

Chapter 1

The Electrical Activity of the Heart

Basic concepts

The heart is a pump that sends blood to every organ in the human body. This is carried out through an electrical stimulus that originates in the sinus node and is transmitted through the specific conduction system (SCS) to contractile cells.

During the rest period, myocardial cells present an equilibrium between the positive electrical charges outside and the negative charges inside. When they receive the stimulus propagated from the sinus node, the activation process of these cells starts. **The activation** of myocardial cells is an electro-ionic mechanism (as explained in detail in Chapter 5) that involves two successive processes: **depolarization**, or loss of external positive charges that are substituted by negative ones, and **repolarization**, which represents the recovery of external positive charges.

The process of depolarization in a contractile myocardial cell starts with the formation of a **depolarization dipole** comprising a pair of charges ($-+$) that advance through the surface cell like a wave in the sea, leaving behind a wave of negativity (Figure 1.1A). When the entire cell is depolarized (externally negative), a new dipole starting in the same place is formed. This is known as a **dipole of repolarization** ($+ -$). **The process of repolarization**, whereby the entire cell surface is supplied with positive charges, is then initiated (Figure 1.1B).

The expression of the dipoles is a vector that has its head in the positive charge and the tail in the negative one. An electrode facing the head of the vector records positivity (+), whereas when it faces the tail, it records negativity ($-$) (Figures 1.1–1.3; see also Figures 5.24, 5.25, and 5.28). The deflection originating with the depolarization process is quicker because the process of depolarization is an active one (abrupt entry of Na ions, and later Ca) and the process of repolarization is much slower (exit of K) (see Chapter 5, Section “Transmembrane action potential”).

If what happens in one contractile cell is extrapolated to the left ventricle as the expression of all myocardium, we will see that the **repolarization process in this case starts in the opposite place to that of depolarization**. This explains why the repolarization of a single contractile cell is represented by a negative wave, whereas the repolarization of the left ventricle expressing the human electrocardiogram (ECG) is represented by a positive wave because the repolarization process of all left ventricles starts in the zone less ischemic, the subepicardium compared with the subendocardium that is the more ischemic zone (Figure 5.28) (see Chapter 5, Section “From cellular electrogram to human ECG”).

How can we record the electrical activity of the heart?

There are various methods used to record the electrical activity of the heart. The best known method, the one we examine in this book, is **electrocardiography**. **An alternative method**, rarely used in clinical practice today but **very useful in understanding ECG curves and therefore helpful in learning about ECGs**, is **vectorcardiography**.

The latter and other methods will be briefly discussed in Chapter 25. These include, among others, body mapping, late potentials, and esophageal and intracavitary electrocardiography. In addition, normal ECGs can be recorded during exercise and in long recordings (ECG monitoring, Holter technology, telemetry, etc.) (Braunwald et al. 2012; Camm et al. 2006; Fuster et al. 2010). For more information about different techniques, consult our books *Clinical Arrhythmology* (Bayés de Luna and Baranchuk 2017) and *Electrocardiography in Ischemic Heart Disease* (Fiol-Sala et al. 2020), and other ECG reference books (Macfarlane and Lawrie 1989; Wagner 2001; Gertsch 2004; Surawicz et al. 2008) (see “Recommended Reading”).

