Rapid ECG Interpretation

Third Edition

M. Gabriel Khan

MD, FRCP (LONDON), FRCP (C), FACP, FACC

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Rapid ECG Interpretation
CONTemporary Cardiology

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Rapid ECG Interpretation

Third Edition

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With a Foreword by
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Humana Press
Totowa, New Jersey
To my wife, Brigid
The electrocardiogram (ECG) is the first test performed on most cardiac patients—one that helps make the first part of the diagnosis and one that can frequently direct treatment decisions. Thus, for any physician, a solid understanding of the ECG is critical. Learning the basics and subtleties of the ECG is a right of passage for all physicians and healthcare providers during their training.

So, what would we want from a book on ECGs? Ideally, such a book would be comprehensive, yet concise, practically oriented, and explain pathophysiology and its application to practice.

Dr. Khan has written such a book. *Rapid ECG Interpretation* is comprehensive, yet concise, and very practically oriented. More important, it takes a step-by-step approach, walking the reader through a thorough evaluation of the ECG. This, as many of us have been taught, is the “right” way to look at an ECG. This edition includes a new opening chapter that covers basic concepts. This quickly orients the reader to the physiology, anatomy, and geometry of the electrical system of the heart.

After reviewing each component of the ECG, the next section describes the unique ECG patterns of specific cardiac conditions, including pulmonary embolism and long QT syndrome. This is followed by a chapter with each of the arrhythmias. Finally, Dr. Khan includes an invaluable section—an ECG Board Review and Self-Assessment Quiz. With this, the reader can really see if the basic concepts and ECG fundamentals have been learned.

Dr. Khan is to be congratulated on an outstanding text that will help readers at all levels become very familiar and facile in rapid interpretation of the ECG.

*Christopher P. Cannon, MD*  
TIMI Study Group, Brigham and Women’s Hospital  
Harvard Medical School, Boston, MA
A new approach for the interpretation of the electrocardiogram (ECG), *a step-by-step method for the accurate interpretation of the ECG*, is outlined in this text.

The most important addition in the second edition of *Rapid ECG Interpretation* was a new chapter, Basic Concepts. This chapter gives considerable practical details with 16 instructive illustrations so that the reader can fully understand the genesis of each wave and deflection of the ECG and the reason 12 carefully positioned leads are needed to capture 12 views of the heart’s electrical currents and vector forces. Also, more than 35 new ECG tracings were added to the chapters that discuss topics that will be of value to postgraduates and internists.

The major addition in this third edition is a new chapter: ECG Board Self-Assessment Quiz. The chapter provides 90 selected ECG tracings that should sharpen the skills of all who wish to interpret ECGs. This small-volume text contains more than 320 ECGs and instructive illustrations.

The ECG is the oldest cardiologic test, but even 100 years after its inception, it continues as the most commonly used cardiologic test. Despite the advent of expensive and sophisticated alternatives, the ECG remains the most reliable tool for the confirmation of acute myocardial infarction (MI). The ECG—not CK-MB, troponins, echocardiogram, or SPECT or PET scan—dictates the timely administration of lifesaving PCI or thrombolytic therapy. There is no test to rival the ECG in the diagnosis of arrhythmias, which is a common and bothersome clinical cardiologic problem. Also, the clinical diagnosis of pericarditis and myocardial ischemia is made mainly by ECG findings.

This text gives a systematic step-by-step approach but departs somewhat from the conventional sequence and gives steps that are consistent with the changes in cardiology practice that have evolved over the past decade. The early diagnosis of acute MI depends on astute observation for ST segment changes. New terms have emerged: ST elevation MI and non–ST elevation MI (non–Q wave MI). The ST segment holds the key to the diagnosis. Currently, ambulance crews are being trained in Europe, the United States, and Canada to recognize ST segment abnormalities and to make the diagnosis of ST elevation MI (STEMI)
and non–ST elevation MI. Thus, patients can be rapidly shuttled to special cardiac centers for coronary angiography and angioplasty/stent or thrombolytic therapy; rapid triage in emergency rooms is crucial. These lifesaving measures depend on the accurate and rapid interpretation of the ECG by clinicians who must be adequately trained to interpret tracings.

This text describes ST segment abnormalities in detail. For example, the recent observation that ST segment elevation in lead aVR (a commonly ignored lead) is a marker for left main coronary artery (LMCA) occlusion is of lifesaving value. Because LMCA occlusion is a serious condition, any noninvasive diagnostic clue represents a valuable addition to our armamentarium. Thus, only after detailed assessment of the ST segment is completed are the QRS complex, T waves, atrial and ventricular hypertrophy, and lastly the axis assessed. This change in the analytical sequence is necessary so that the most crucial diagnoses can be made accurately and rapidly.

In addition, the standard teaching is for the interpreter to assess all leads and all deflections and waves before entertaining diagnoses. This text gives presumptive diagnoses as soon as a clue is uncovered in the tracing. Also, a few rare but life-threatening conditions are excluded early in the assessment sequence. For example, although Wolff-Parkinson-White (WPW) syndrome is uncommon, it is an important diagnosis that may be missed by computer analysis and by physicians. Because WPW syndrome is a result of widening of the QRS complex, it is logical to consider this diagnosis in the same framework as bundle branch blocks; this approach avoids the danger and embarrassment of missing the diagnosis. No text considers WPW syndrome in the assessment of the 10 essential ECG features. Most important, it is imperative to exclude mimics of MI early in the sequence. WPW syndrome may mimic MI. Right bundle branch block (RBBB) may reveal Q waves in leads III and aVF that may be erroneously interpreted as MI. Left bundle branch block (LBBB) may mimic MI and must be quickly documented because its presence hinders the accurate diagnosis of acute coronary syndromes. Furthermore, the ECG manifestation of acute MI may be a new LBBB pattern. Thus, the assessment for blocks is performed early, in step 2 of the 11 steps outlined.

Because RBBB and LBBB are best revealed in leads V1 and V2, the clinician is advised to screen these leads before assessing other leads. The text advises the clinician or senior resident that the assessment of V1 and V2 may assist with the diagnosis of Brugada syndrome and right ventricular dysplasia, which may display particular forms of right
bundle branch block and recently have been shown to be causes of sudden death in young adults. Many rare syndromes are described in medicine, but those that cause sudden death should be made familiar to trainees and clinicians. We should not fear divulging information about such rare syndromes at an early stage to students and residents, because these topics may serve to motivate them to higher levels of excellence.

This text presents a unique 11-step method for accurate and rapid ECG interpretation in a user-friendly synopsis format. Medical house staff should welcome this step-by-step method, because it simplifies ECG interpretation and provides for greater accuracy than the approaches given in texts published over the past 50 years. The succinct writing style allows a wealth of information to be presented in a small text that is highlighted with bullets to allow for rapid retrieval. The 11 steps are illustrated in algorithms and outlined in Chapter 2 with references to later chapters, each of which expands on one of the steps and provides advanced material for senior internal medicine residents, cardiology residents, and internists. The text moves rapidly from basics to advanced material.

All diagnostic ECG criteria are given with relevant and instructive ECGs, providing a quick review or refresher for proficiency tests and for physicians preparing for the ECG section of the Cardiovascular Diseases Board Examination. This text can be a valuable tool for all those who wish to be proficient in the interpretation of ECGs.

M. Gabriel Khan
I had the privilege of borrowing several ECG tracings from *Electrocardiography in Clinical Practice* by the late Dr. Te-Chuan Chou and from *Practical Electrocardiography* by Dr. Henry J.L. Marriott; I am grateful to these authors. A special note of thanks to my editor, Paul Dolgert.
Dr. M. Gabriel Khan is a cardiologist at the Ottawa Hospital and an Associate Professor of Medicine, at the University of Ottawa. Dr. Khan graduated MB, BCh, with First-Class Honours at The Queen’s University of Belfast. He was appointed Staff Physician in charge of a Clinical Teaching Unit at the Ottawa General Hospital and is a Fellow of the American College of Cardiology, the American College of Physicians, and the Royal College of Physicians of London and Canada. He is the author of On Call Cardiology, 3rd ed., Elsevier, Philadelphia (2006); Heart Disease Diagnosis and Therapy, 2nd ed., Humana Press (2006); Cardiac and Pulmonary Management, Elsevier, Philadelphia, PA (1993), Medical Diagnosis and Therapy (1994), Heart Attacks, Hypertension and Heart Drugs (1986), Heart Trouble Encyclopedia (1996), and Encyclopedia of Heart Diseases (2006), Academic Press/Elsevier, San Diego; and Cardiac Drug Therapy, 7th ed., Humana Press (2007).

Dr. Khan’s books have been translated into Chinese, Czech, Farsi, French, German, Greek, Italian, Japanese, Polish, Portuguese, Russian, Spanish, and Turkish. He has built a reputation as a clinician-teacher and has become an internationally acclaimed cardiologist through his writings.

His peers have acknowledged the merits of his books by their reviews of Cardiac Drug Therapy: Review of the 5th edition in Clinical Cardiology: “this is an excellent book. It succeeds in being practical while presenting the major evidence in relation to its recommendations. Of value to absolutely anyone who prescribes for cardiac patients on the day-to-day basis. From the trainee to the experienced consultant, all will find it useful. The author stamps his authority very clearly throughout the text by very clear assertions of his own recommendations even when these recommendations are at odds with those of official bodies. In such situations the ‘official’ recommendations are also stated but clearly are not preferred.”

And for the fourth edition a cardiologist reviewer states that it is “by far the best handbook on cardiovascular therapeutics I have ever had the pleasure of reading. The information given in each chapter is up-to-date, accurate, clearly written, eminently readable and well referenced.”
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1

Basic Concepts

CONTENTS

- Electrical Activity of the Heart
- Electrocardiogram
- How Are the Waves of the Electrocardiogram Produced?
- Why Use 12 Leads to Record the Electrocardiogram?
- Leads and Electrodes
- Genesis of the QRS Complex
- Vector Forces
- QRS Normal Variants and Abnormalities

ELECTRICAL ACTIVITY OF THE HEART

Each contraction of the heart is preceded by excitation waves of electrical activity that originate in the sinoatrial (SA) node. Figure 1-1 depicts the radial spread of activation from the SA node. The waves of electrical activity spread through the atria and reach the atrioventricular (AV) node. Note that the SA node tracing shows no steady resting potential, as does the ventricular muscle tracing. The SA node’s spontaneous depolarization and repolarization provides a unique and miraculous automatic pacemaker stimulus that activates the atria and the AV node, which conducts the activation current down the bundle branches to activate the ventricular muscle mass. Cardiac cells outside the SA node normally do not exhibit spontaneous depolarization; thus they must be activated.

Depolarization

In a resting cardiac muscle cell, molecules dissociate into positively charged ions on the outer surface and negatively charged ions on the
inner surface of the cell membrane; the cell is in an electrically balanced or polarized resting state (Fig. 1-2).

