The Use of Body Surface Potential Map for Identifying Sites of Accessory Pathway in Patients with Wolff-Parkinson-White Syndrome

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SUMMARY

The body surface potential map (BSPM) may reflect regional myocardial electrical activity. This technique can thus provide information regarding the excitation of ventricles. This study is an attempt to evaluate the usefulness of BSPM in determining the sites of the atrioventricular (AV) accessory pathway (AP) in patients with Wolff-Parkinson-White (W-P-W) syndrome.

The BSPMs were obtained from 40 consecutive patients with W-P-W syndrome in a fasting state, using the heart potential map system designed by Toyama et al. Unipolar electrocardiograms were recorded simultaneously from 87 lead points on the chest surface, including 59 lead points on the anterior chest and 28 on the back. Wilson's central terminal was used as a voltage reference and BSPMs in an isopotential distribution pattern were made every millisecond throughout ventricular activation from these unipolar ECGs with the use of a microcomputer system.

All patients underwent an electrophysiologic study (EPS) at cardiac catheterization. We analyzed the potential distribution during ventricular depolarization and compared the results between EPS and BSPM findings.

The following results were obtained: (1) seven types of BSPM pattern were identified in accordance with the sites of the AV AP confirmed by EPS; (2) the location of the potential minimum of ventricular depolarization and the direction of the excitation wavefront during early ventricular depolarization, the reversal pattern of ventricular potential distribution, the epicardial right ventricular breakthrough and the dynamic change of ventricular potential distribution were useful for the detection of the ventricular pre-excitation site; (3) epicardial right ventricular breakthrough occurred in nearly all patients with left ventricular free wall accessory AV connections; (4) the abnormal early reversal pattern of ventricular potential distribution did not occur in patients with left ventricular AV connections but did appear in most patients with right
ventricular free wall AV connections. Accordingly, BSPM is a reliable non-invasive procedure to determine the ventricular pre-excitation sites of patients with W-P-W syndrome. (Jpn Heart J 1998; 39: 445-455)

**Key words:** Potential minimum, Potential maximum, Right ventricular breakthrough, W-P-W syndrome

WOLFF-PARKINSON-WHITE (W-P-W) syndrome is considered to be caused by the early excitation of some ventricular sites through accessory atrio-ventricular (AV) pathways.\(^\text{1}\) The accessory pathway (AP) can exist anywhere along the AV ring except where the left atrium is attached to the aortic ring.\(^\text{2}\) Identification and localization of the AP in patients with this syndrome may be imperative for management. This is particularly true for patients with significant tachyarrhythmia requiring interruption of the APs. A reliable non-invasive procedure to determine the location of APs in patients with this syndrome would be clinically useful. The body surface potential map (BSPM) may reflect regional myocardial electrical activity and can provide information regarding the ventricular excitation which can not be derived from conventional electrocardiography.\(^\text{3,4}\) This technique has been reported to be potentially applicable in identifying the location of APs in patients with W-P-W syndrome.\(^\text{5-17}\) However, the identifying parameter remains to be established.

In this report, we will present the BSPM data which are useful in identifying the site of the AP in patients with W-P-W syndrome.

**MATERIALS AND METHODS**

Forty consecutive patients with W-P-W syndrome were studied. Their ages ranged from 16 years to 70 years. There were 28 males and 12 females. Electrophysiological (EPS) studies at cardiac catheterization were performed after BSPM recording to identify the earliest ventricular activation. The BSPM was obtained in a fasting state using the heart potential map system designed by Toyama et al.\(^\text{18}\) (HPM-6500 Fukuda Denshi Co., Ltd., Nagoya, Japan). Unipolar electrocardiograms were recorded simultaneously from 87 lead points on the chest surface. There were 59 lead points on the anterior chest wall and 28 on the back. The positions of electrodes in this system are shown in Figure 1. The points A, I and E were at the right mid-axillary line, left mid-axillary line and mid-sternal line, respectively. The distance from A to E was divided into 4 equal parts and the positions of lead points B, C and D were thus ascertained. Similarly, the distance from E to I was also divided into 4 equal parts and the points F, G and H were positioned. Points J and M were determined at the same interval using the method for determining points described above. The distance from J to M was
Figure 1. Position of electrodes in the recording system of the body surface potential map.