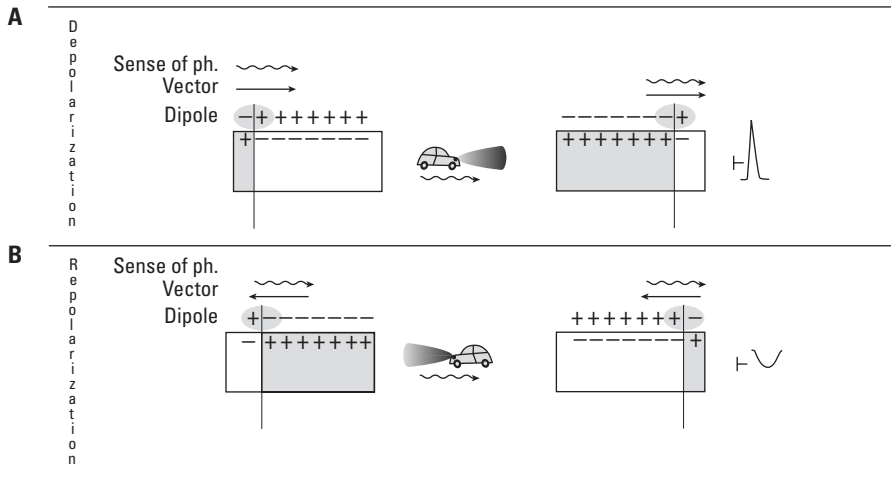


Figure 1.1 Depolarization (A) and repolarization (B) of the dipole in an isolated myocardium cell. We see the onset and end of the depolarization and repolarization processes and how this accounts for the positivity and negativity of corresponding waves (see text and Chapter 5).

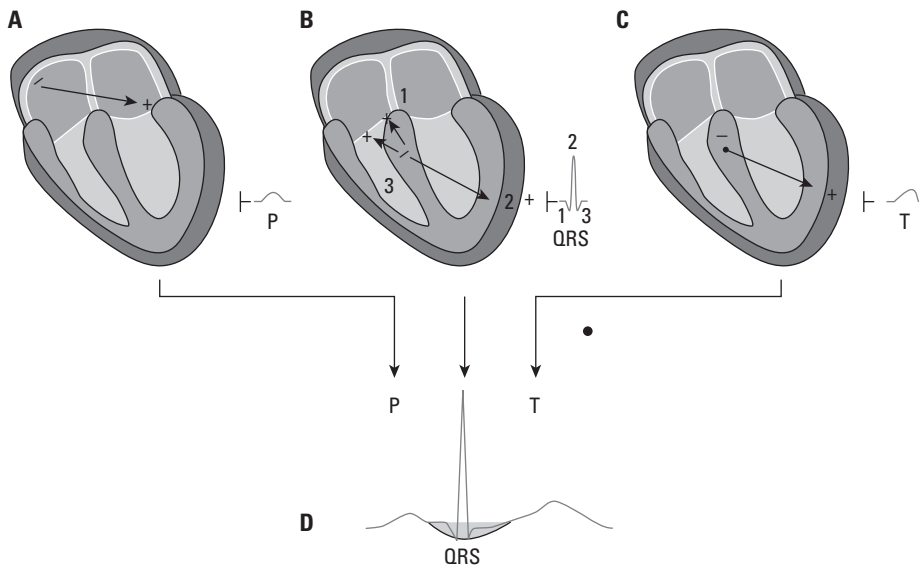


Figure 1.2 The origin of P, QRS, and T deflections (A, B and C). When an electrode located in some zone of LV faces the head (+) of a vector of depolarization (P, QRS) or repolarization (T), it records positivity. When an electrode faces the tail of a vector (-), it records negativity. Atrial repolarization is hidden in the QRS (shadow area) (see text and Chapter 5).

What is the surface ECG?

The ECG is the standard technique used for recording the electrical activity of the heart. **We can record the process of depolarization and repolarization** through recording electrodes (leads) located in various places.

The depolarization process of the heart, atria, and ventricles (see Chapter 5 and Figures 5.16 and 5.18) starts with the formation of a dipole of depolarization (+), which has a vectorial expression (→) that advances through the surface of the myocardium and seeds the entire surface of the myocardial cells with negative charges. A recording electrode facing the head of the vector records positivity (Figure 1.2). Later, **the repolarization process starts** with the formation of a repolarization dipole (+), which also has a vectorial expression. During

this process, the positive charges of the outside surface of the cells are restored.

These two processes relate to specific characteristics of the atria and ventricles (Figure 1.2). The process of atrial depolarization, when recorded on the surface of the body in an area close to the left ventricle (Figure 1.2), presents as a small positive wave called the P wave (∧). This is the expression of the atrial depolarization dipole (vector). **The process of ventricular depolarization**, which occurs later when the stimulus arrives at the ventricles, usually presents as three deflections (↓), known as the QRS complex, caused by the formation of three consecutive dipoles (vectors). The first vector appears as a small and negative deflection because it represents the depolarization of a small area in the septum the first part of LV to be depolarized, and is usually directed upward and to

the right and therefore, recorded from the left ventricle as a small negative deflection ("q wave"). Next, a second important and positive vector is formed, representing the R wave. This is the expression of depolarization in most of the left ventricular mass. The head of this vector faces the recording electrode. Finally, there is a third small vector of ventricular depolarization that depolarizes the upper part of the septum and right ventricle. It is directed upward and to the right and is recorded by the recording electrode in the left ventricle zone as a small negative wave ("s wave") (Figure 1.2).