- When the cell is stimulated by an excitatory electrical wave, the negative ions migrate to the outer surface of the cell and the positively charged ions pass into the cell; this reversal of polarity is called depolarization (see Fig. 1-2).
- If an electrode is placed so that the depolarization wave flows toward the electrode, a galvanometer will record an upward or positive deflection (Fig. 1-3).

**Fig. 1-1.** Electrical activation of the heart by the sinoatrial (SA) node. The current of activation (arrows) spreads radially from the SA node across the atria to the atroventricular (AV) node and down the bundle branches to the ventricular muscle and Purkinje network. The SA node tracing shows no steady resting potential and is characterized by spontaneous depolarization.
**Fig. 1-2.** A, Resting cell: Positive ions on the outer surface and negative ions inside equal an electrically balanced or polarized cell. B, Depolarized cell: Negative ions on the outer surface and positive ions inside. C, Repolarization of cell: Positive ions return to the outside.

**Fig. 1-3.** Recording of the effects of electrical activation process. A, Current flows toward the electrode produce a positive upward deflection. B, Current flows away from the electrode produce a negative deflection. C, Current flows toward an electrode placed at a distance produce a positive but smaller amplitude deflection than in (A).
• When a depolarization current is directed away from an electrode, a negative or downward deflection is recorded (see Fig. 1-3).

**Repolarization**

• During a recovery period, positively charged ions return to the outer surface and negatively charged ions move into the cell. The electrical balance of the cell is restored; this process is called repolarization (see Fig. 1-2).
• The transfer of sodium (Na\(^+\)) and potassium (K\(^+\)) ions across the cell membrane plays an important role in generating cardiac electrical activity. In Fig. 1-4, the relative magnitudes of the concentration of Na\(^+\) and K\(^+\) ions are indicated. Intracellular concentration of K\(^+\) is 30 times greater than extracellular K\(^+\). Na\(^+\) concentration is 30 times less inside the cell than outside. Because of this ionic composition, the membrane of the resting cardiac fiber is in an electrically balanced or polarized state. The potential difference across the cell membrane can be measured by a microelectrode and is observed on an oscilloscope to be \(-90\) mV.

**The Action Potential**

• The inward Na\(^+\) current results in a change in transmembrane potential; results in depolarization; and is shown as the upstroke, phase 0 of the action potential. With a decrease in Na\(^+\) and K\(^+\) permeability, the membrane potential remains close to 0; this represents phases 1 and 2 of the action potential (see Fig. 1-4). The Na\(^+\)-K\(^+\)-ATPase (adenosine triphosphatase) sodium pump, depicted in Fig. 1-4, pumps Na\(^+\) from the intracellular to the extracellular fluid compartment; K\(^+\) passes from the extracellular fluid to the intracellular fluid.
• Phase 3 is the phase of rapid repolarization and is followed by a period of stable resting potential, phase 4 of the action potential.

The appreciation of these four phases is important for the understanding of abnormal heart rhythms (arrhythmias) and the therapeutic actions of antiarrhythmics. For example, digoxin or excess catecholamines increase the slope of spontaneous phase 4 depolarization and therefore increase automaticity of ectopic pacemakers (Fig. 1-5); β-blockers cause inhibition or depression of spontaneous phase 4 diastolic depolarization and thus suppress catecholamine-induced arrhythmias, particularly those related to ischemia. Digitalis causes inhibition of the cellular Na\(^+\) pump, which causes increased intracellular Na\(^+\), which is then exchanged for calcium via the Na\(^+\)-calcium exchanger. Increased intracellular calcium during cardiac systole increases myocardial muscle contractility. Digitalis toxicity causes cellular calcium overload that potentiates arrhythmias.
<table>
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| Na⁺ | Na⁺ Na⁺ | Na⁺ |
| K⁺ to Na⁺ ions 30:1 | K⁺ K⁺ K⁺ | Na⁺ |

**Fig. 1-4.** A simplified concept of ionic exchange; the polarized, depolarized, and repolarized state of a myocardial cell; and the action potential. An electrical current arriving at the cell causes positively charged ions to cross the cell membrane, which causes depolarization, followed by repolarization, which generates an action potential: phases 0, 1, 2, 3, and 4. This electrical event traverses the heart and initiates mechanical systole, or the heartbeat (see also Fig. 1-7).

**Fig. 1-5.** Effects of catecholamines, digoxin, and β-blockers on spontaneous phase 4 depolarization. β-Blockers inhibit or decrease spontaneous phase 4 depolarization caused by catecholamines, especially that caused by ischemia.
Sinoatrial Node

The SA node is unique and has no steady resting potential. After repolarization, slow, spontaneous depolarization occurs during phase 4 that causes the automaticity of the SA node fibers (see SA node waveform in Fig. 1-1). Thus, the unique pacemaker provides individuals with an automatic infinitesimal current that sets the heart’s electrical activity and contractions. The SA discharge rate, usually 50 to 100 per minute, is under autonomic, chemical, and hormonal influence.

Atrioventricular Node

The AV node provides a necessary physiologic delay of the electrical currents, which allows the atria to fill the ventricles with blood before ventricular systole.

• From the AV node and bundle of His, the excitatory electrical current rapidly traverses the right and left bundle branches, the specialized conductive tissues of the ventricles, and the Purkinje system, and the entire ventricular muscle is depolarized (see Fig. 1-1).

• Depolarization spreads down the intraventricular septum toward the apex of the heart and then along the free wall of the left ventricular myocardium; it always proceeds from the endocardium toward the pericardium. The specialized fine arborization of branches that constitute the Purkinje network spreads over the endocardial surfaces of the ventricles.

• The transient halt and slowing of conduction through the specialized AV node fibers play an important protective role in patients with atrial flutter and atrial fibrillation. In these common conditions, a rapid atrial rate of approximately 300 to 600 beats/min reaches the AV node; this AV “tollgate” reduces the electrical traffic that reaches the superhighway that traverses the ventricles to approximately 120 to 180 beats/min, and serious life-threatening events are prevented.

ELECTROCARDIOGRAM

The heart muscle is made up of several thousand muscle elements, about $10^{10}$ cells. Each instant of depolarization or repolarization represents different stages of activity for a large number of cells. The electrical activity of each element can be represented by a vector force.

• A vector is defined as a force that can be represented by direction and magnitude. The sum total of cardiac vectors is considered the electrical activity of the entire heart (Fig. 1-6). The ECG records the sequence of such instantaneous vectors.
The heart muscle is arranged in three muscle masses: the intraventricular septum, a large left ventricular muscle mass, and a small right ventricular muscle mass. The magnitude or amplitude of the deflections recorded is influenced by the size of the muscle mass depolarized and the distance from the recording electrode (see Figs. 1-3 and 1-6).

The graphic representation of the heart’s electrical activity recorded through electrodes positioned at strategic points on the body constitutes the electrocardiogram (ECG). The recording of the electrical currents, their direction, and their magnitude, as well as the rate of the heart’s contractions, is made by the machine and electrocardiograph, which is essentially a galvanometer whose deflections are recorded on moving, specially prepared paper.

The ECG is the recording obtained, and to simplify interpretation, it suffices to state that the ECG displays the following:

• Three major deflections or waves: the P wave, the QRS complex, and a T wave (Fig. 1-7).
• Two time intervals of clinical importance: the PR interval and QRS duration (see Fig. 1-7).
The ST segment, a most important ECG component. The study of abnormalities of the ST segment reveals the early diagnosis of acute myocardial infarction (MI) and myocardial ischemia. Thus, this text devotes an in-depth chapter to abnormalities of the ST segment and does so early in the interpretive sequence; that is, before analysis of abnormalities of the P wave, ventricular hypertrophy, QRS abnormalities, and the electrical axis, all of which are discussed early in other textbooks. This approach simplifies ECG interpretation and is a strategy that is now embraced by physicians who render acute care to patients with acute MI and those with myocardial ischemia.

**HOW ARE THE WAVES OF THE ELECTROCARDIOGRAM PRODUCED?**

**P Wave**

The early part of the P wave represents the electrical activity generated by the right atrium; the middle portion of the P wave represents
completion of right atrial activation and initiation of left atrial activation; and the late portion is generated by the left atrium. The P wave is the first deflection recorded and is a small, smooth, rounded deflection that precedes the spiky-looking QRS complex (Fig. 1-8). (See Chapter 3 for an in-depth discussion of P waves.)

**PR Interval**

The PR interval involves the time required for the electrical impulse to advance from the atria through the AV node, bundle of His, bundle branches, and Purkinje fibers until the ventricular muscle begins to depolarize (see Figs. 1-7 and 1-8).

**QRS Complex**

The QRS complex represents the spread of electrical activation through the ventricular myocardium; the resultant electrical forces generated from ventricular depolarization is recorded on the ECG as a spiky deflection (see Figs. 1-7 and 1-8). The sharp, pointed deflections are labeled QRS regardless of whether they are positive (upward) or negative (downward).

Figure 1-9 indicates the conventional labeling of the QRS complex: q or Q, r or R, s or S, depending on the size of the components that

![Fig. 1-8](image-url)

**Fig. 1-8.** Relationship of P wave, PR interval, and QRS complex to activation from the sinoatrial (SA) node, atrioventricular (AV) node, bundle of His (HIS), and bundle branches. Note that the normal ST segment curves imperceptibly into the ascending limb of the T wave and is not a horizontal line.
may be recorded (i.e., those influenced by the electrode position) and the direction of the resultant vector forces. Large deflections are labeled with uppercase letters.

The genesis of the QRS complex is intricate and is better understood after the reader has been presented with information on leads and lead positions and why 12 leads are used to capture 12 views of the heart’s electrical activity. Thus the genesis of the QRS complex is discussed at the end of this chapter.

**ST Segment**

The ST segment is the segment that lies between the end of the QRS complex and the beginning of the T wave (see Figs. 1-7 and 1-8). It represents the period when all parts of the ventricles are in the depolarized state or a stage in which the terminal depolarization and the starting repolarization are superimposed and thus neutralize each other. Early repolarization may encroach on the ST segment to a variable degree. The part at which the ST segment takes off from the QRS complex is called the J, or the junction point. The ST segment normally curves imperceptibly into the ascending limb of the T wave and should not form a horizontal line nor form a sharp angle with the proximal
limb of the T wave. The student must be aware of this important diagnostic point.

This important diagnostic ECG segment is discussed in detail in Chapters 2 and 5.

**T Wave**

The T wave represents electrical recovery, repolarization of the ventricles, and is a broad, rounded wave (see Figs. 1-7 and 1-8). The T wave follows each QRS complex and is separated from the QRS by an interval that is constant for that ECG. Because ventricular recovery proceeds in the general direction of ventricular excitation, the polarity of the resultant T vector is similar to that of the QRS vector. The T wave is recorded during ventricular systole, whereas the QRS occurs immediately before mechanical systole.

