artery Disease in Patients Presenting With Acute Chest Pain

Hyukchan Kwon, PhD; Kiwoong Kim, PhD; Yong-Ho Lee, PhD; Jin-Mok Kim, PhD; Kwon Kyu Yu, BSc; Namsik Chung, MD, PhD; Young-Guk Ko, MD, PhD

Background: Accurate identification of patients with acute coronary syndrome (ACS) is often difficult especially when an electrocardiogram (ECG) does not show typical elevation of ST segment. The aim of the present study was therefore to evaluate the efficacy of magnetocardiography (MCG) for diagnosis of ACS in patients with acute chest pain presenting without ST segment elevation.

Methods and Results: In the present retrospective study 364 patients with the suspected ACS without ST segment elevation were selected. Significant coronary artery disease (CAD) was defined as a stenosis ≥50% in at least one of 16 segments of the 3 major coronary arteries and their branches. The MCG recordings were obtained at resting state using a 64-channel MCG system in a magnetically shielded room. The patients were classified on the basis of the probability distribution. The presence of significant CAD was identified with a sensitivity of 84.0% and a specificity of 85.0%, compared to 44.7% and 89.8% on ECG. In the subgroup of patients without specific findings on ECG or biomarker test, MCG had a sensitivity of 73.5% and a specificity of 82.3%.

Conclusions: MCG was acceptably sensitive and specific in identifying patients with ACS even in the absence of specific findings on ECG and positive biomarker tests. Thus, MCG seems beneficial for the early triage of patients with acute chest pain. (Circ J 2010; 74: 1424–1430)

Key Words: Acute coronary syndrome; Coronary artery disease; Magnetocardiography; Non-invasive detection

P atients presenting with acute chest pain without ST elevation on electrocardiogram are a diverse and heterogeneous group representing a wide spectrum of disease severity from acute myocardial infarction (MI) to non-ischemic chest pain syndromes. Despite development of various diagnostic approaches and modern treatments, the rates of mortality, MI and readmission in acute coronary syndrome (ACS) remain very high.1 In general, initial risk assessment in patients with acute chest pain is performed on the basis of the clinical, electrocardiographical and biochemical data to identify patients with acute myocardial ischemia.2 In high-risk patients, coronary angiography is required to confirm the clinical diagnosis, and subsequent coronary revascularization either on percutaneous coronary intervention or coronary artery bypass graft is considered, depending on the severity of coronary artery disease (CAD). Because of inherent risks and relatively high costs, however, the use of coronary angiography is restricted to patients judged to be at high risk for myocardial progression to MI or death.3 Thus, if magnetocardiography (MCG) can effectively identify patients with myocardial ischemia before coronary angiography, it will be a valuable non-invasive technique for risk stratification at an early stage of admission.

MCG has been proposed as a non-invasive and contact-free technique for functional diagnosis of the heart such as cardiac ischemia and arrhythmia.4–8 MCG records the magnetic field induced by the same bioelectric currents recorded on electrocardiography (ECG).8 During myocardial ischemia, ST segment and T wave changes often appear on ECG. Acute ischemia usually results in transient horizontal or downsloping ST segment depression and T wave flattening or inversions in ECG.9,10 Because these changes originate from the abnormal current flow in the myocardium during ventricular repolarization, magnetic field orientation correlated with the current flow was used for ischemia decision in resting11–16 and stress-testing MCG.17,18 In principle, MCG is more sensi-
tive to tangential currents than ECG. MCG is also sensitive to circular vortex currents not detectable on ECG, and it has been shown to contain information complementary to ECG. Thus MCG may show pathological deviations from the normal activation direction induced by, for example, myocardial ischemia, with better accuracy than ECG.

For CAD detection in the resting state, several studies have reported that MCG is superior to ECG in that it has a diagnostic accuracy of 60–90% in various patient populations. Comparing spatial distribution of QT dispersion, a sensitivity of 74% and a specificity of 80% were observed in 23 CAD patients and a control group, in which 13 out of 20 subjects in the control group were normal subjects. In comparison between patients with and without hemodynamic relevant stenosis, a sensitivity of 62.8% and a specificity of 61.3% were reported on analysis of the current density vector. A sensitivity and a specificity of >90% were recently reported in patients admitted to the intensive care unit, using four variables derived during ST–T interval.

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In a preliminary study we proposed a new classification method of MCG parameters depending on the severity of ischemia to improve diagnostic accuracy. In a preliminary study we proposed a new classification method of MCG parameters depending on the severity of ischemia to improve diagnostic accuracy. In a preliminary study we proposed a new classification method of MCG parameters depending on the severity of ischemia to improve diagnostic accuracy. In a preliminary study we proposed a new classification method of MCG parameters depending on the severity of ischemia to improve diagnostic accuracy.