After depolarization of the atria and ventricles, the **process of repolarization starts**. The repolarization of the atria is usually a smooth curve that remains hidden within the QRS complex. The ventricular repolarization curve appears after the QRS as an isoelectric ST segment and a T wave. This T wave is recorded as a positive wave from the left ventricle electrode because the process of ventricular repolarization, as already mentioned and later explained in detail (see Chapter 5, Section "From cellular electrogram to the human ECG" and Figures 5.24 and 5.25),

appears very differently from what happens in an isolated contractile cell (see Figure 5.9). Repolarization starts on the opposite side to that of depolarization, because it starts in the zone less ischemic of LV that is the subepicardium. Thus, the recording electrode faces the positive part of the dipole, or head of the vector, and will record a positive deflection, even though the dipole moves away from it (Figure 1.2C; see also Figures 5.24 and 5.25). Therefore, repolarization of the left ventricle in a human ECG (the T wave) is recorded as a positive wave, just as occurs with the depolarization complex (QRS) in leads placed close to the left ventricle surface (—).

The successive recording of the ECG is linear and the distance from one P–QRS–T to another can be measured in time. The frequency of this sequence is related to heart rate.

The heart is a three-dimensional organ. In order to see its electrical activity on a two-dimensional piece of paper or screen, it must be projected from at least two planes, the frontal plane (FP) and the horizontal plane (HP) (Figure 1.3).