- The T wave process is energy consuming, but the QRS process is not. During repolarization, cellular metabolic work and energy consumption occurs to accomplish the ionic flux associated with repolarization. Thus several metabolic, hemodynamic, and physiologic factors may affect the repolarization process and alter the morphology of the T wave. The student or clinician interpreting ECGs should be aware of the normal variations in T wave morphology and the influence of a host of factors that may alter the T wave and lead to erroneous diagnoses.
- Levine listed approximately 67 causes for T wave changes, which include the patient drinking ice water, eating, exercising, or fasting or having infections, fever, tachycardia, anoxia, shock, electrolyte derangements, acidemia, alkalemia, hormonal imbalances, subarachnoid hemorrhage, or drug or alcohol abuse.

Because of the unreliable diagnostic yield derived from the scrutiny of T waves, further details on this topic are relegated to Chapter 8.

**U Wave**

The U wave is a wave that follows the T wave and is observed only in the ECG tracings of some individuals. It is a small, often indistinct wave, and its source is uncertain (see Chapter 8).

**WHY USE 12 LEADS TO RECORD THE ELECTROCARDIOGRAM?**

Einthoven’s discovery in 1901 was of paramount importance. His landmark paper was published in 1901, and a further paper on the galvanometric registration of the human electrocardiogram was published
in 1903. However, the initial work of Galvani (1791), Muller (1856), and Waller (1887) initiated Einthoven’s accomplishment. Einthoven recognized that the heart possessed electrical activity, and he recorded this activity using two sensors attached to the two forearms and connected to a silver wire that ran between two poles of a large permanent magnet. He noted that the silver wire moved rhythmically with the heartbeats, but to visualize the small movements Einthoven shone a light beam across the wire, and the wavy movements of the wire were recorded on moving photographic paper. Einthoven recorded the waves and spiky deflection and labeled the first smooth, rounded wave, P; the spiky deflection, QRS; and the last recorded wave, T.

- Einthoven labeled the waves P, Q, R, S, and T; his lettering obeyed the convention used by geometricians: curved lines were labeled beginning with P, and points on straight lines were labeled beginning with Q.

Einthoven, Sir Thomas Lewis, and others correlated the ECG waves with the contracting heart and correlated that the P wave was related to atrial contraction and that the QRS deflection was associated with ventricular contraction. Improvements in the quality of recordings resulted from the immense work and technique of Frank Wilson, who studied with Lewis and, in Michigan (1934), described the unipolar leads that include the precordial V leads and VR, VL, and VF.

**LEADS AND ELECTRODES**

*Why Are 12 Leads Necessary?*

Figure 1-10 shows the infinite number of electrode positions arranged in a continuous circle, at the center of which is the origin of the depolarization wave. The illustration indicates that the electrode position has a profound influence on the size, or amplitude, of the recording.

- Twelve ECG leads are used to obtain 12 views of the heart’s electrical activity. The heart may be considered to lie at the center of an equilateral triangle (Fig. 1-11). The leads attached to the limbs, the limb leads, act as linear conductors and have virtually identical voltages at all points along their lengths. The limbs can be regarded as extensions of a lead wire. Thus, the left arm electrode placed at the wrist, arm, or shoulder displays the same ECG record. Because the limb leads act as linear conductors, the effective sensing points and electrode locations are at the left and right shoulders and left groin, but are usually positioned and labeled as follows:
**Fig. 1-10.** Effect of varied electrode positions on the amplitude and direction of deflections recorded: Leads between C and A or C and B give positive deflection less than at C. Leads at D and A or D and B record negative deflection of varying size. The line AB is perpendicular to the electrical current.

**Fig. 1-11.** The heart depicted as a three-muscle mass that lies in the center of an equilateral triangle. The two shoulders and left groin are sensing positions.
Rapid ECG Interpretation

- R = right arm lead
- L = left arm lead
- F = foot = left leg lead

These leads lie along the frontal plane of the body and display action potential only in the frontal plane. (See discussion of frontal plane axis in Step 9 in Chapter 2 and in Chapter 9.)

Two important concepts must be reemphasized:

- If the excitatory depolarization head of the current (vector force) flows toward a unipolar electrode, a galvanometer will record an upward or positive deflection (see Fig. 1-3).
- When an excitatory depolarization process is directed away from the electrode, a downward or negative deflection is recorded (see Fig. 1-3).

Figure 1-11 displays deflections that can be recorded by limb leads R, L, and F. The main electrical current of activation flows toward the F (left leg) electrode and records an upward or positive deflection of large amplitude. The current flows away from the right shoulder (R) electrode and records a downward deflection. The right shoulder lead (R) looks into the interior of the heart toward the endocardium, and as mentioned previously, the current of activation flows from the endocardium and traverses the myocardium toward the pericardium and thus displays a negative deflection. The student should notice that aVR is always relatively negative and aVF is always relatively positive.

- Lead L at the left shoulder or left arm usually displays a small positive or equiphasic deflection, but the heart hangs in the chest and is subject to rotational changes, and the main current direction may be altered; thus, this lead may show a large-amplitude positive deflection in some individuals, and a negative deflection if the heart’s position is vertical.

**Why Augmented Leads?**

- Why is a V added to the R, L, and F? These leads are termed unipolar limb leads, but voltage measurements are virtually never unipolar. The connection formed by attaching the R, L, and F electrodes together acts as a reference connection, and the lead formed is termed a V lead (V = voltage); thus the convention VR, VL, and VF, and the V is also used for the leads positioned on the chest, V₁ to V₆.
- Goldberger (1942) augmented Wilson’s unipolar extremity leads that gave low-amplitude records; Goldberger’s strategy increased the amplitude of the deflections by 50%. Thus, the letter a is used to denote the
augmented lead (e.g., $aVL = \text{augmented-voltage left arm lead \ [V = voltage]}$).

**Standard Bipolar Limb Leads I, II, and III**

Figure 1-12 shows the views of the heart obtained by leads I, II, and III.

- Lead I connects the two arms and is formed by connecting L to the positive terminal and R to the negative terminal of the galvanometer; thus, $I = aVL = aVR$. Lead I looks at the heart from the left, inferior to lead $aVL$, the lead of the left shoulder (arm), and displays the electrical tracing produced by a combination of the right arm and left arm electrodes. The right leg electrode is an earth (or ground) and minimizes interference.

- Lead II looks at the heart from a position to the left of the left groin, foot lead F (see Fig. 1-12).

- Lead III looks at the heart from a position to the right of the left groin, foot lead F. Thus, leads II, III, and $aVF$ look at the inferior surface of the heart from different angles, and they usually show some similarities. Lead III is the most unreliable of the leads II, III, and $aVF$. Thus, many errors are made from the observation of the QRS and T wave in lead

![Fig. 1-12. Standard limb leads I, II, and III. Note that $aVF$ leads II and III look at the inferior surface of the heart and deflections show minor variation. Leads I and $aVL$ look at the anterolateral aspect of the heart.](image-url)
III. Normal yet pathologic-appearing Q waves and T wave inversion may be observed frequently in lead III as a normal variant (see Chapters 6 and 8).

- The six leads display six photographs of the heart’s electrical activity taken from six angles (one every 30 degrees). The six leads can be visualized as traversing a flat plane over the chest of the patient (i.e., the frontal plane). Importantly, if only two of the six leads are recorded, the most informative pair are I and aVF.

**Vertical Versus Horizontal Heart Position**

Figure 1-13 shows the changes in QRS waveform caused by alteration of the position of the heart:

- Both aVR and aVL face the ventricular cavity and show a QS complex.
- A qR complex in lead aVL indicates a horizontal heart position, and the QRS morphology in aVL resembles that in V$_5$.
- A qR complex in aVF and a QS complex in aVL indicate a vertical heart position, and the QRS morphologies in leads aVF and V$_5$ resemble each other.
- The position of the heart varies between horizontal and vertical.

**Chest Leads/Precordial or V Leads**

The six chest leads give six more views of the heart’s electrical activity and vector forces; they are positioned around the anterior and left chest wall in a horizontal plane. Figures 1-14 and 1-15 indicate the position of the precordial chest leads that overlie the right and left ventricles.

V$_1$ and V$_2$ face and lie close to the wall of the right ventricle. V$_2$ and V$_3$ lie near the intraventricular septum. V$_4$ and V$_3$ look at the anterior parts of the left ventricle, with V$_4$ close to the apex. V$_5$ and V$_6$ (leads I and aVL) view the anterolateral region of the left ventricle and often appear similar to each other. The recording in lead aVL, however, varies depending on a horizontal or vertical heart position. If V$_7$ is taken, it is positioned in the posterior axillary line.

The precordial electrodes V$_1$ to V$_6$ are so close to the electrical currents of the heart that no augmentation is necessary. Lead V$_6$ is far around (in the axilla) and is separated from the free wall of the left ventricle by a significant distance. Figure 1-15 indicates the approximate relationship of the ventricular myocardium and the precordial chest leads V$_1$ to V$_6$. 
Fig. 1-13. Changes in deflections with the heart in a vertical (A) and a horizontal (B) position. In the vertical position (A), both the aVR and aVL face the cavity of the ventricles and record a QS complex. A QRS complex in aVF indicates a heart that is positioned close to vertical; qRS in aVL indicates a horizontal heart position (B).
Figure 1-14. Position of precordial chest leads.

Figure 1-16 reemphasizes that the position of leads aVL and aVF and other limb leads are in the same frontal plane. The chest leads V₁ to V₆ encircle the left thorax in a horizontal plane (see Fig. 1-15).

Caution: The entire chest, with the heart within it, acts as a volume conductor, and thus voltage varies appreciably at locations only a centimeter apart. Therefore, the leads placed on the chest wall V₁ to V₆

Fig. 1-15. Magnetic resonance image of heart to illustrate approximate relationship of chest electrodes to cardiac chambers. Points 1 to 6 represent sites of the six precordial electrodes V₁ to V₆. RA, right atrium; RV, right ventricle; LV, left ventricle; RL, right lung; LL, left lung; A, aorta. (From Marriott HJL: *Practical Electrocardiography*, 8th ed., Philadelphia, 1988, Williams & Wilkins.)
must be positioned meticulously so that when the ECG is repeated days or years later, accurate comparison can be made. Caution is required so that the $V_5$ and $V_6$ electrodes are not placed too anteriorly. $V_5$ must be placed in the anterior axillary line; $V_6$ should be placed in the midaxillary line at the level of $V_4$ in the fifth intercostal space or in line with the apex beat.

*Fig. 1-16.* aVF and aVL are in the same frontal plane. The chest leads encircle the left thorax in a horizontal plane.
If lead V₃ is placed too close to V₂ or is positioned near the left third intercostal space, no positive deflection or a reduced-amplitude R wave may be recorded, which can falsely simulate an anterior MI. If lead V₂ is positioned too close to V₁, no R wave may be recorded in lead V₂, and the erroneous diagnosis of anteroseptal MI may be made. These errors are made commonly in the ECGs of females, and they may be interpreted as “loss or poor R wave in V₃, consider anteroseptal MI.” An ECG with faulty recording may lead to serious errors in interpretation.