divided into 3 equal parts and the positions of lead points K and L were ascertained. Accordingly, the anterior chest wall was divided into 8 columns (from A to I) and the back into 5 columns (from I to A). In addition, the thorax was divided horizontally into 7 rows. The point E6 was set at the level of the 2nd intercostal space, and E4 at the 5th intercostal space. The position of E5 was centered between E4 and E6. The points E1, E2, E3 and E7 were set at the same intervals as those between E6 and E5 or E4 and E5.

Wilson’s central terminal was used as a voltage reference and BSPMs were made in an isopotential distribution pattern every millisecond throughout ventricular activation from these unipolar ECGs using a microcomputer system. To illustrate the BSPMs, a planar map was drawn with a cut along the right midaxillary line. The solid line illustrated an equipotential line at an interval of 0.4 mV. The line with zero potential, representing the potential of Wilson’s central terminal, was called the zero line. The symbol “+” indicates a potential maximum and the symbol “−” indicates a potential minimum. We analyzed the potential distribution during ventricular depolarization. The parameters for analysis included potential distribution in the early period of ventricular depolarization, the epicardial right ventricular breakthrough (RVBT) and dynamic change of potential distribution.

We studied the relationship between BSPM findings and sites of AP as confirmed by EPS at catheterization.
RESULTS

Location of accessory pathway: A single AP was present in all 40 patients. These 40 patients were classified into 7 groups according to the pre-excited sites as determined by EPS at cardiac catheterization (Figure 2). There were 9 patients with anterior right ventricular (RV) AP in group I; 12 with lateral RV AP in group II; 2 with posterior RV AP in group III; 4 with posteroseptal AP in group IV; 5 with posterior left ventricular (LV) AP in group V; 5 with lateral LV AP in group VI; and 3 with anterior LV AP in group VII.

The potential distribution on the thorax of patients with Wolff-Parkinson-White syndrome during the early period of ventricular depolarization: There were 7 types of potential distribution patterns during the early period of ventricular depolarization (the initial 40 msec period with constant potential distribution). This classification was made in accordance with the anatomic site of APs along the A-V ring as shown in Figure 2.

In patients with the AP in the anterior RV, the potential minimum was located higher than the potential maximum during the initial period of the QRS complex. The potential minimum appeared in the middle or right superior portion of the forechest and the potential maximum was in the left-inferior portion of the thorax, either anterior or posterior (type I BSPM pattern, Figure 3A). An example is illustrated in Figure 4A.

In patients with the AP in the lateral RV, the potential minimum was located lower than the potential maximum during the initial period of the QRS complex. The potential minimum appeared in the left-inferior portion of the thorax, either anterior or posterior (type I BSPM pattern, Figure 3A). An example is illustrated in Figure 4A.
Figure 3. The potential distribution of Wolff-Parkinson-White syndrome during the initial period of ventricular depolarization.

located lower than the potential maximum during the initial period of the QRS complex. The potential minimum was found in the middle or right-inferior portion of the forechest, while the potential maximum was located in the left-superior portion of the forechest (type II BSPM pattern, Figure 3B). An example is shown in Figure 4B.

