Validation of the Ventricular Gradient Comparing Sinus Beats and Ectopic Beats

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Abstract

Introduction. Wilson postulated that the ventricular gradient (VG) is independent of the cardiac activation order. This concept has been criticized several times, but these studies had small numbers of patients, selective pathology, were mostly done in two dimensions, and did not take restitution effects into account. The purpose of our study was to validate this concept by intra-individual comparison of the VG of sinus beats and ectopic beats, and thus to estimate the effects of altered ventricular conduction and of restitution.

Methods. We studied standard diagnostic ECGs of 118 patients with accidental extrasystoles, who had either normally conducted supraventricular ectopic beats (SN, N=6), aberrantly conducted supraventricular ectopic beats (SA, N=20), or ventricular ectopic beats (V, N=92). We computed the ventricular gradient vectors of the predominant beat, VGp, of the ectopic beat, VGe, the VG difference vector, VGpe, and compared their sizes.

Results. The sizes of the VGe vectors were significantly larger than those of the VGp vectors in the SA and V ectopic beats. The sizes of the VGpe vectors were three times larger than the difference in size of the VGe and VGp vectors, a manifestation of differences in the spatial directions of the VGp and VGe vectors. Prematurity of the ectopic beats had no influence on these results.

Discussion. This outcome can mechanistically be explained by the electrotropic interactions during ventricular repolarization that change with a changing ventricular activation order. Because of this electrophysiological mechanism, the concept of a conduction-independent ventricular gradient as put forward by Wilson is untenable. Therefore, the VG cannot straightforwardly be used in ECG diagnostics.

1. Introduction

The ventricular gradient (VG), defined as the integral of the ECG amplitude over the QT interval, has intrigued many investigators ever since it was introduced by Wilson et al. in 1931 (1). Originally defined for separate ECG leads, the VG evolved into a three-dimensional concept when vectorcardiography emerged: the spatial ventricular gradient (2). Already from the beginning, the VG was thought to be independent of the intraventricular conduction order, only depending on “local variations in the excitatory process” (1, 3), i.e., on action potential heterogeneity. Hence, VG was considered as an individual intrinsic property of the ventricular myocardium. In 1957, Burger mathematically derived a formula for the VG, showing that it was proportional to the volume integral of the action potential duration gradients over the heart (4, 5):

$$\vec{G} = \int_{Q_T} \vec{H} \cdot dt = -\int_{vol} \vec{\nabla} (h \cdot \tau) \cdot dv$$

$\vec{G}$ = spatial ventricular gradient (VG)
$\vec{H}$ = heart vector
$h$ = resting potential (maximal diastolic potential)
$\tau$ = action potential duration

As this equation does not figure the excitation order of the heart, VG could be useful in ECG diagnostics. When, in a patient, the ECG changes while the VG remains the same, these changes should have been caused by differences in the excitation order but not in the intrinsic electrophysiological properties of the cardiac myocytes. When both the ECG and the VG change, the electrophysiological properties of the myocardium must have changed too. This property of the VG could, e.g., be helpful to detect acute ischemia in patients with a new left bundle branch block (BBB). If the new BBB is caused by the acute ischemia, not only the ECG but also the VG of this patient should have changed due to ischemia-based action potential changes. If the new BBB is caused by a local defect in the conduction system that is unrelated to acute ischemia, the VG is expected to remain unchanged.

Since decades the concept of conduction-independency of the VG has met criticisms. Simonson et al. (6) did not agree with Wilson et al. (3) that the VG was “reasonably constant in spite of the greatly different pathways of excitation and grossly different patterns and electrical axes”. They calculated that the amount of variability in Wilson’s experiments was 25 percent of the mean, and therefore should not have been interpreted as “a small variability” (3). Angle (7) determined the VGs in the ECGs with intermittent right BBB as published by Wilson et al. (8) and by White (9), and in the ECGs with intermittent left BBB as published by Segers and Boyadjian (10). The VGs in the complexes exhibiting BBB were about half the size of the VGs in the normally conducted complexes in right
BBB, and about two thirds the size of the VGs in the normally conducted complexes in left BBB.

Calculations in the above studies were done in the frontal plane. Berkun et al. (11) investigated conduction independency of the VG in 3D. In 1966, Cosma et al. (12) stated that a 3D approach of the VG requires time-integral calculations in an orthonormal vectorcardiographic lead system. They compared the VG of normal sinus beats with the VG of premature ventricular contractions in patients with various pathology and concluded that the VG did not remain constant when the ventricular activation changes.

Other investigators were more positive about the presumed independence of the ventricular activation order. Lux et al. (13, 14) investigated this using body surface potential mapping in dogs. They concluded that “QRST iso-area maps were shown to be independent of ventricular activation sequence with the exception of features which are explicable on the basis of electrotonic interaction during ventricular repolarization” (15): “it appears that the influence of activation order on recovery properties should be considered in evaluating the significance of a given value of the ventricular gradient” (16).