GENESIS OF THE QRS COMPLEX

Understanding the genesis of the QRS complex is a fundamental step. Knowledge of the normal sequence of activation or depolarization of the ventricles is crucial to an understanding of the normal and abnormal QRS complex. The accurate diagnosis of acute and old MI, right and left bundle branch block, hemiblocks, and ventricular hypertrophy depends on knowledge of resultant vectors that dictate the components of the QRS complex.

The electrical impulse that proceeds from the SA node activates the atria, producing the P wave, the first wave of the ECG. The electrical impulse is briefly slowed in the AV node, then progresses rapidly down the bundle of His, the right and left bundle branches, and the Purkinje fibers of the ventricular myocardium. The spread of the electrical impulses through the septum and ventricular muscle is called depolarization, which produces the QRS complex of the ECG.

VECTOR FORCES

The electrical impulses that activate each area of heart muscle have direction and magnitude and can be represented by a vector force. The direction of the resultant force can be represented by an arrow, the length of which represents the magnitude of the force. The term vector does not imply vector cardiography.

Three Caveats

A vector describes a force in terms of its duration and magnitude. The following three caveats must be considered:

1. An electrical impulse traveling toward an electrode causes a positive deflection or R wave (Fig. 1-17).
2. When the impulse is traveling away from the electrode, a negative deflection occurs (i.e., an S, a small Q, or a QS wave is recorded).
3. Three resultant vectors dictate the inscription of the QRS complex.
Vector I

- The ventricular septum is activated from left to right; electrodes or leads positioned over the right ventricle (V₁ or V₂) face the wave of depolarization and inscribe a positive wave, a small R wave (see Fig. 1-17).
- Because the force of the activation impulse (vector I) is small, the positive deflection is small; the R wave recorded in V₁ and V₂ is small and ranges from 1 to 4 mm in V₁ and from 1 to 7 mm in V₂ in normal individuals older than age 30 years (see Table 2-1). Incorrect lead placement of V₁, V₂, and V₃, especially in women, may cause the ECG tracing to falsely show diminished or loss of R or r waves in V₂ and V₃, which is often incorrectly interpreted as anteroseptal MI.
- The initial depolarizing current travels away from leads V₅ and V₆ and thus inscribes a small negative deflection, a small Q wave in leads V₅, V₆, and I.

Vector II

- After septal depolarization, both ventricular walls are activated simultaneously.
- The impulse depolarizes the thin-walled right ventricle; however, the magnitude of the forces is small in comparison with the forces that activate the thick left ventricular free wall. Thus, the resultant force, vector II, is directed toward and through the left ventricular free wall (see Fig. 1-17).

Fig. 1-17. Genesis of the normal QRS complex. V(I), vector I produces a small r wave in leads V₁ and V₂, Q in leads V₅ and V₆; V(II), vector II produces an S wave in lead V₁ and an R wave in lead V₅ or V₆; V(III), vector III produces the terminal S in leads V₅ and V₆ and the terminal r or r' in V₁, V₂, and aVR; V₁, lead V₁ electrode; V₆, lead V₆ electrode; R, right ventricle muscle mass; L, left ventricle muscle mass; S, septum. (From Khan, M. Gabriel: On Call Cardiology, 3rd ed., Philadelphia, 2006, WB Saunders, Elsevier Science.)
• The resultant force, vector II, is indicated by an arrow directed toward the left; the electrodes V5 and V6 face the left ventricle and show a positive wave, an R wave, the height of which depends on the thickness of the left ventricular muscle. The height of the R wave in V4 through V6 ranges from 10 to 25 mm and may exceed 30 mm in individuals with left ventricular hypertrophy and in normal subjects younger than age 25 years. The R wave in V4 through V6 is lost or is reduced to less than 3 mm in height in patients with anterior MI.
• Because the electrical current represented by vector II travels away from an electrode overlying the right ventricle, V1 and V2 record a negative deflection, an S wave.
• The larger the left ventricular muscle, the deeper the S wave in V1 and V2.

**Vector III**

• Activation of the posterobasal right and left ventricular free walls and the basal right septal mass, including the crista supraventricularis, represents vector III.
• The resultant force is directed to the right, is small in magnitude, and may record a small S wave in V5 and V6 and a terminal r′ wave in lead V1 or V2; thus, an Rsr′ pattern in V1 may occur in normal individuals.

**QRS NORMAL VARIANTS AND ABNORMALITIES**

*Clockwise and Counterclockwise Rotation*

• Variations in the normal QRS configuration are shown in Fig. 1-18. If the heart undergoes strong clockwise or counterclockwise rotation, changes in QRS morphology occur. Failure to recognize these normal variants may result in incorrect interpretation of the ECG.
• With clockwise rotation, the V1 electrode, like aVR, faces the cavity of the ventricle and records a QS complex; therefore Q waves can occur as a normal finding if there is extreme clockwise rotation of the heart (see Fig. 1-18). The normal Q wave in V6 disappears because the resultant force of the initial vector I is not directed toward the electrode V1.

*Tall R Waves*

• If the left ventricle is hypertrophied, the magnitude of vector force II increases; thus, a tall R wave is recorded in V5 and V6 (see Fig. 2-25). With right ventricular hypertrophy, the magnitude of vector force I increases and tall R waves occur in V1 and V2 (see Fig. 2-26 and Table 2-3).
A myocardial infarct is an area of necrotic cells caused by the blood supply to that area of heart muscle being cut off. The necrotic area is an electrical window:

- If there is necrosis of the left ventricular muscle facing electrodes $V_4$ through $V_6$, no R waves (i.e., Q waves) will be produced (see Figs. 1-18, 2-18, and 2-19) or the R in $V_3$ through $V_5$ may be considerably decreased; this is termed poor R wave progression (see Chapter 6). Loss of R waves or poor R wave progression in leads $V_3$ through $V_5$ may indicate anterior MI (see Fig. 2-18).
- R waves should increase in amplitude from $V_2$ through $V_4$. If R waves are present in leads $V_1$ and $V_2$ and are not present in $V_4$ through $V_6$, a diagnosis of anterolateral MI should be considered (see Fig. 2-19).

**Q Waves**

A myocardial infarct is an area of necrotic cells caused by the blood supply to that area of heart muscle being cut off. The necrotic area is an electrical window:
• Infarction of the ventricular septum causes the loss of vector I, as well as loss of the normal R wave in leads V₁ and V₂ (i.e., pathologic Q waves), indicating anteroseptal infarction (see Fig. 2-18A).
• Normal Q waves are less than 0.04 second in duration and are less than 3 mm deep. These small Q waves are recorded when a small activation current is directed away from the electrode. Small Q waves are found normally in leads V₅, V₆, and I (see Fig. 2-2). Changes in the position of the heart may cause small Q waves in leads III, aVF, and aVL; with extreme counterclockwise rotation, small Q waves occur in V₁ through V₆ (see Fig. 1-18).
• Leads III and aVL may record narrow Q waves up to 10 mm deep in normal individuals. In lead III, the Q wave can be normally ≤0.04 second wide (see Fig. 2-2D). In all other leads, Q waves should be considered normal if they are less than 0.04 second wide and less than 3 mm deep. If Q waves are not observed in leads II or aVF, a Q wave in lead III should be considered normal (see Table 2-1).
• Hypertrophy of the interventricular septum occurs in hypertrophic cardiomyopathy, and the ECG often reveals deep Q waves that can mimic MI (see discussion of pathologic Q waves and QS patterns given under Step 6 in Chapter 6).
• When the arm leads are inadvertently placed on the legs and vice versa, Q waves are recorded in leads II, III, and aVF; consider this technical error if there is no deflection in lead I (see Fig. 2-39).
• Replacement of ventricular muscle by tumor; fibrosis; or amyloid, sarcoid, or other granuloma may cause an electrical window and Q waves that simulate infarction.
• Lead aVR normally records a negative QRS or QS complex because aVR looks into the cavity of the ventricle and faces the endocardial surface; the activating current flows from endocardium to pericardium (see Fig. 2-2B).
• See Chapter 5 for the recently observed importance of ST segment elevation in aVR and the diagnosis of acute MI.
INTRODUCTION

Conventional Sequence Regarding Interpretation

The time-honored advice to students and staff is as follows: In every ECG, the following features should be examined systematically:

From: Contemporary Cardiology: Rapid ECG Interpretation, 3e
by: M. Gabriel Khan © Humana Press Inc., Totowa, NJ

2
• Rate
• Rhythm
• P wave morphology
• PR interval
• QRS interval, QRS complex morphology
• ST segment
• T wave
• Electrical axis
• U wave, and QT duration

Some authors, advise the following sequence:
Assess: rate, rhythm, axis, hypertrophy, infarction

but this is not the conventional teaching of cardiology tutors.

**New Sequence for Interpretation**

This text departs somewhat from the conventional sequence and gives a new approach consistent with the changes in cardiology practice that have evolved over the past decade. The early diagnosis of acute MI depends on astute observation for abnormal changes in the ST segment. Determination of creatinine kinase MB (CK-MB) and troponins is not relevant in the early phase of acute MI, because these cardiac enzymes are not elevated and are nondiagnostic within the crucial first hour of onset of MI. The door-to-needle or balloon time must be minimized if maximal life-saving is to be achieved. Diagnosis depends on symptoms and ST segment changes. Thus, this text rushes the interpreter to the assessment of ST segment morphology and suggests an 11-step method or sequence for the rapid yet accurate interpretation of ECGs.

**BRIEF HIGHLIGHTS OF AN 11-STEP METHOD**

Figure 2-1 defines the ECG waveform; Fig. 2-2A–F shows features of the normal ECG; and Table 2-1 gives normal ECG intervals and parameters.

An 11-step method is advised to ensure accurate, yet rapid, interpretation of the ECG. Algorithms, illustrations, and many sample ECGs make the 11 steps easy to understand and apply. The 11 steps are briefly outlined in this chapter, and each step receives in-depth coverage in later chapters, which also give advanced diagnostic features for postgraduates.
Fig. 2-1. Sodium influx, potassium efflux, the action potential, and the ECG. (From Khan, M. Gabriel: On Call Cardiology, 3rd ed., Philadelphia, 2006, WB Saunders, Elsevier Science.)

Fig. 2-2. A, Chest leads V₁ through V₆.
Fig. 2-2.  Continued  

B, Limb leads I through aVF. Sinus rhythm, rate 65 beats/min; PR interval, 0.14 second; QRS duration, 0.08 second; QT interval, 0.36 second; axis, +30 degrees.  