In a preliminary study we proposed a new classification method for detection of CAD on the basis of different distribution of the MCG parameters between CAD patients and asymptomatic patients or healthy volunteers. The aim of the present study was to validate the method in a relatively large population as a useful tool for the early diagnosis of CAD (stenosis ≥50%) in patients with ACS without ST elevation.

### Table 1. Patient Characteristics

<table>
<thead>
<tr>
<th></th>
<th>CAD (n=237)</th>
<th>Minimal CAD (n=127)</th>
<th>Normal (n=89)</th>
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</thead>
<tbody>
<tr>
<td>Female, n (%)</td>
<td>80 (34)</td>
<td>74 (58)</td>
<td>42 (47)</td>
</tr>
<tr>
<td>Age (years)</td>
<td>61.3±10.3</td>
<td>55.0±10.9</td>
<td>24.5±5.2</td>
</tr>
<tr>
<td>Coronary status, n (%)</td>
<td>1-vessel disease 89 (38)</td>
<td>0 (0)</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>2-vessel disease 79 (33)</td>
<td>0 (0)</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>3-vessel disease 69 (29)</td>
<td>0 (0)</td>
<td>–</td>
</tr>
<tr>
<td>Abnormal ECG, n (%)</td>
<td>106 (45)</td>
<td>13 (10)</td>
<td>0 (0)</td>
</tr>
<tr>
<td>RWMA, n (%)</td>
<td>99 (42)</td>
<td>2 (2)</td>
<td>–</td>
</tr>
<tr>
<td>Disease</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Atypical chest pain, n (%)</td>
<td>18 (8)</td>
<td>99 (78)</td>
<td>–</td>
</tr>
<tr>
<td>Unstable angina, n (%)</td>
<td>128 (54)</td>
<td>28 (22)</td>
<td>–</td>
</tr>
<tr>
<td>MI, n (%)</td>
<td>91 (38)</td>
<td>0</td>
<td>–</td>
</tr>
<tr>
<td>Ejection fraction, (%)</td>
<td>59.4±12.4</td>
<td>67.7±7.4</td>
<td>–</td>
</tr>
<tr>
<td>Diabetes mellitus, n (%)</td>
<td>49 (21)</td>
<td>7 (6)</td>
<td>0 (0)</td>
</tr>
<tr>
<td>Hypertension, n (%)</td>
<td>75 (32)</td>
<td>36 (27)</td>
<td>0 (0)</td>
</tr>
</tbody>
</table>

CAD, coronary artery disease; ECG, electrocardiography; RWMA, regional wall motion abnormality; MI, myocardial infarction.

### Table 2. MCG Parameters

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>TT_CAMx</td>
<td>Maximum value of the main current angle in the ST period.</td>
</tr>
<tr>
<td>TT_CMD</td>
<td>Maximum value of a dynamic change of the strength of the main current vector within a time interval of 30ms in the ST period.</td>
</tr>
<tr>
<td>T_FMA</td>
<td>Orientation of magnetic field map at T_max.</td>
</tr>
<tr>
<td>R_FMA</td>
<td>Orientation of magnetic field map at R_peak.</td>
</tr>
</tbody>
</table>

MCG, magnetocardiography; TT, ST period; T, peak of T-wave; R, peak of R-wave.

### Methods

#### Subjects

The present study was performed with the approval of the institutional review board at Yonsei University Medical College. Written informed consent was obtained from all subjects. For this retrospective study, 364 consecutive patients, who were admitted to the cardiovascular center with suspected ACS between May 2004 and April 2006, were enrolled. Exclusion criteria were persistent ST elevation, suspected variant angina, any specific abnormalities of bundle branch block, atrial fibrillation, arrhythmia on ECG or left ventricular hypertrophy on echocardiography. Known CAD patients with old MI or previous revascularizations, and MI patients with insignificant coronary stenosis (<50%) were also excluded. Out of 364 patients, 329 patients underwent coronary angiography during the initial hospitalization. Abnormal ECG changes indicative of myocardial ischemia without persistent ST elevation such as ST depression, T inversion and pathologic Q waves were observed in 119 patients. On echocardiogram, regional wall motion abnormality was observed in 101 out of 317 patients. On biomarker tests, 91 patients had elevated troponin T level (>0.1 ng/ml) level or peak creatine kinase >2 times the upper limit of normal, indicating MI. Unstable angina (UA) was diagnosed in 156 patients. UA was defined as angina pectoris or an equivalent type of ischemic discomfort with ≥1 of the following 3 features and negative results of cardiac biomarker tests (such as troponin T and creatine kinase-MB): (1) prolonged (usually ≥20 min) angina occurring at rest; (2) new-onset (within 1 month) angina of Canadian Cardiovascular Society class ≥III severity; and (3) previously diagnosed angina that had become distinctly more frequent, longer in duration, or lower in threshold.