Another mechanism that may affect VG is restitution, the relationship between the duration of a cardiac action potential and the length of the preceding diastolic interval. When studying the VG in the setting of ectopic beats, it should be taken into account that not only the ectopic ventricular activation order is different from that of the normal beat, but also the preceding diastolic interval (in supraventricular as well as in ventricular ectopic beats). Premature beats have shorter action potential durations, but the amount of shortening is known to be not homogeneously distributed over the heart: epicardial action potentials shorten more with prematurity than endocardial action potentials do (17). In this way, such inhomogeneous restitution alters the VG of ectopic beats.

For clinical use, a robust measure of the recovery properties of the heart is crucially important. Independence of ventricular activation order would render such a measure useful for intrapatient comparison of ECGs with and without conduction disorders. In our current study, we addressed the dependence of the VG on ventricular activation order and on restitution by retrospectively analyzing standard 10-second 12-lead ECG recordings with a spontaneous supraventricular or ventricular ectopic beat, to further quantify and qualify the differences between the VGs of the predominant and the ectopic beats.

2. Methods

We searched the historical ECG database of the Leiden University Medical Center (LUMC) over a period of ten successive months for technically sound digital standard 10-second 12-lead ECGs of patients who were 18 years or older, who showed sinus rhythm and contained at least one spontaneous supraventricular or ventricular ectopic beat.

2.1. ECG analysis

The ECGs were analyzed and generally characterized by the Glasgow Royal Infirmary (GRI) software (18), thus labelling each ECG as normal or abnormal. We additionally classified the GRI detected abnormalities as long QT, axis deviations, BBBs, intraventricular conduction disorders, hypertrophy, old infarctions, pericarditis, ST abnormalities, or low QRS voltage.

The ECGs were additionally analyzed by the Leiden ECG Analysis and Decomposition Software (LEADS) (19). Briefly, LEADS combines automated analysis and human interaction to identify the heartbeats of interest in an ECG recording, and to verify and when necessary to adjust the essential landmarks in time (onset of the QRST complex and end of the T wave, together defining the interval for the computation of the VG). Initially, LEADS computes a vectorcardiogram (VCG) by multiplying the ECG by the Kors transformation matrix (20). Then, heartbeats are detected in the VCG-derived spatial velocity signal (19). Next, ectopic beats are excluded, thus focusing on the analysis of the predominant QRST complex. Baseline correction is done before averaging the predominant beats, by cubic spline interpolation through the PQ segments immediately preceding the predominant QRS complexes.

For the purpose of this study, ectopic beats had to be analyzed as well. To facilitate the analysis of a separate ectopic beat, a special experimental version of the LEADS program was developed, in which the ectopic beats were baseline-corrected with the baseline as calculated from the surrounding predominant beats in the ECG recording.

From the variables produced by LEADS, we calculated the prematurity of the ectopic beat (the ratio of the coupling interval of the ectopic beat and the mean interval between consecutive predominant beats). From the VG vectors of the predominant and the ectopic beats, $\overrightarrow{VG}_p$ and $\overrightarrow{VG}_e$, we computed two measures of difference:

1. the magnitude of the difference vector between the VGs of the ectopic and predominant beats: $|\Delta \overrightarrow{VG}_{ep}|$
2. the magnitude of the difference between the VG magnitudes of the ectopic and predominant beats: $|\overrightarrow{VG}_e| - |\overrightarrow{VG}_p|$

Finally, we divided the ectopic beats into 3 categories: supposed supraventricular origin and normal conduction (SN), supposed supraventricular origin and aberrant conduction (SA), and supposed ventricular origin (V).

2.2. Statistics

To compare $\overrightarrow{VG}_p$ and $\overrightarrow{VG}_e$, we calculated the means and standard deviations of the $\overrightarrow{VG}_p$ and $\overrightarrow{VG}_e$ magnitudes, and tested the difference of the means by a two-sided paired T test. Additionally, we computed the mean and standard
deviation of the VG difference vector magnitude, $|\Delta VG_{ep}|$. Values were computed for all patients together, and for the subgroups of patients with ectopic beats of the SN, SA and V categories. Finally, we statistically compared the values of these subgroups by two-sided unpaired T tests, except for the comparison of $|\Delta VG_{ep}|$, for which a one-sided unpaired T test was used. Additional linear regression analyses were performed in the context of several scatterplots, as presented in the Results section.

3. Results

Table 1 shows the anthropomorphic characteristics of the study group; cardiovascular diagnoses are described in Table 2. An average of 2.2 cardiovascular conditions was present per patient. Table 3 shows the ECG qualification by the GRI software. The majority of the ECGs (72.0%) was interpreted as abnormal or borderline abnormal.