C, Chest leads of a normal ECG with a QRS complex in V2 that is positive, indicating early transition. Compare with (A), in which transition is normal, occurring in lead V5; tall R waves in V1 and V2 are not caused by posterior infarction (see Table 2-3). Heart rate, 75 beats/min (see Table 2-2). Note normal small Q wave in V4 through V6.
Fig. 2-2. D, Limb leads of a normal ECG showing a deep but normal Q wave in lead III (see Table 2-1 for normal parameters). E, Leads V_4 through V_6 show small, normal Q waves less than 4 mm deep; leads V_1 through V_3 show normal R wave progression.
Continued. F. Normal ECG, sinus rhythm 75 beats/min; PR interval, 0.16; QRS duration, 0.08; normal QRS axis +60 degrees; QT interval, 0.35. The small notch on the R wave of leads II, III, and aVF is a normal finding in some individuals and does not indicate intraventricular conduction delay (see Chapter 4).
Table 2-1

Important Normal ECG Intervals and Parameters*  

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>PR interval</td>
<td>0.12 to 0.2 second (up to 0.22 second in adults).</td>
</tr>
<tr>
<td>P waves</td>
<td>&lt;3 small squares (0.12 second) in duration, and amplitude &lt;3 mm. Upright in lead I, inverted in aVR (if opposite, suspect reversed arm leads† or dextrocardia) (see Step 6, Figs. 2-21 and 2-36).</td>
</tr>
<tr>
<td>QRS duration</td>
<td>0.05 to 0.1 second; ≥0.1 second, consider incomplete LBBB, incomplete RBBB, or WPW syndrome (see Steps 2 and 3, Figs. 2-4, 2-9, and 2-10).</td>
</tr>
<tr>
<td>Q waves</td>
<td>Normally present in aVR; occasionally in V₁ or in aVL (vertical heart) (see Chapter 6). Often present in lead III: should be ≤0.04 second duration. Other leads except lead I: &lt;0.04 second duration and ≤3 mm deep; lead I ≤1.5 mm in patients older than age 30. Q waves may be up to 5 mm deep in several leads in individuals age &lt;30.</td>
</tr>
<tr>
<td>R waves</td>
<td>V₁: 0 to 15 mm, age 12 to 20 (see Table 2-3). 0 to 8 mm, age 20 to 30. 0 to 6 mm, age &gt;30.‡ V₂: 0.2 to 12 mm, age &lt;30.‡ (see Step 5, Fig. 2-16). V₃: 1 to 20 mm, age &gt;30.‡</td>
</tr>
<tr>
<td>ST segment</td>
<td>Isoelectric or &lt;1 mm elevation in limb leads and &lt;1 mm in precordial leads except for normal variant (see Step 4, Fig. 2-12).</td>
</tr>
<tr>
<td>T wave</td>
<td>Inverted in aVR; upright in I, II, and V₃ through V₆. Variable in III, aVF, aVL, V₁, and V₂ (see Step 8, Fig. 2-27).</td>
</tr>
<tr>
<td>Axis</td>
<td>O degrees to +110 degrees age &lt;40. −30 degrees to +90 degrees age &gt;40 (see Step 9, Fig. 2-30).</td>
</tr>
<tr>
<td>QT interval</td>
<td>See Table 2-5.</td>
</tr>
</tbody>
</table>

*ECG paper speed 25 mm/s.  
†Precordial leads remain normal.  
‡Age >30 is relevant to the diagnosis of myocardial infarction (see Fig. 2-20 and compare with Fig. 2-18, poor R wave progression).
Step 1: see Fig. 2-3.
Assess:
• Rhythm, then the rate.
• Note that rhythm is assessed before rate, because it is clinically more important, and a normal rate of 60 to 100 beats/min is easily spotted.

Step 2: see Fig. 2-4.
Assess:
• PR and QRS intervals for blocks.
• Widening of the QRS duration suggests right bundle branch block (RBBB) or left bundle branch block (LBBB) (see Table 2-1 and Fig. 2-4).

Step 3: see Fig. 2-9.
If the QRS duration is increased in the absence of LBBB or RBBB, assess:
• For nonspecific intraventricular conduction delay (IVCD), a cause of which is Wolff-Parkinson-White (WPW) syndrome (see Fig. 2-9).
• Although WPW syndrome is uncommon, it is an important diagnosis that may be missed by computer analysis and by physicians. Because WPW syndrome is a cause of widening of the QRS complex, it is logical to consider this diagnosis in the same frame as bundle branch blocks; this approach avoids the embarrassment of missing the diagnosis. No other text considers WPW syndrome in the assessment of the 10 essential ECG features, and conventional teaching does not give the approach outlined in Step 3.
• Most importantly, it is imperative to exclude mimics of MI early in the assessment sequence. WPW syndrome may mimic MI. RBBB may reveal Q waves in leads III and aVF that may be erroneously interpreted as MI. The diagnosis of LBBB must be documented quickly, because the presence of LBBB obviates many diagnoses, particularly ischemia and hypertrophy, and the diagnosis of MI is difficult.
• Because ECG changes of bundle branch block may be observed in V1 and V2, the reader is requested to first focus on V1 and V2. Importantly, V1 usually reveals the morphology of P waves and is an excellent lead for the assessment of sinus rhythm and arrhythmia. Thus, sinus rhythm or rhythm disturbances can be rapidly documented; in addition, the PR interval can be assessed, and left atrial enlargement may be revealed.
• The thorough assessment of V₁ and V₂ provides considerable information.
• In addition, the assessment of V₁, V₂/V₃ may assist with the diagnosis of Brugada syndrome and right ventricular dysplasia, which may display particular forms of RBBB and have been shown to be causes of sudden death in young adults. We should not fear to divulge rare syndromes at an early stage to students, because these topics may serve to motivate them to higher levels of excellence. The steps that discuss these rare but important topics are directed to senior trainees and internists. It is logical to discuss basics mixed with advanced material because this may appeal to students and medical residents.

**Step 4: see Fig. 2-12.**

Assess:

• The all-important ST segment.
• The early diagnosis of acute MI depends on observation for ST segment changes. New terms have emerged: ST elevation MI (STEMI) and non–ST elevation MI (previously termed non–Q wave MI). The ST segment holds the key to the diagnosis. This text describes ST segment abnormalities in detail in this chapter and provides further discussion in Chapter 5.

**Step 5: see Fig. 2-16.**

Assess:

• For pathologic Q waves, which, with the prior assessment of the ST segment should determine the presence or absence of new or old MI.
• Search the V leads for the loss of R waves or poor R wave progression, which may indicate MI, lead placement errors, or other cause (see later discussion, figures in this chapter, and Chapter 6).

**Step 6: see Figs. 2-21 and 2-22.**

Assess:

• P waves for atrial hypertrophy.

**Step 7: see Fig. 2-24.**

Assess:

• For left ventricular hypertrophy (LVH) and right ventricular hypertrophy (RVH).
Step 8: see Fig. 2-27.
Assess:
• T waves for inversion, which can have many causes (see later discussion in this chapter and Chapter 8).

Step 9: see Fig. 2-30.
Assess:
• The axis and for fascicular blocks.
• The axis provides no specific diagnosis and is of ancillary assistance only. In the 21st century, I believe conventional teaching should change a little. We should not lose sight of the fact that medical students and interns are bright individuals who desire to move quickly to clinical problem solving. Thus boring topics, particularly difficult ones to grasp such as axis determination, which provides little diagnostic yield, should be assessed after most others. Thus determination of the axis is relegated to Step 9.

Step 10: see Fig. 2-32.
Assess:
• Miscellaneous conditions, such as long QT, pericarditis, pacing, and pulmonary embolism (see later discussion and Chapter 10).

Step 11: see Fig. 2-37.
Assess:
• For arrhythmia.
• Step 11 is indeed Step 1 if an abnormal rhythm is revealed in Step 1: assessment of rhythm (see later discussion in this chapter and detailed coverage in Chapter 11).

Switching the Sequence

Most importantly, these steps can be switched. After the assessment of the important ST segment in Step 4 and for Q waves indicative of acute or old MI in Step 5, Step 7 can switch with Step 9. Thus, the conventional approach is restored, with assessment of the P wave followed by that of the T wave, axis, hypertrophy, and miscellaneous conditions. Therefore, in essence, this text covers the 11 ECG features systematically with minor changes to the conventional approach and
offers relevant and important diagnoses during the sequence, which allows the reader to interpret ECGs with greater accuracy.

Close attention to the 11 steps for ECG interpretation outlined in this chapter and reference to detailed explanations given in subsequent chapters should allow students, staff, and practicing clinicians to be competent interpreters of most ECGs. Accurate, yet rapid, interpretation of the ECG requires a methodic approach.

**THE NORMAL ELECTROCARDIOGRAM**

Figure 2-2A–F shows normal ECG tracings. Figure 2-1 and Table 2-1 list important ECG intervals and parameters. The ECG interpretation should end with one of the following statements:

- Normal ECG
- ECG within normal limits
- Borderline ECG
- Abnormal ECG

**STEP 1: ASSESS RHYTHM AND RATE (FIG. 2-3)**

Focus on leads V1, V2, and II (see Fig. 2-2). Leads V1 and II are best for visualization of P waves to determine the presence of sinus rhythm or an arrhythmia, and V1 and V2 are best to observe for bundle branch block. If P waves are not clearly visible in V1, assess them in lead II, which usually shows well-formed P waves. Identification of the P wave and then the RR intervals allows the interpreter to discover immediately whether the rhythm is sinus or other and to take the following steps:

- Confirm, if the rhythm is sinus, that the RR intervals are equidistant (see Fig. 2-2A), that the P wave is positive in lead II, and that the PP intervals are equidistant and equal to the RR interval.
- Do an arrhythmia assessment if the rhythm is abnormal (see Fig. 2-3, Step 11 [Fig. 2-37], and Chapter 11).
- Determine the heart rate (Table 2-2).

**STEP 2: ASSESS INTERVALS AND BLOCKS (FIG. 2-4)**

- Determine the PR interval; if it is abnormal (>0.2 second), consider first-degree atrioventricular (AV) block (Table 2-1).
- Assess the QRS duration for bundle branch block; if it is ≥0.12 second, bundle branch block is present; assess both V1 and V6. Understanding the genesis of the QRS complex is an essential step and clarifies the ECG manifestations of bundle branch blocks (see Figs. 2-5 to 2-8 and Chapter 4).
**STEP 1**

Look at P waves and RR intervals in leads II and V1. Look at leads V1 and V2; best for bundle branch block.

Determine

Rhythm

Sinus?

Yes

No

Abnormal rhythm

Rate

(see Table 2-2)

Do arrhythmia assessment

(see Step 11 and Chapter 11)

VPBs or APBs* Narrow QRS tachycardia

(Figure 2-37) Wide QRS tachycardia

(Figure 2-38) Bradyarrhythmia

(Chapter 11)

*Ventricular premature beats, atrial premature beats

**Fig. 2-3.** Step-by-step method for accurate ECG interpretation. Step 1: Assess rhythm and rate.

---

**Right Bundle Branch Block**

The ECG criteria for RBBB are as follows:

- QRS duration ≥0.12 second.
- M-shaped complex in V1 and V2.
- Slurred S wave in leads 1, V5, V6; and an S wave that is of greater amplitude (length) than the preceding R wave (*see* Figs. 2-4, 2-6, and 2-7 and Chapter 4, Fig 4-2).