In patients with the AP in the posterior RV, the potential minimum was located lower than the potential maximum during the initial period of the QRS complex. The potential minimum appeared in the inferior portion of the right
Figure 4. Examples of the potential distribution of Wolff-Parkinson-White syndrome during the initial period of ventricular depolarization.
Various parameters of body surface potential maps (Table): There were differences in various parameters of the absolute value of the BSPMs between

<table>
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<tr>
<th>BSPM classification</th>
<th>Right ventricular breakthrough</th>
<th>Abnormal early reversal pattern of ventricular depolarization</th>
</tr>
</thead>
<tbody>
<tr>
<td>I (9)</td>
<td>No: 0 %: 0</td>
<td>No: 8 %: 88.9</td>
</tr>
<tr>
<td>II (12)</td>
<td>No: 0 %: 0</td>
<td>No: 10 %: 83.3</td>
</tr>
<tr>
<td>III (2)</td>
<td>No: 0 %: 0</td>
<td>No: 2 %: 100.0</td>
</tr>
<tr>
<td>IV (4)</td>
<td>No: 4 %: 100.0</td>
<td>No: 0 %: 0</td>
</tr>
<tr>
<td>V (5)</td>
<td>No: 4 %: 80.0</td>
<td>No: 0 %: 0</td>
</tr>
<tr>
<td>VI (5)</td>
<td>No: 5 %: 100.0</td>
<td>No: 0 %: 0</td>
</tr>
<tr>
<td>VII (3)</td>
<td>No: 3 %: 100.0</td>
<td>No: 0 %: 0</td>
</tr>
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BSPM = body surface potential map. Figures in parentheses indicate the numbers of patients per group.

In patients with the AP in the posteroseptum, the potential minimum was slightly lower than the potential maximum in all cases except one, during the initial period of the QRS complex. The potential minimum was found in the right-inferior portion of the back and the potential maximum was in the middle portion of the forechest in all cases except one (type IV BSPM pattern, Figure 3D). An illustration is given in Figure 4D.

In the group with the AP in the posterior LV, the potential minimum was located higher than the potential maximum during the initial period of the QRS complex. The potential minimum appeared in the middle-superior portion or left-superior portion of the back while the potential maximum was in the middle portion of the forechest (type V BSPM pattern, Figure 3E). An example is shown in Figure 4E.

In the group with the AP in the lateral LV, the potential minimum was located lower than the potential maximum during the initial period of the QRS complex. The potential minimum appeared in the left-inferior portion of the back while the potential maximum was found in the middle portion of the forechest (type VI BSPM pattern, Figure 3F). An example is shown in Figure 4F.

In patients with the AP in the anterior LV, the potential minimum was located lower than the potential maximum during the initial period of the QRS complex. The potential minimum was found in the left-inferior portion of the forechest and the potential maximum appeared in the right-superior portion of the forechest (type VII BSPM pattern, Figure 3G). An example is shown in Figure 4G.
right-sided and left-sided APs.

None of the patients with RV APs demonstrated RVBT. However, the RVBT was found in 16 of 17 patients with LV APs.

An abnormal early reversal pattern of ventricular depolarization in which the absolute value of the potential minimum was greater than that of the potential maximum during the early phase of ventricular depolarization (the initial 40 msec period with constant potential distribution), was observed in 20 of 23 patients with the RV AP, but not in patients with LV AP.

**DISCUSSION**

This study identified 7 types of BSPM according to the sites of APs determined by EPS. Yamada et al. first identified 3 types of BSPM in patients with W-P-W syndrome, according to the potential distribution during the QRS complex. There then followed several reports addressing the question of the usefulness of BSPM in identifying the insertion site of the accessory A-V pathway. De Ambroggi et al. classified 6 types of BSPM based on the location of the potential minimum and potential maximum during the delta wave. Iwa et al. observed 5 types of BSPM according to the location of the potential minimum at 40 milliseconds of the QRS complex and stressed the value of this criterion in predicting the sites of APs. Benson et al. observed 8 types of BSPM according to the different sites of APs while Kamakura et al. found 7 types. Liebman et al. observed 6 types of BSPM and emphasized the value of the QRS map and RVBT in predicting the location of APs.