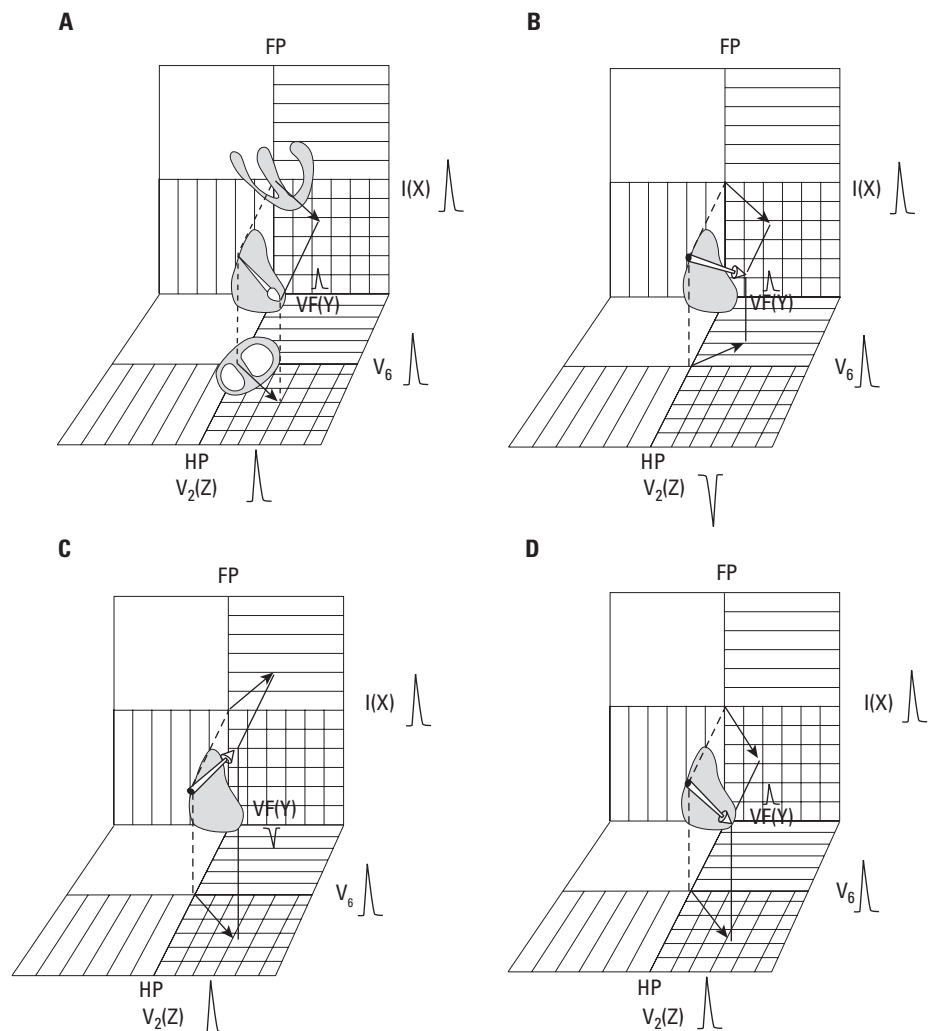


Figure 1.3 Four locations of a vector and their projection in frontal (FP) and horizontal planes (HP). **A** and **B** have the same projection in FP but not in HP. **C** and **D** have the same projection in HP but not in FP. Different positive and negative morphologies appear according to these projections. The locations of the orthogonal leads X, Y, and Z perpendicular to each other are similar to I, aVF, and V2 leads. Vertical lines correspond to the positive hemifields of aVF and V2, and horizontal lines correspond to the positive hemifields of leads I and V6. FP lead I (X) = 0°; VF (Y) = +90°; HP V2 (Z) = +90°; V6 = 0°.

The shape of the ECG varies according to the location (lead) from which the electrical activity is recorded. In general, the electrical activity of the heart is recorded using 12 different leads: six on the FP (I, II, III, aVR, aVL, aVF), located from $+120^\circ$ (III) to -30° (aVL). The aVR is usually recorded in the positive part of the lead that is located in -150° (see Figures 6.10 and 6.11), and six on the HP (V1–V6) located from $+120^\circ$ to 0° (see Chapter 6, Section “Leads” and Figures 6.10 and 6.13).

Each lead has a line that begins where the lead is placed, 0° for lead I or $+90^\circ$ for lead aVF in the FP and 0° for lead V6 and $+90^\circ$ for lead V2 in the HP, for example (see Figure 6.10), and ends at the opposite side of the body, passing through the center of the heart. By tracing each perpendicular line that passes through the center of the heart, we may divide the electrical field of the body into two hemifields for each lead, one positive and one negative (Figure 1.3). A vector that falls into the positive hemifield records positivity, while one that falls into the negative hemifield records negativity. When a vector falls on the line of separation between hemifields, an isodiphasic curve is recorded that is $+-$ or $-+$ according to the sense of rotation of the curve (loop) (see later) (see Figures 6.14 and 6.16).

The different vectors are recorded as positive or negative depending on whether they are projected onto positive or negative hemifields of different leads (Figures 1.3 and 1.5). This is a key concept for understanding the morphology of ECG curves in different leads and is explained in Chapter 6 in more detail (Figure 6.14).

The ECG and its different morphologies can be explained using the following sequence:

Dipole → **Spatial vectors** → **Projection in frontal (FP) and horizontal (HP) planes**

What is vectorcardiography? (See Chapter 25 for more extensive information)

The vectorcardiogram (VCG) is the closed curve or loop that records the entire pathway of an electrical stimulus from the depolarization of the atria (P loop) and ventricles (QRS loop) to the repolarization of the ventricles (T loop). These loops are recorded in FP and HP, as well as in the sagittal plane. The VCG is the result of the joined heads of the multiple vectors that are formed during the consecutive processes of depolarization and repolarization of the heart (Figure 1.4). VCG loops are obtained from three orthogonal (perpendicular to each other) leads, X, Y, and Z, which are placed in positions similar to those of leads I, aVF, and V2, respectively (see Figure 1.4 and Chapter 25).

The VCG curve is a plot of voltage against voltage of the different waves generated by the heart (P, QRS, T loops), and therefore in the VCG loops, it is not possible to measure the time between the beginning of the P loop and the beginning of the QRS loop (PR interval), or the beginning of QRS and the end of the T loop (QT interval). However, we can interrupt the loops of P, QRS, and T by