**Left Bundle Branch Block**

The ECG criteria for LBBB are as follows:

- QRS duration ≥0.12 second.
- A small R or QS wave in V1 and V2.
Table 2-2

**Determination of Heart Rate**

<table>
<thead>
<tr>
<th>Heart rate (bpm)</th>
<th>Number of large squares (bold boxes) in one RR interval*</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>300</td>
</tr>
<tr>
<td>1.5</td>
<td>200</td>
</tr>
<tr>
<td>2</td>
<td>150</td>
</tr>
<tr>
<td>3</td>
<td>100</td>
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<td>4</td>
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<td>60</td>
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<tr>
<td>6</td>
<td>50</td>
</tr>
<tr>
<td>7</td>
<td>42</td>
</tr>
<tr>
<td>8</td>
<td>38</td>
</tr>
<tr>
<td>9</td>
<td>33</td>
</tr>
<tr>
<td>10</td>
<td>30</td>
</tr>
<tr>
<td>Number of QRS complexes in 6 seconds†</td>
<td></td>
</tr>
<tr>
<td>5 × 10</td>
<td>50</td>
</tr>
<tr>
<td>6</td>
<td>60</td>
</tr>
<tr>
<td>7</td>
<td>70</td>
</tr>
<tr>
<td>10</td>
<td>100</td>
</tr>
<tr>
<td>15</td>
<td>150</td>
</tr>
<tr>
<td>20</td>
<td>200</td>
</tr>
</tbody>
</table>

*Normal paper speed 25 mm/s. One large box or five small squares (0.2 second) = 300 bpm (see Fig. 2-2C); four large boxes = 75 bpm.

†If the ECG paper has markers at 3-second intervals, count the number of QRS complexes in two of these 3-second periods (6 seconds) and multiply by 10 (see Fig. 2-2C). This method is advisable if there is bradycardia or irregular rhythm. For 5-second interval, multiply the number of QRS complexes by 12.

For regular rhythm: start with a complex that lies on a bold vertical grid line.

Rate = 300 bpm ÷ number of large boxes (0.2 second) in one RR interval.

Normal rate is between 60 bpm (five boxes) and 100 bpm (three boxes); therefore, no need to calculate exact rate.

Or: rate = 1,500 ÷ number of small (1 mm, 0.04 second) squares in one RR interval.

- A notched R wave in leads 1, V₅, and V₆ (see Figs. 2-4 and 2-8 and Chapter 4).

In the presence of LBBB, vector forces are deranged and the ECG cannot be used for the diagnosis of ischemia or ventricular hypertrophy. The diagnosis of acute MI in the presence of LBBB is difficult to make and can be erroneous (see discussion of LBBB and acute MI in Chapter 6).
STEP 2

INTERVALS
(see Table 2-1)

QRS duration
PR > 0.2 sec

= 0.12 sec

Yes
No
Normal

First-degree AV block

BLOCKS

QRS configuration

V₁ and V₂

V₁

RBBB

V₆

slurred S

(see also Figures 2-6 and 2-7 and text on RBBB, Chapter 4)

V₁

LBBB

(see also Figure 2-8 and text on LBBB, Chapter 4)

V₆

notched R

Fig. 2-4. Step-by-step method for accurate ECG interpretation. Step 2: Assess intervals and blocks.
V(I) = vector I produces a small r wave in leads V1 and V2, Q in leads V5 and V6.
V(II) = vector II produces an S wave in lead V1 and an R wave in lead V5 or V6.
V(III) = vector III produces the terminal S in leads V5 and V6 and the terminal
r or r’ in V1, V2, and aVR.

V1 = lead V1 electrode.
V5 = lead V5 electrode.
R = right ventricle muscle mass.
L = left ventricle muscle mass.
S = septum.

**Fig. 2-5.** Vectors I, II, and III, labeled V(I), V(II), and V(III), underlie the genesis of the normal QRS complex. (From Khan, M. Gabriel: *On Call Cardiology*, 3rd ed., Philadelphia, 2006, WB Saunders, Elsevier Science.)

**Fig. 2-6.** Genesis of the QRS complex in right bundle branch block. (From Khan, M. Gabriel: *On Call Cardiology*, 3rd ed., Philadelphia, 2006, WB Saunders, Elsevier Science.)
Fig. 2-7. A, QRS duration in $V_1 \geq 0.12$ second; RSR’ (M-shaped complex) in $V_1$; and wide, slurred S wave in leads 1, $V_5$, and $V_6$ indicate right bundle branch block.
Fig. 2-7. B. Limb leads; slurred, wide S wave in lead I, and the amplitude (length or duration) of the S wave is greater than the preceding R wave.
Fig. 2-8. A, The contribution of vectors I, II, and III, labeled V(I), V(II), and V(III), to the genesis of left bundle branch block. (From Khan, M. Gabriel: On Call Cardiology, 3rd ed., Philadelphia, 2006, WB Saunders, Elsevier Science.). B, QRS duration >0.12 second; small R waves in V₁ to V₃; and notched R wave in V₅ indicate left bundle branch block.

STEP 3: ASSESS FOR NONSPECIFIC INTRAVENTRICULAR CONDUCTION DELAY AND WOLFF-PARKINSON-WHITE SYNDROME (FIG. 2-9)

- If the QRS duration is prolonged ≥0.11 second and bundle branch block appears to be present but is atypical, consider WPW syndrome, particularly if there is a tall R wave in V₁ and V₂ (Table 2-3; see also Figs. 11-27 and 11-28).
- Assess for a short PR interval ≤0.12 second and for a delta wave (Fig. 2-10).
Fig. 2-8. Continued
**STEP 3**

QRS ≥0.11 But not typical RBBB or LBBB configuration

1. Atypical RBBB
2. Atypical LBBB

CONSIDER

Spot for Delta Wave + PR ≤0.12* (see Figure 2-10)

Present

3. WPW SYNDROME*

1, 2, and 3 excluded

Diagnosis

INTRAVENTRICULAR CONDUCTION DELAY (IVCD)

Consider Brugada syndrome and right ventricular dysplasia, rare forms of RBBB patterns; see Chapter 4

* = in ~20% the QRS is <0.11 seconds; in ~23% the PR interval is 0.12 second or slightly longer

Fig. 2-9. Step-by-step method for accurate ECG interpretation. Step 3: Assess for nonspecific intraventricular conduction delay and Wolff-Parkinson-White (WPW) syndrome (see Figs. 2-10 and 2-11). (See Chapter 4 for Brugada syndrome and right ventricular dysplasia, rare forms of RBBB patterns, and Chapter 11 for Wolff-Parkinson-White syndrome.)

WPW syndrome may mimic an inferior MI (see Chapters 6 and 11 for discussion of WPW syndrome). If WPW syndrome, RBBB, or LBBB is not present, interpret as nonspecific intraventricular conduction delay (IVCD) and assess for the presence of electronic pacing (see Figs. 2-7, 2-8, 2-11, and 10-16).
Table 2-3
Causes of Tall R Waves in V\textsubscript{1} and V\textsubscript{2}

1. Thin chest wall or normal variant, age <20, early transition (see Fig. 2-2C)
2. Right bundle branch block (see Fig. 2-7)
   \textit{Note}: Slurred S wave in leads I, V\textsubscript{5}, and V\textsubscript{6}
3. Right ventricular hypertrophy (see Fig. 7-8)
   No slurred S wave in leads I, V\textsubscript{5}, and V\textsubscript{6}
4. Wolff-Parkinson-White syndrome (see Fig. 2-10)
5. True posterior infarction (see Fig. 6-19)
   \textit{Note}: Associated inferior MI, no slurred S in V\textsubscript{5} and V\textsubscript{6}, and T upright in V\textsubscript{1} and V\textsubscript{2}
6. Hypertrophic cardiomyopathy
7. Duchenne muscular dystrophy
8. Low placement of leads V\textsubscript{1} and V\textsubscript{2}
9. Dextroposition (see Fig. 10-9)

Fig. 2-10. Tall R waves in leads V\textsubscript{1} and V\textsubscript{2}; QRS duration ≥0.11 second; and delta wave in V\textsubscript{3} through V\textsubscript{5} indicate Wolff-Parkinson-White syndrome.
Fig. 2-11. Sinus rhythm 72/min; ventricular premature beats; QRS duration 0.14 s, 140 ms: Intraventricular conduction delay (IVCD). Abnormal ECG.
STEP 4: ASSESS FOR ST SEGMENT ELEVATION OR DEPRESSION (FIG. 2-12)

- Focus on the ST segment for elevation or depression (see Fig. 2-12). ST elevation $\geq 1$ mm (0.1 mV) in two or more contiguous ECG leads in a patient with chest pain indicates ST elevation MI (STEMI). The diagnosis is strengthened if there is reciprocal depression (Fig. 2-13).
- Figure 2-13A shows marked ST elevation in leads II, III, and aVF, with marked reciprocal depression in leads I and aVL, diagnostic of acute inferior MI.
- Figure 2-13B shows marked ST segment elevation in V$_1$ through V$_5$, caused by extensive acute anterior MI.

**STEP 4**

![STEP 4 Diagram](image)

*Reciprocal depression increases probabilities of acute myocardial infarction (MI).

**Fig. 2-12.** Step-by-step method for accurate ECG interpretation. Step 4: Assess for ST segment elevation or depression.
A, Marked ST segment elevation in leads II, III, and aVF with reciprocal depression in leads I and aVL indicate acute inferior infarction. B, Marked ST segment elevation in leads V₁ through V₅ indicates acute anterior infarction. C, Electrocardiogram of a 79-year-old woman with an apparent acute subendocardial myocardial infarction attributed to subtotal occlusion of the left main coronary artery, associated with global hypokinesis and an estimated left ventricular ejection fraction of 10%. ST segment is depressed in leads I, II, III, aVL, aVF, and V₂ through V₆. Apparent “reciprocal” ST segment elevation is seen in leads aVR and V₁. (From Surawicz B, Knilans TK: Chou’s Electrocardiography in Clinical Practice, 5th ed., Philadelphia, 2001, WB Saunders, Elsevier Science.)

- Figure 2-13C shows the ECG of a patient with a subtotal occlusion of the left main coronary artery. Note the ST elevation in aVR is greater than the ST elevation in V₁, a recently identified marker of left main coronary disease. (See Chapter 5, particularly Figs. 5-11 and 5-12, for an in-depth discussion of Step 4: ST segment elevation.)
- Figure 2-14A shows features of non–ST elevation MI (non–Q wave MI).
- Figures 2-14B and 2-14C illustrate ECG features diagnostic of myocardial ischemia.
Fig. 2-13. Continued
A. Upsloping ST depression is nonspecific; commonly seen with tachycardia.

B. Marked ST segment depression and elevated creatine kinase (CK) and CK-MB indicate non–Q wave myocardial infarction. ECG patterns of myocardial ischemia. (From Khan, M. Gabriel: On Call Cardiology, 3rd ed., Philadelphia, 2006, WB Saunders, Elsevier Science.)

C. Leads V₄ through V₆ show ST segment depression; V₄ through V₆ are in keeping with myocardial ischemia from a patient known to have unstable angina.