Based on the severity of CAD, patients were divided into 2 groups: those with stenosis ≥50% in at least 1 of 16 segments in the 3 major coronary arteries and their branches, termed the CAD group (n=237, mean age 61.3±10.3 years); and those with stenosis <50% at all 16 segments or who were discharged without undergoing coronary angiography, termed the minimal CAD group (n=127, mean age 55.0±10.9 years). In the CAD group 89 patients had single-vessel disease, 79 had 2-vessel disease, and the remaining 69 patients had 3-vessel disease. As a reference, the normal group (mean age 24.5±5.2 years) consisted of 89 healthy volunteers with no history of any cardiovascular disease and with normal ECG at rest. The clinical characteristics of all subjects are listed in Table 1. Some of the MCG data have been used in previous studies.

#### MCG Recording

In patients who underwent coronary angiography, MCG re-
cordings were obtained prior to coronary angiography. All patients were treated and stabilized according to the clinical presentation. All patients were chest pain free at the time of MCG recording. MCG was recorded in the resting state using a multichannel MCG system inside a magnetically shielded room. MCG measures cardiac magnetic fields tangential to the chest surface at 64 locations using 1st-order planar gradiometers in an area of 162 mm × 162 mm with a sensor interval of approximately 35 mm. The total recording time was <10 min. The MCG signals were digitally recorded for 30 s at the sampling rate of 500 Hz. After baseline correction, data were averaged using R-peaks. The averaged signals were analyzed after DC offset correction using a baseline during the P–Q interval.

**MCG Parameters**

Three MCG parameters were derived from the spatial distribution of the measured magnetic fields during the repolarization period (Table 2). Magnetic field map (MFM) angle at T\text{max}, T\text{FMA}, was measured in respect to the patient’s right–left line as shown in Figure 1. After the source current map was reconstructed from the spatial distribution of the cardiac fields using a distributed current source model and minimum norm estimation algorithm, angle and strength of the main current vector were calculated at each time instant during the ST period between the J-point and T\text{max}. Among

![Figure 1](image1.png)

**Figure 1.** (A) Magnetic field map and (B) current map, and angle notations used in the present study. Examples of field map angle (–56.8°) and current angle (46.2°), and maximum current moment (71.9 nA·m) are also shown.

![Figure 2](image2.png)

**Figure 2.** T\text{FMA}–R\text{FMA} (orientation of magnetic field map at T\text{max} relative to the orientation of magnetic field map at R\text{peak}) in the (A) CAD, (B) minimal CAD and (C) normal groups, respectively. CAD, coronary artery disease.
them, the maximum of the current angle was TT_CAMx and its dynamic change within a 30-ms interval was TT_CMD. The early time window of the ST period, however, when the main current vector could not be definitely evaluated from the MFM due to the small signals or circulating current pattern, was excluded from the analysis. Additionally, MFM angle at R_peak was used as a reference angle to reduce the effect of different electrical axis between individuals.

Five variables were therefore used for diagnosis: T_FMA, T_FMA–R_FMA, TT_CAMx, TT_CAMx–R_FMA and TT_CMD.

Algorithm of Weighted Maximum a Posteriori (wMAP)

In the wMAP method, the probability that a patient has CAD is represented as a function of 5 variables. The probability was calculated from the prior knowledge on the probability distribution, which was estimated from MCG data collected in 2004 (86 CAD patients, mean age 60.3±10.1 years; 59 minimal CAD patients, mean age 52.9±10.2 years; 66 healthy volunteers, mean age 22.8±3.3 years). As an example, Figure 2 shows the histogram of T_FMA–R_FMA (the orientation of MFM at T_max relative to the orientation of MFM at R_peak) in the CAD, minimal CAD and normal group, respectively.