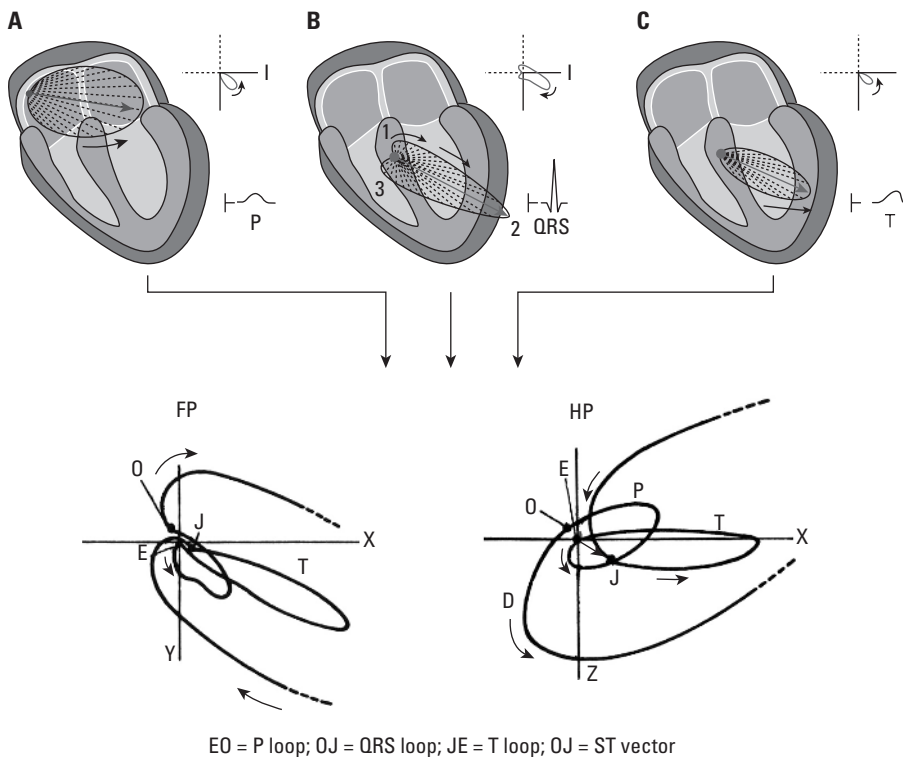


Figure 1.4 The origin of P, QRS, and T loops (A, B and C). The vectorcardiographic curve (D) is the union of the heads of multiple vectors that form during the consecutive processes of depolarization and repolarization (see text and Figure 5.23).

cutting the tracing every 2.5ms, which allows the duration of each loop to be measured (see Figures 10.6–10.10 and 10.22–10.25).

One advantage of the VCG is that the different rotations of the loop can be visualized. It is important to know if the stimulus follows a clockwise or counterclockwise rotation when one complex or wave is biphasic. Figure 1.5B shows how the mean vector of a loop directed to $+0^\circ$ that falls within the limit between the positive or negative hemifields in lead “Y” (aVF) may present a $+-$ ($\downarrow$) or a $-+$ ($\uparrow$) deflection. The direction of the mean vector of the loop does not solve one important problem: a $+-$ deflection is normal, but a $-+$ deflection may be the expression of myocardial infarction (MI). The correct morphology will be shown, however, by the direction of loop rotation (Figure 1.5). In addition, the mean vector of the QRS loop, which expresses the sum of all vectors of depolarization, does not indicate the direction of the small initial and final forces when these forces are opposed to a mean vector (Figure 1.5). However, the small part of the loop (beginning and end) that falls in the opposite hemifield of the main vector can explain the complete ECG morphology with initial (q) and final small (s) deflections (Figures 1.5 and 1.6; see also Figures 7.10 and 7.11).

The VCG can be described using the following sequence:

The head of multiple vectors $\rightarrow$ Spatial loops $\rightarrow$ Projection in FP and HP

ECG–VCG correlation

Bearing in mind the abovementioned information, it is clear that **to better understand the morphology of an ECG, we must consider the stimulus pathway through the heart (VCG loop) in different normal and pathological conditions and identify the projection of these loops in FP and HP**. It is important to understand how the different parts of the loop that fall into the positive or negative hemifields of each lead correspond to the different deflections of an ECG curve (Figures 1.5 and 1.6; see also Figure 5.23) (ECG–VCG correlation). **This allows the ECG curves to be drawn from the VCG loops and vice versa.**

The key concepts around how ECG curves can be obtained from the VCG loops and vice versa (ECG–VCG correlation) are defined using the following sequence: Dipole $\rightarrow$ Vector $\rightarrow$ Loop $\rightarrow$ Projection in different hemifields $\rightarrow$ ECG patterns