**Fig. 2-14.** A, Marked ST segment depression and elevated creatine kinase (CK) and CK-MB indicate non–Q wave myocardial infarction. B, ECG patterns of myocardial ischemia. *Upsloping ST depression is nonspecific; commonly seen with tachycardia. (From Khan, M. Gabriel: On Call Cardiology, 3rd ed., Philadelphia, 2006, WB Saunders, Elsevier Science.) C, Leads V₄ through V₆ show ST segment depression; V₄ through V₆ are in keeping with myocardial ischemia from a patient known to have unstable angina.
• Elevation of the ST segment may occur as a normal variant (Fig. 2-15). See Chapters 5 and 6 for further discussion of ST segment abnormalities and MI.

Note: This text advises scrutiny of the ST segment before assessment of T waves, electrical axis, QT interval, and hypertrophy because the diagnosis of acute MI or ischemia is vital and depends on careful assessment of the ST segment.
Exclude other causes of ST elevation:

- Normal variant: 1- to 2-mm ST segment elevation, mainly in leads V\(_2\) through V\(_4\), nonconvex, and with fishhook appearance. Common in African Americans: even 4-mm ST segment elevation (see Fig. 2-15 and sections on acute myocardial infarction in Chapters 5 and 6).
- Coronary artery spasm: ST returns to normal with nitroglycerin or with pain relief.
- LBBB: QRS >0.12 second and typical configuration (see Fig. 2-8B, and Chapter 4).
- Left ventricular aneurysm and known old infarct with old Q waves (see Chapter 6).

**STEP 5: ASSESS FOR PATHOLOGIC Q WAVES (THAT IS, LOSS OF R WAVES) (FIG. 2-16)**

- Assess for the loss of R waves—pathologic Q waves—in leads I, II, III, aVL, and aVF (see Figs. 2-17A and 2-17B and Chapter 6 for detailed discussion).
a. Assess for Q waves, leads I, II, III, aVF, and aVL.

Normal if <0.04 second (one millimeter square = 0.04 second) and ≤3 mm deep, except lead III normal up to 0.04 second and up to 7 mm deep in III and aVL; lead I ≤1.5 mm deep (see Table 2-1)

If abnormal Q II, III, aVF consider inferior MI

I, aVL, V5, V6 determine age of infarct

anterolateral MI (see Chapter 6, Figure 6-15) recent (see Figure 2-17, A) old (see Figure 2-17, B) exclude mimics

hypertrophic cardiomyopathy (see Chapter 6, Figure 6-23) WPW syndrome (see Figure 6-24) borderline Qs, II, III, aVF

b. Assess for R wave progression in V1 through V6 or pathologic Q waves.

R should be

0 to 6 mm in V1*

>0.2 mm in V2 (normal 0.3 to 12 mm)

≥1 mm in V3 (normal 1 to 24 mm)

If poor R progression, consider

late transition (see Figure 2-20)

normal variant (Figure 6-5)

anterior MI LVH (see Figures 2-25 and 6-27 and Chapter 7)

LBBB (QRS ≥0.12) (see Figures 2-8, B, and 6-6 and Chapter 4)

emphysema (see Figures 6-8 and 6-28 and Chapter 6) exclude mimics of MI (see Chapters 5 and 6)

lateral MI (see Figures 2-19 and 6-15)
hypertrophic cardiomyopathy (see Figure 6-23)

recent (see Figures 2-13, B, 2-18, A, and 6-10)

indeterminate† (see Figures 2-18, B, and 6-15)

old (see Figures 2-18, C, and 6-9)

*Age >30; see Chapter 6 and Table 2–1 for exceptions and normal parameters.

†Compare old ECGs.

Fig. 2-16. Step-by-step method for accurate ECG interpretation. Step 5: Assess for pathologic Q waves (i.e., loss of R waves).
Fig. 2-17. A, Loss of R wave in leads III and aVF (i.e., pathologic Q waves associated with marked ST segment elevation in leads III and aVF) and minimal elevation in lead II and reciprocal depression in leads I and aVL indicate typical acute Q wave inferior infarction. B, Wide, deep pathologic Q waves in leads II, III, and aVF and isoelectric ST segment indicate old inferior myocardial infarction. C, Variation in QRS configuration caused by rotation. (From Khan, M. Gabriel: *On Call Cardiology*, 2nd ed., Philadelphia, 2001, WB Saunders, Elsevier Science.)
Fig. 2-17. Continued
• Assess for R wave progression in V2 through V4. Figure 2-17C illustrates the variation in the normal QRS configuration that occurs with rotation. The R wave amplitude should measure from 1 mm to at least 20 mm in V3 and V4 (see Table 2-1). Loss of R waves in V1 through V4 with ST segment elevation indicates acute anterior MI (Fig. 2-18A).

• Loss of R wave in V1 through V3 with the ST segment isoelectric and the T wave inverted may be interpreted as anteroseptal MI age indeterminate (i.e., infarction in the recent or distant past) (Fig. 2-18B). Features of old anterior MI are shown in Fig. 2-18C and lateral infarction is shown in Fig. 2-19.

* = with clockwise rotation, the V1 electrode, like aVR, faces the cavity of the heart and records a QS complex; no initial q in lead V6.

† = qR complexes, q < 0.04 second, < 3 mm deep, therefore not pathologic Q waves.

‡ = loss of R wave V3 to V5 pathologic Q waves: signifies anterior myocardial infarction.

Fig. 2-17. Continued

V1 V2 V3 V4 V5 V6
Normal intermediate position

Extreme clockwise rotation*

Extreme counterclockwise rotation†
qR complexes

Loss of R waves V3–V5 pathologic Q waves‡
Fig. 2-18. A, Loss of R waves in V2 through V5 (i.e., pathologic Q waves associated with abnormal ST segment elevation) indicates acute anterior infarction. B, Loss of R wave in V1 through V3 (i.e., pathologic Q waves associated with an isoelectric ST segment) and T wave inversion indicate anteroseptal infarction, age indeterminate, infarction occurring approximately 1 to 12 months before the recording of this tracing; comparison with previous ECGs and clinical history required to determine the age of infarction. C, Loss of R waves in V2 through V5 (i.e., pathologic Q waves in V2 through V4 not associated with acute ST segment changes) indicates old anterior infarction. D, Loss of R waves in V4 and V5 indicate anterior myocardial infarction, age indeterminate.
Fig. 2-18. Continued
Fig. 2-18. Continued
Fig. 2-19. Pathologic Q waves in V₄ through V₆ and ST segment in keeping with an old anterolateral infarct; clinical correlation necessary to confirm the presence of an old infarction.

Poor R wave progression in V₂ through V₄ may be caused by the following:

- Improper lead placement.
- Late transition (Fig. 2-20).
- Anteroseptal or anteroapical MI.
- LVH (see Chapter 7).
- Severe chronic obstructive pulmonary disease, particularly emphysema—emphysema may cause QS complexes in leads V₁ through V₄,
Fig. 2-20. Poor R wave progression in leads V₂ through V₅. Note: The negative QRS complex in V₅ is caused by late transition and not by other causes of poor R wave progression such as anterior infarction. ECG within normal limits.

which may mimic MI; a repeat ECG with recording electrodes placed one intercostal space below the routine locations should cause R waves to be observed in leads V₂ through V₄ (see Chapter 6)

• Hypertrophic cardiomyopathy.
• LBBB (Fig. 2-8B).

In women, albeit rarely, the R wave in V₂ or V₃ may be less than 1 mm tall; this may cause an erroneous diagnosis of anteroseptal infarction.

In summary: An abnormal, pathologic Q wave is defined in adults as one that has a duration of >40 ms, but the definition does not apply to leads aVR and V₁, which may normally lack the initial R wave. In
addition, in leads III, aVF, and aVL, the initial R wave may be absent; the resultant QS or QR pattern represents a normal variant. A QS pattern is disturbing to students and clinicians. The student is warned: Sound knowledge of normal variants and features of normal ECG deflections that look abnormal but are indeed normal must be mastered by the competent interpreter of ECGs. A QS is often observed in lead aVL in thin subjects with a vertical heart. A QS in lead III is common in individuals with a horizontal heart position, some of whom are obese. See Chapter 6 for an in-depth discussion of Step 5: Q wave abnormalities.

**STEP 6: ASSESS P WAVES** (FIG. 2-21)

- Assess the P waves for abnormalities including atrial hypertrophy (Figs. 2-22 and 2-23; see also Fig. 3-3).

---

**STEP 6**

Assess P waves in leads II and V1 for hypertrophy.

Peaked ≥3 mm amplitude

- Yes
  - Probable right atrial enlargement (see Figure 2-22 and Chapter 7)
  - Check for RVH and causes of RVH, right ventricular strain, and other features of pulmonary embolism (see Chapter 10)

- No

Broad >2.5 mm (≥0.11 second) or bifid in lead II

- Yes
  - Left atrial enlargement.*
  - Check for mitral stenosis
  - mitral regurgitation
  - LV failure
  - dilated cardiomyopathy
  - LVH and causes of LVH (see Figure 2-23 and Chapter 7)

- No

or diphasic in V1 (see Figures 2-22 and 2-23)

---

* = Left atrial abnormality: enlargement, hypertrophy, or increased atrial volume or pressure.

**Fig. 2-21.** Step-by-step method for accurate ECG interpretation. Step 6: assess P waves.
Fig. 2-22. A, Left atrial enlargement: P wave duration greater than three small squares (0.12 second) in lead 2; in lead V1 the negative component of the P wave occupies at least one small box: 1 mm $\times$ 0.04 second = P terminal force $\geq$ −0.04 mm s. B, Right atrial enlargement: lead 2 shows P amplitude $>3$ mm; in V1, the first half of the P wave is positive and $>1$ mm wide (see Figs. 2-23, 7-2, and 7-3). (From Khan, M. Gabriel: On Call Cardiology, 3rd ed., Philadelphia, 2006, WB Saunders, Elsevier Science.)

Fig. 2-23. Lead V1 shows right and left atrial hypertrophy. Lead II shows peaked P waves caused by right atrial enlargement.
STEP 7: ASSESS FOR LEFT AND RIGHT VENTRICULAR HYPERTROPHY (FIG. 2-24)

- Assess for LVH (see Figs. 2-24 and 2-25 and Chapter 7) and RVH (see Fig. 2-26 and Chapter 7 for further details).
- Criteria for LVH and RVH are not applicable if bundle branch block is present. Thus, it is essential to exclude LBBB and RBBB early in the interpretive sequences as delineated previously in Steps 2 and 3.

**STEP 8: ASSESS T WAVES (FIG. 2-27)**

- Assess the pattern of T wave changes (see Fig. 2-27). T wave changes are usually nonspecific (Fig. 2-28). T wave inversion associated with ST segment depression or elevation indicates myocardial ischemia (Fig. 2-29). See Chapter 8 for further information on T wave abnormalities.