<table>
<thead>
<tr>
<th>Diagnosis</th>
<th>n</th>
<th>%</th>
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<tbody>
<tr>
<td>Coronary atherosclerotic disease</td>
<td>49</td>
<td>41.5</td>
</tr>
<tr>
<td>Healed myocardial infarction</td>
<td>28</td>
<td>23.7</td>
</tr>
<tr>
<td>Chronic heart failure</td>
<td>15</td>
<td>12.7</td>
</tr>
<tr>
<td>Cardiomyopathy</td>
<td>13</td>
<td>11.0</td>
</tr>
<tr>
<td>Systemic/pulmonary hypertension</td>
<td>60</td>
<td>50.8</td>
</tr>
<tr>
<td>Hypertrophy (ventricular/atrial)</td>
<td>2</td>
<td>1.7</td>
</tr>
<tr>
<td>Valvular disease</td>
<td>34</td>
<td>28.8</td>
</tr>
<tr>
<td>Congenital heart disease</td>
<td>6</td>
<td>5.1</td>
</tr>
<tr>
<td>Rhythm disorders</td>
<td>45</td>
<td>38.1</td>
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<tr>
<td>Conduction disorders</td>
<td>14</td>
<td>11.9</td>
</tr>
<tr>
<td></td>
<td>266</td>
<td>225</td>
</tr>
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</table>

Table 2. Cardiovascular diagnoses. On the average, 2.2 cardiovascular conditions were present per patient.

<table>
<thead>
<tr>
<th>Interpretation</th>
<th>n</th>
<th>%</th>
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<tbody>
<tr>
<td>Normal</td>
<td>2</td>
<td>1.7</td>
</tr>
<tr>
<td>Borderline normal</td>
<td>31</td>
<td>26.3</td>
</tr>
<tr>
<td>Borderline abnormal</td>
<td>17</td>
<td>14.4</td>
</tr>
<tr>
<td>Abnormal</td>
<td>68</td>
<td>57.6</td>
</tr>
<tr>
<td></td>
<td>118</td>
<td>100.0</td>
</tr>
</tbody>
</table>

Table 3. Automated GRI classification of the ECGs.

Figure 1. Magnitudes of the ventricular gradient vectors of the predominant beat ($|VG_p|$), of the ectopic beat ($|VG_e|$), and of the difference vector ($|\Delta VG_{ep}|$). Statistical comparisons were done with unpaired (columns) and paired (rows) two-sided T tests. SN = patients with a normally conducted supraventricular ectopic beat; SA = patients with an aberrantly conducted supraventricular ectopic beat; V = patients with a ventricular ectopic beat.

Table 4 shows the distribution of the ECGs in our study group (All; N=118). Data are displayed as mean±SD [range]. BMI = body mass index.

|          | $|VG_p|$ (mV∙ms) | $|VG_e|$ (mV∙ms) | $|\Delta VG_{ep}|$ (mV∙ms) |
|----------|----------------|----------------|---------------------------|
|          | Mean SD        | Mean SD        | Mean SD                   |
| All (N=118) | 47.78 ±24.62 | 53.72 ±25.03 | 19.94 ±9.76 |
| SN (N=6) | 57.01 ±27.46 | 59.5 ±32.77 | 0.66 ±17.83 |
| SA (N=20) | 48.45 ±31.69 | 54.3 ±27.89 | 0.04 ±18.87 |
| V (N=92) | 47.03 ±22.85 | 53.22 ±24.12 | <0.001 ±20.31 |

Table 4. Means and standard deviations of the magnitudes of the ventricular gradient vectors of the predominant beat ($|VG_p|$), of the ectopic beat ($|VG_e|$), and of the difference vector ($|\Delta VG_{ep}|$). Statistical comparisons were done with unpaired (columns) and paired (rows) two-sided T tests. SN = patients with a normally conducted supraventricular ectopic beat; SA = patients with an aberrantly conducted supraventricular ectopic beat; V = patients with a ventricular ectopic beat.
4. Discussion

We investigated the dependence of the VG on activation order and on restitution by analyzing standard diagnostic ECGs with a spontaneous ectopic beat, measuring intraindividual differences between the VGs of predominant and ectopic beats in 118 patients (Table 1) with various cardiovascular pathology (Table 2) and with, for the greater part, abnormal ECGs (Table 3). Prematurity varied widely (0.43–1.03; Figure 9), thus covering almost the entire electrophysiologically possible range.

Ectopic VG vector magnitudes were significantly larger than predominant VG magnitudes, for SA and V, but not for SN ectopic beats (Table 4). VG difference vectors were larger than the differences between the predominant and ectopic VG magnitudes, signalling predominant-ectopic differences in spatial VG orientation (Table 4). The magnitudes of the VG difference vectors were uncorrelated with the prematurity of the ectopic beats (Figure 1).

From the times in which the concept of the VG was formulated, it was considered important because of its presumed independency of ventricular activation order. In this way, the VG could help to diagnose ECG changes in a given subject as either changes in the ventricular conduction system, or changes in the electrophysiological properties of the ventricular myocardium. Later, additional use of the VG in risk assessment was proposed (for an overview, see the publication by Waks and colleagues [21]). Our study addresses the premise that the VG forms an individual ‘electrocardiographic constant’ for the electrophysiological properties of the ventricular myocardium. Our results demonstrate that the VG may change considerably with a deviating ventricular activation pattern, although prematurity is not playing a major role in this effect. Our results can mechanistically be explained by the electrotonic interactions during ventricular repolarization that change with the ventricular activation order (15); and this precludes diagnostic use of the VG.

References


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