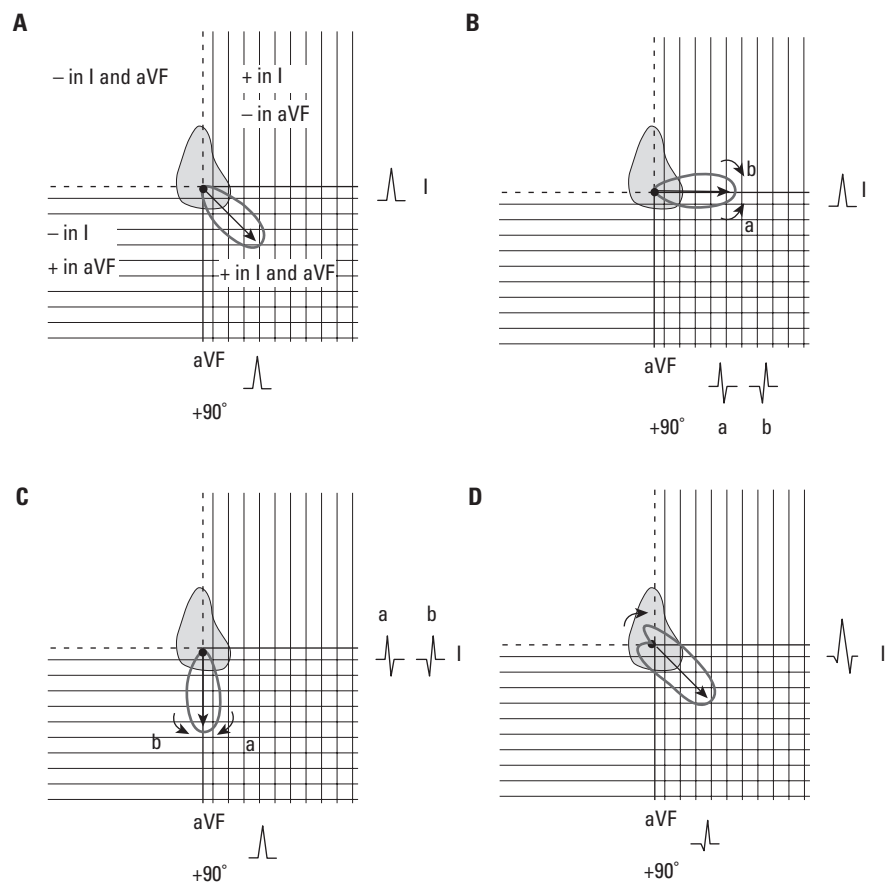


Figure 1.5 The concept of the hemifield. We see how a morphology may be $+-$ or $-+$ with the same vector but a different loop rotation (B and C) (A and B). The recording of the initial and terminal deflections of qRs are well understood with the correlation of the loop and hemifields in D (see I and aVF).