Interpolating the histograms and normalizing by the total number of subjects in each group, a probability distribution function was estimated and it was defined as the conditional probability

$$P(x_j|g_i) = \frac{1}{\sum_k P(x_j|g_k) \cdot P(g_k)} 
$$

Figure 3. Conditional probability $P(g_i|x_j)$ that a subject belongs to the group $g_i$ (normal, thick; minimal CAD, gray; CAD, thin) given that he/she has the measured value $x_j$ for the variables (A) T_FMA (orientation of magnetic field map at T_max), (B) T_FMA–R_FMA (orientation of magnetic field map at T_max relative to the orientation of magnetic field map at R_peak), (C) TT_CAMx (maximum of the main current angle in the ST period), (D) TT_CAMx–R_FMA and (E) TT_CMD (maximum of a dynamic change of the strength of the main current vector within a time interval of 30ms in the ST period), respectively. Straight radius line, measured variable. For example, when $x_1$ (T_FMA) = -56.8° in (a), $P(g_i|x_1) = 0.448, 0.409$ and $0.144$ in the normal, minimal CAD and CAD groups, respectively, as plotted in the bar chart in the lower right corners. CAD, coronary artery disease.
Finally, the presence of significant CAD is determined by the weighted sum of the posterior probability of belonging to CAD in comparison to the probability of belonging to the minimal CAD or normal groups. A subject is classified as having significant CAD if \( P(CAD | x_1, x_2, x_3, x_4, x_5) \) is larger than \( P(\text{minimal CAD} | x_1, x_2, x_3, x_4, x_5) \) and \( P(\text{normal} | x_1, x_2, x_3, x_4, x_5) \). Otherwise, he/she is classified as not having significant CAD. The weighting factors were adjusted experimentally so that the diagnostic accuracy is maximized. The best classification result was obtained with the weighting factors of 0.3, 1.0, 2.0, 1.7 and 2.5 for T_FMA, T_FMA–R_FMA, TT_CAMx, TT_CAMx–R_FMA and TT_CMD, respectively.

\[
P(g | x_1, x_2, x_3, x_4, x_5) = \sum_j w_j P(g | x_j)
\]  

(2)

Figure 4. Diagnostic accuracy of magnetocardiography (MCG), electrocardiography (ECG) and echocardiography (Echo) in determining ischemia. NPV, negative predictive value; PPV, positive predictive value.

### Table 3. Diagnostic Accuracy of MCG in a Conditioned Population

<table>
<thead>
<tr>
<th>Including criteria</th>
<th>No. patients</th>
<th>Tool</th>
<th>Sensitivity (%)</th>
<th>Specificity (%)</th>
<th>PPV (%)</th>
<th>NPV (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>All patients</td>
<td>237</td>
<td>MCG</td>
<td>84.0</td>
<td>85.0</td>
<td>91.3</td>
<td>74.0</td>
</tr>
<tr>
<td>Negative biomarker</td>
<td>146</td>
<td>MCG</td>
<td>78.1</td>
<td>82.6</td>
<td>87.7</td>
<td>70.4</td>
</tr>
<tr>
<td>Angiogram available</td>
<td>102</td>
<td>MCG</td>
<td>73.5</td>
<td>82.3</td>
<td>84.3</td>
<td>70.7</td>
</tr>
</tbody>
</table>

PPV, positive predictive value; NPV, negative predictive value. Other abbreviations see in Tables 1, 2.

P was calculated for the comparison of diagnostic accuracy between MCG and ECG.

Results

All patients were classified according to the wMAP method using the prior probability distribution (Figure 3) and weighting factors. As a result, 199 out of 237 patients in the CAD group were classified as having CAD on MCG, and 108 out of 127 patients in minimal CAD group had negative MCG signs, giving a sensitivity, specificity, PPV and NPV of 84.0%, 85.0%, 91.3% and 74.0%, respectively (P<0.001). In the normal group, 87 out of 89 subjects had negative MCG signs (97.8%). The diagnostic accuracy of MCG was compared with the mean of the cosines and sines of each angle. In order to estimate a probability distribution function, the histogram was represented by a combination of normal distribution function and gamma distribution function. Sensitivity, specificity, positive predictive value (PPV) and negative predictive value (NPV) were determined in comparison between the CAD group and minimal CAD group according to the contingency table. Differences in categorical data were assessed on chi-square test. Statistical significance was defined as \( P<0.05 \).
that of the 12-lead ECG and echocardiography in Figure 4, showing a significantly higher sensitivity and NPV for MCG than ECG and echocardiography (P<0.001). In contrast, the specificity of MCG was lower than for ECG (P=0.02) or echocardiography (P<0.001), and the PPV of MCG was lower than that of echocardiography (P<0.03), but the differences were not so large. Here an ECG was defined as positive in the case of pathologic Q wave, amplitude of ST depression ≥0.05 mV or T inversion ≥0.2 mV.25 The diagnostic accuracy of MCG was also tested in a conditioned population. In the subgroup of 238 patients who had no indications of ischemia on biochemistry but who underwent coronary angiography due to suspected CAD, MCG still had a higher sensitivity of 78.1% and NPV of 70.4% (P<0.001), compared to that of 30.1% and 43.6%, respectively, on ECG. But the specificity and PPV of MCG were similar to those of ECG (P>0.05). When 57 patients with indications of ischemia on ECG were further excluded from the subgroup, MCG had a sensitivity of 73.5% and NPV of 70.7% in the remaining 181 patients (P<0.001). The results are summarized in Table 3.