Several indices of BSPM were considered as useful criteria in predicting the site of APs. Iwa et al. did not accept the analysis of the potential distribution during 40 milliseconds after the onset of the QRS complex as a reliable criterion for determining the location of APs in patients with W-P-W syndrome. Benson et al. reported the predictive reliability of potential distribution in the first 40 milliseconds of the QRS complex for the sites of APs. Kamakura et al. found that stability of the QRS potential distribution during the delta wave always occurred both in the initial 40 milliseconds and at a peak negative potential of $-150$ to $-200 \mu V$. They therefore recommended that an amplitude-based map rather than a time-based map was useful in localization of APs. The predictive ability of their criterion ranged from 30 to 35 of 41 cases. Liebman et al. stressed the importance of both negative and positive, not just negative potentials, and disagreed with the recommendation of Kamakura et al. Benson et al. reported the superiority of the ST-T map over the QRS map in predicting the location of APs. Some reports also demonstrated the close relationship between BSPM of repolarization abnormalities and the location of the accessory
pathway in patients with manifest W-P-W syndrome. However, Liebman et al.\textsuperscript{12})
documented the usefulness of this potential distribution, pointing out that it pro-
vided limited accuracy only in some cases with marked ventricular pre-excitation.
It is well known that the potential maximum in a given area on the chest surface
represents the area that an intracardiac excitation wavefront is pointing toward.
The potential minimum indicates the negative aspect of a wavefront where the
excitation is initiated.\textsuperscript{19}) The location of the potential minimum during the delta
wave is thus an ideal index for predicting the pre-excitation site of APs and has
proven to be useful in the previous literature, and in our report. Liebman et al.\textsuperscript{12})
even found that the ventricular insertion sites of APs determined by QRS analysis
of BSPM and surgical EPS mapping were identical or within 1.5 cm in all of their
cases with a single AP. The distribution of heart potentials on the body surface
depends not only on the sites of the excitation waves, but also on the geometry of
the chest and the conductivity of body tissue. It is not surprising that overlap sites
of potential minimum during the delta wave of the QRS complex were found in
different locations of the AP. This means that the potential minimum alone
cannot be a useful criterion for identifying the sites of the AP. If we take into
consideration the direction of the excitation impulse as well as the location of the
potential minimum, identification of AP sites will be improved. In our study, this
combination of indices of BSPM clearly classified the 7 types of BSPM useful in
identifying the sites of APs.

Epicardial RVBT is an indication of conduction through the AV node and
His purkinje fiber.\textsuperscript{20}) Liebman et al.\textsuperscript{12}) did not find RVBT in their patients with
RV free wall or anteroseptal APs. In patients with the AP in the RV, the RV is
not activated in the usual way by the specialized AV conduction system and the
activation sequence of the RV is abnormal. Consequently, there is no RVBT. In
patients with LV APs, however the activation of the RV develops via the special-
ized AV conduction system and results in a typical RVBT pattern. This was also
true of the patients in the present study.

In eccentrically located dipole with radial direction, the surface potential
maximum and potential minimum are located at the ends of a diameter with a
higher absolute value at one extreme.\textsuperscript{6}) The greater strength of the anterior
minimum as compared to the posterior maximum suggests that the pre-excitation
wavefront was closer to the anterior chest surface than the posterior one. The
location of pre-excitation could be either in the RV wall or in the right side of the
interventricular septum.\textsuperscript{6}) The abnormal early reversal pattern of ventricular de-
polarization was found in nearly all patients with RV APs in our study, demon-
strating that this index is useful in differentiating between right-sided and left-
sided APs.

Accordingly, the BSPM is a reliable, noninvasive procedure to determine
the ventricular pre-excitation sites in patients with W-P-W syndrome. The following BSPM findings have been shown to be useful in identifying sites of APs: (1) the location of the potential minimum and direction of excitation wavefront during early ventricular depolarization; (2) epicardial RVBT; and (3) abnormal early reversal pattern of ventricular depolarization.

REFERENCES