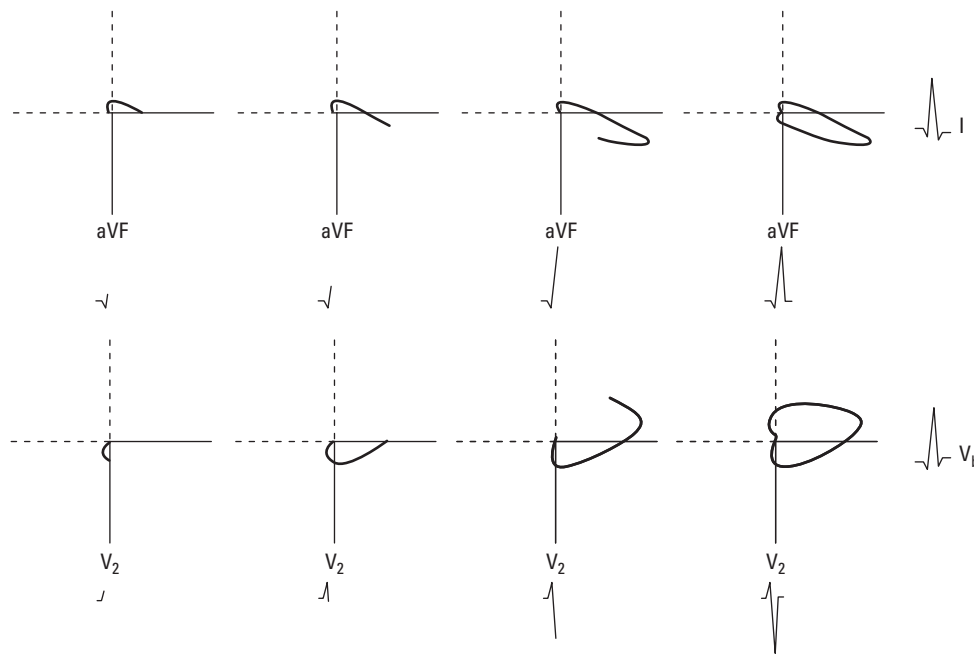


Figure 1.6 Correlation of a vectorcardiographic loop with an electrocardiographic morphology in aVF and V2.

Why do we record ECG curves and not VCG loops?

Although ECG–VCG correlation is used in this book to explain how the different ECG patterns appear, the recording of vectorcardiographic loops for diagnostic purposes is rarely used in clinical practice at present. There are many reasons for the superiority of ECG curves over VCG loops, the main ones being as follows:

- The established diagnostic criteria of ECG in different pathologies are more defined and agreed-upon, compared with the VCG criteria. They are also easier to apply. Furthermore, it has not been clearly demonstrated by experts in ECG/VCG interpretation that VCG criteria provide more diagnostic information than that taken from ECGs, even in an era when VCG criteria were most used (Simonson *et al.* 1967; Rautaharju 1988; Van Bommel *et al.* 1992) (see later).
- VCG loops do not show an appreciation of time (PR and QT interval), as previously mentioned.
- The ECG curves–VCG loop correlation gives us all the detailed information obtainable from VCG loops. In fact, if the origin of the ECG curve interpretation based on the projection of VCG loops in the positive and negative hemifields of different leads (ECG–VCG correlation) is understood and used, we are able to derive the same information that VCGs provide by just looking at the ECG. For example, it has been reported (Benchimol *et al.* 1972) that VCGs are essential for the diagnosis of superoanterior fascicular block (hemiblock) associated with inferior MI. However, as discussed in Chapter 13,

the same information can be obtained from the ECG if we recognize the exact pathway of the stimulus in both cases (inferior MI alone or associated with hemiblock) and we make a good ECG–VCG correlation (see Figure 13.99). Furthermore, many details provided by amplified VCG loops (the degree of ST shifts, onset of pre-excitation, characteristics of the P wave, etc.) can also be obtained from surface ECGs through amplification of the ECG waves, if necessary (see Figure 13.24) (Fiol-Sala *et al.* 2020).

- The VCG is not useful in the diagnosis of arrhythmias. Even information about the ectopic P wave may be correctly obtained from ECG curves. However, the recording of P loops may help to know better how is the way of stimulus in situations such as the presence of advanced interatrial block (Bayés de Luna *et al.* 2012).
- Computerization of ECG data and not of VCG has become dominant and, despite current limitations, it has a great future. However, as we see later on (Chapter 3, Section “Limitations”), it is necessary to improve the results of ECG computerization with better technology and the inclusion of new data (clinical setting, different scores, etc.), and we hope that in the near future become a substitute of individualized interpretations.
- The ECG is used more than the VCG for estimating the size of an MI (Selvester QRS scoring system) (Selvester *et al.* 1972; Wagner *et al.* 1982). However, in the era of ECG-imaging correlations, it is necessary to improve the methodology of QRS score measurement to obtain a better correlation with contrast-enhanced cardiovascular magnetic resonance measurements (see later and Chapter 3, Section “Limitations”).