Discussion

The present retrospective study analyzed MCG datasets of 364 patients with suspected ACS without ST elevation and was focused on the diagnostic accuracy in the heterogeneous population for practical use in a general clinical environment. Diagnostic accuracy was then determined in comparison between patients with and without severe CAD. Furthermore, severe CAD was defined as stenosis ≥50% at not only major coronary arteries but also their branches for the detection of CAD in an early stage of stenosis. Therefore, 23 patients, who would have been diagnosed as having minimal CAD on general criteria of a stenosis ≥70% in a major coronary artery, were additionally included in the CAD group. Among them, 18 patients were classified as having CAD on MCG (78.3%).

It is well known that when there is ischemia, biological injury currents are generated that can be detected on MCG. Therefore an abnormality in cardiac depolarization or repolarization is reflected in an abnormality in the MFM.5 But a diagnostic method for CAD detection has not been established. Here, we used an abnormality of several MCG parameters, such as the orientation angle derived from MFM and the current vector, and a dynamic change in the strength of the main current vector during the time interval between the beginning of the T wave and T wave apex. In general, abnormality of MCG parameters was determined dichotomously so that the probability of having CAD is 1 or 0 depending on whether the parameter exceeds the normal range or not. Such binary decision might be useful in discriminating between a high-risk group (UA or MI) and a healthy group.10 As shown in Figure 2, however, it is limited in the heterogeneous population because a number of CAD patients have MCG parameters in the intermediate region between normal and abnormal ranges. Using our method, the presence of CAD is determined by comparing the probability, which was calculated from the prior knowledge on the probability distribution of the MCG parameters depending on the presence of CAD. In this way we obtained better diagnostic accuracy than the previous studies,11–14 showing that the proposed method is more effective than the conventional binary decision method. This WMAP method was tested with a small population (86 CAD patients, 59 minimal CAD patients, 66 healthy volunteers) and was applied to all patients to confirm the validity in a relatively large population. The results demonstrate that the diagnostic criteria proposed in the present study may not lose the ability to correctly classify newly added patients.

Compared with ECG and echocardiography, MCG had distinctly higher sensitivity and NPV (Figure 4). The results show that MCG may be useful to avoid the inadvertent discharge of the patient who truly has myocardial ischemia. The difference of diagnostic accuracy between MCG and ECG was further tested in a conditioned population (Table 3). First, 91 patients, who clearly had high probability of CAD due to positive biomarkers, and 35 patients, who were discharged without undergoing coronary angiography due to high probability of minimal CAD, were excluded. In this subgroup of 238 patients, MCG still had higher sensitivity and NPV with similar specificity and PPV, compared to ECG. When 57 patients with indications of ischemia in ECG were further excluded from the subgroup, MCG had a sensitivity of 73.5% and NPV of 70.7% in the remaining 181 patients. These results demonstrate the efficacy of MCG as a complementary tool to ECG in patients with no indication of ischemia according to various diagnostic tools before angiography.

Study Limitations

First, MCG measures abnormal electrophysiological changes caused by ischemia, and the severity of ischemia was known to be affected by the degree and morphology of stenosis.26 In the present study the severity of CAD was defined by the degree of stenosis on coronary angiography and we need a clear gold standard for ischemia. Second, our classification method is based on the probability distribution function, examined in a retrospective study and the findings have not been prospectively validated. For clinical application of the method, the diagnostic accuracy should be accepted reliably without loss of the classification accuracy when new cases are added in the future. Finally, the presence of CAD is represented as a probability without electrophysiological or anatomical information. Further work to localize the ischemic region in a realistic heart model is needed.

In conclusion, MCG was acceptably sensitive and specific in identifying patients with ACS even in the absence of specific findings on ECG and positive biomarker tests. Thus, MCG seems beneficial for the early triage of patients with acute chest pain.

References