- **ECG and not VCG patterns have already been correlated with imaging techniques**, especially coronarography and contrast-enhanced cardiovascular magnetic resonance (CE-CMR). The correlation of ECG patterns with coronarography has allowed us to better locate the occlusion and determine the severity of ischemia in different types of acute coronary syndromes (leads with different ST shifts) (Sclarovsky 1999; Wellens *et al.* 2004; Fiol-Sala *et al.* 2020). It is also possible to obtain better classification and location of Q-wave MI (leads with abnormal Q or R waves as mirror image) using CE-CMR-ECG correlation (Bayés de Luna *et al.* 2006a, 2006b; Cino *et al.* 2006; Bayés de Luna 2007; Rovai *et al.* 2007; Van der Weg *et al.* 2009; Fiol-Sala *et al.* 2020). Similar correlations have not been done with VCG loops. Currently, a good estimation of infarction size using CE-CMR has been obtained (Kim *et al.* 1999; Moon *et al.* 2004). However, the correlation of infarct size measurement performed by surface ECG (QRS scoring system) (Selvester *et al.* 1972) with CE-CMR is not very consistent, and the CE-CMR shows larger values than the QRS score estimation (Weir *et al.* 2010). We hope that in the future it will be possible to improve these results with new equations (see Chapter 3, Section “The future”). Good results have also recently been shown by Montant *et al.* (2010) using a contrast-enhanced three-dimensional echocardiography compared with CE-CMR to identify and quantify myocardial scars (positive and negative predictive value (PV) > 90%).

- **Young physicians should realize that ECG–VCG correlation is a basis for better understanding ECG curves.** This does not mean that they need to know specific VCG criteria, such as the number of milliseconds the loop is going up and down, because these data obtained from the VCG does not add too much diagnostic information. Therefore, a recorded VCG loop is not clinically necessary. However, **it is important to remember that the ECG–VCG correlation is a key point for better understanding of how ECG curves are originated** (see below).

- **Currently, there are very few devices that still correctly record VCG curves.** At the Electrophysiology Congress held in 2009, titled with the subheading “VCG symposium,” it was decided that this subheading should be suppressed (Macfarlane 2009). “Signum temporis” stated the first organizers (Sobieszczkańska and Jagielski 2010). The number of VCG papers published in Medline in the 1970s and 1980s reached more than 800 per decade; today, in the first decade of this century, there are fewer than 60 and in the last decade the number is probably still lower.

- It appears that the VCG loops taken from the orthogonal leads do not give much more information of clinical interest than a conventional 12-lead surface ECG. They are also time consuming and need special devices. Although we consider that VCG is not very useful for diagnosis, we are sure that the VCG loops are very useful

for teaching and research purposes (Kors *et al.* 1990, 1998; Rautaharju *et al.* 2007; Pérez Riera 2009; Lazzara 2010). It may be that incorporating VCG loops synthesized directly from 12-lead surface ECG recordings would be an interesting option (see Chapter 3).

Why do we use ECG–VCG correlations to understand ECG patterns?

Electrocardiography and vectorcardiography are two methods for recording the electrical activity of the heart. As explained above, the **ECG is a linear curve** based on the positive and negative deflections recorded when an electrode faces the head or the tail of a depolarization and repolarization dipole, the expression of which is a vector, from leads placed in frontal and horizontal planes. The **VCG is a loop** that represents the outline of the joining of multiple dipoles (vectors) formed along the electrical stimulus pathway through the heart. The projection of these loops in frontal and horizontal planes is a closed curve that is different in morphology from the linear curves of an ECG. Both ECG curves and VCG loops, however, are completely connected so that the **ECG curve may be easily deduced from the VCG loop, and vice versa** (see ECG–VCG correlation, Figures 1.5 and 1.6). As already mentioned, this approach is considered to be the best way to understand both the normal ECG and all the morphological changes that different pathologies introduce to the ECG.

The correlation between VCG loops and projection of this on different hemifields to understand the ECG pattern (**dipole → vector → loop → hemifield sequence**) will no doubt remain a cornerstone of the teaching of the ECG (Grant and Estes 1952; Sodi-Pallares and Calder 1956; Cooksey *et al.* 1957; Cabrera 1958; Gertsch 2004).

1. **The deduction of the ECG from the VCG loops is crucial** to better recognizing how both normal ECG curves and the many ECG changes found in different heart diseases and under special circumstances originate.
2. Although the deductive method for teaching electrocardiography is fundamentally based on the correlation that exists between ECG curves and VCG loops, VCG criteria are not used for diagnosis.
3. **In the majority of diagrams** used to show the usefulness of VCG loop–ECG wave correlations, the pathway of the electrical stimulus is represented as a curve with a **continuous line**. When we **record original VCG tracings**, however, dashes every 2.5 ms in the VCG loops are shown. Examples of this may be seen throughout this book (see, for example, Figures 11.25, 11.36, and 11.40).

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