---

**STEP 7**

**a. Assess for left ventricular hypertrophy (LVH).**

1. S wave in V1 + R wave in V5 or V6 ≥35 mm = LVH ≈90% specificity; sensitivity <40%
2. R wave in aVL + S wave in men ≥24 mm and in women ≥18 mm = LVH ≈90% specificity; sensitivity <40%
3. Specificity of (1) or (2) increased to ≈98% in presence of
   (a) Left atrial enlargement or
   (b) ST segment depression and T wave inversion (strain pattern) in V5 or V6 (see Figures 2-23 and 2-25)

**b. Assess for right ventricular hypertrophy (RVH).**

1. R wave in V1 ≥7 mm†
2. S wave in V5 or V6 ≥7 mm
3. R/S ratio in V1 ≥1
4. R/S ratio in V5 or V6 ≤1
5. Right axis deviation ≥ +110°
   Any two of above = RVH likely (see Figure 2-26)
6. Specificity increased if ST depression and T wave inversion in V1 to V3 or right atrial hypertrophy (see Figures 2-26, 7-7, and 7-8

---

*Age >30; ≥40 mm, age 20 to 30.
†Age >30; see Table 2-1.

**Fig. 2-24.** Step-by-step method for accurate ECG interpretation. Step 7: Assess for left ventricular hypertrophy (LVH) and right ventricular hypertrophy (RVH) (not applicable if QRS duration ≥0.12 second or in presence of LBBB or RBBB).
Fig. 2-25. Note that the standardization at half voltage in V₁ through V₆ is markedly increased; ST-T strain pattern in V₅ and V₆ and left atrial enlargement are typical features of left ventricular hypertrophy.
Fig. 2-26. A, Leads V₁ through V₆: A tall R wave in V₁, R/S ratio in V₁ >1, and R/S ratio in V₅ or V₆ <1 are features of right ventricular hypertrophy. B, Limb leads: right-axis deviation +140 degrees, peaked P wave in lead II, and right atrial enlargement are all in keeping with right ventricular hypertrophy.
Fig. 2-27. Step-by-step method for accurate ECG interpretation. A, Step 8: Assess T wave changes. B, Step 8: Alternative approach for the assessment of T wave changes.
Fig. 2-28. T wave inversion in V2 through V5 not associated with ST segment depression or elevation; nonspecific ST-T wave changes; cannot exclude ischemia, but the tracing is not diagnostic. Abnormal ECG.
Fig. 2-29. The deep T wave inversion in V2 through V5, which is associated with an abnormal ST segment that is hitched up in V3 and abnormally shaped in V3 and V4, is in keeping with myocardial ischemia and likely left anterior descending artery obstruction. Tracing from a 52-year-old woman with unstable angina; tracing taken in the absence of chest pain.

**STEP 9: ASSESS ELECTRICAL AXIS** (FIG. 2-30)

Assess the electrical axis (see Fig. 2-30 and Table 2-4) using two simple clues:

1. If leads I and aVF are upright, the axis is normal.
2. The axis is perpendicular to the lead with the most equiphasic or smallest QRS deflection (see Fig. 2-30B). Figure 2-31 shows left-axis deviation and the commonly associated left anterior fascicular block (see Chapter 9 and Figs. 9-5, 9-8, 9-10).
**Rapid ECG Interpretation**

**Axis**

**STEP 9**

Rule I

- QRS upright leads I and aVF?
  - Yes
  - QRS positive in lead I and negative in aVF
  - Normal: 0° to +110° age <40, -30° to +90° age >40
  - Left: -30° to -90° (see Figure 2-30, B)
  - Right: +110° to +180°

- No
  - QRS negative in lead I and positive in aVF
  - Axis is perpendicular to this lead and in quadrant determined in rule I above (see Figure 2-30, B)

Rule II

- Locate the smallest or most equiphasic lead

---

**A**

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<tr>
<th>Axis</th>
<th>Lead I</th>
<th>aVF</th>
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<th>III</th>
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**B** *Most equiphasic lead.*
Table 2-4

<table>
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<tr>
<th>Most equiphasic lead</th>
<th>Lead perpendicular*</th>
<th>Axis</th>
</tr>
</thead>
<tbody>
<tr>
<td>III aVR</td>
<td>Normal = +30 degrees</td>
<td></td>
</tr>
<tr>
<td>aVL II</td>
<td>Normal = +60 degrees</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Lead I positive and aVF negative = Left axis</td>
<td></td>
</tr>
<tr>
<td>II aVL (QRS positive)</td>
<td>Left = −30 degrees</td>
<td></td>
</tr>
<tr>
<td>aVR III (QRS negative)</td>
<td>Left = −60 degrees</td>
<td></td>
</tr>
<tr>
<td>I aVF (QRS negative)</td>
<td>Left = −90 degrees</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Lead I negative and aVF positive = right axis</td>
<td></td>
</tr>
<tr>
<td>aVR III (QRS positive)</td>
<td>Right = +120 degrees</td>
<td></td>
</tr>
<tr>
<td>II aVL (QRS negative)</td>
<td>Right = +150 degrees</td>
<td></td>
</tr>
</tbody>
</table>

*Lead perpendicular (at right angles) to the most equiphasic (isoelectric) lead usually has the tallest R or deepest S wave.

Fig. 2-30. A, Step-by-step method for accurate ECG interpretation. Step 9: Assess the electrical axis. Leads are indicated in parentheses. B, See Table 2-4 and Figs. 2-3 and 9-5.
Fig. 2-31. Lead aVR is the most equiphasic: The lead perpendicular to aVR is lead III, indicating a left axis of −60 degrees. There is a small normal Q wave in lead I and a small R wave in lead III in keeping with left anterior fascicular block (hemiblock). Borderline ECG.

STEP 10: ASSESS FOR MISCELLANEOUS CONDITIONS (FIG. 2-32)

Perform a rapid screen for miscellaneous conditions (see Fig. 2-32). Chapter 10 gives details and relevant ECGs.

• Artificial pacemakers: If electronic pacing is confirmed, usually no other diagnosis can be made from the ECG (see Chapter 10).
• Prolonged QT syndrome: See normal QT parameters listed in Table 2-5. No complicated formula is required for assessment of the QT intervals (see Chapter 10 for further details). Some miscellaneous conditions are illustrated in Figures 2-33 to 2-36.
**Fig. 2-32.** Step-by-step method for accurate ECG interpretation. Step 10: Assess for miscellaneous conditions (see Chapter 10).

| QT Intervals* |
|---|---|---|
| **Clinically useful approximation of upper limit of QT interval (s)** |
| **Heart rate (bpm)** | **Male** | **Female** |
| 45–65 | <0.47 | <0.48 |
| 66–100 | <0.41 | <0.43 |
| >100 | <0.36 | <0.37 |

*ECG paper speed 25 mm/s. No complicated formula required.
Fig. 2-33. Stage 1 electrocardiographic changes from patient with acute pericarditis. Diffuse ST segment elevation, which is concave upward, is present in all leads except aVR and V1. Depression of the PR segment, an electrocardiographic abnormality that is common in patients with acute pericarditis, is not evident because of the short PR interval. (From Braunwald E: Heart Disease: A Textbook of Cardiovascular Medicine, 5th ed., Philadelphia, 1997, WB Saunders, Elsevier Science.)

Fig. 2-34. V leads of a 39-year-old woman who had a large atrial septal defect repaired 5 years earlier. Note the RSR′ in lead V1 and a wide, slurred S wave in V5; the QRS duration is 0.1 second: Incomplete right bundle branch block.
Fig. 2-35. ECG showing electronic pacing and ventricular capture; rate is 60 beats/min. No further analysis is possible because of pacemaker rhythm.

Fig. 2-36. Mirror-image dextrocardia with situs inversus. The patient is a 15-year-old girl. There is no evidence of organic heart disease. A, Tracing recorded with conventional electrode placement. B, Tracing obtained with the left and right arm electrodes reversed. The precordial lead electrodes also were located in the respective mirror-image positions on the chest. The tracing is within normal limits. (From Chou TC: Electrocardiography in Clinical Practice, 4th ed., Philadelphia, 1996, WB Saunders, Elsevier Science.) (continued)
STEP 11: ASSESS ARRHYTHMIAS (FIG. 2-37)

Tachyarrhythmias should be analyzed as the following:

- Narrow complex tachycardia: Figure 2-37A gives the differential diagnosis of narrow QRS complex tachycardia.
- Wide complex tachycardia: Figure 2-37B gives the differential diagnosis of wide QRS complex tachycardia. See Chapter 11 for relevant ECGs.

**Fig. 2-36.** Continued

**Fig. 2-37.** Step-by-step method for accurate ECG interpretation. Step 11: Assess arrhythmias: differential diagnosis of narrow QRS tachycardia (A) and wide QRS tachycardia (B).
and detailed discussion of arrhythmia diagnosis including that of bradyarrhythmias.

**ELECTROCARDIOGRAM TECHNIQUE**

- Ensure the standardization is 1 mV displayed as a 10-mm deflection (10 small squares in amplitude).
- Always record the ECG at a standard paper speed of 25 mm/s.
- Remember that artifacts such as baseline drift are often caused by loose or improperly installed sensors.
- Most ECG machines have two modes of operation: automatic or manual. Familiarize yourself with the procedure in the ECG department of your hospital so that you can do the ECG, if called, when there is no technician or nurse available to do the procedure.
- Attach the electrodes (bulb suction cup or flat sensors) on a smooth, fleshy part of the lower arm or forearm and on the fleshy parts of the lower leg.
- Attach the chest lead sensors as indicated in Fig. 2-38 (bulb sensor suction cups or flat sensors).

Ensure that electrodes are properly placed. Incorrect lead placement can lead to serious errors with interpretation (Fig. 2-39).
Fig. 2-38. Chest leads placement. V₁, 4th interspace at the right margin of the sternum; V₂, 4th interspace at the left margin of the sternum; V₃, midway between positions for V₂ and V₄; V₄, 5th interspace at junction of left midclavicular line (apex); V₅, at horizontal level of position V₄ at left anterior axillary line; V₆, same horizontal line as for position V₄ but in the midaxillary line.

Fig. 2-39. A, Atrial fibrillation and pseudo–inferior infarction resulting from electrode misplacement. With Q waves and ST elevation in leads 2, 3, and aVF and with reciprocal depression of the ST segment in aVL and chest leads, this tracing suggests acute inferior infarction. However, lead 1, with virtually no deflections, is the tip-off: The two arm electrodes are on the two legs (and the leg electrodes are on the arms). B, Limb leads with the electrodes attached correctly. (From Marriott JLH: Practical Electrocardiography, 8th ed., Baltimore, 1988, Williams & Wilkins.)
Arm electrodes interchanged. Otherwise ECG within normal limits.

Fig. 2-40.
Untrained technicians often place leads V₅ and V₆ too anteriorly; this may not give a true recording of the left ventricular muscle mass. The leads must be placed in the anterior and midaxillary line (see Fig. 1-15). Incorrect placement of V₂ and V₃ may render a false interpretation of old anteroseptal MI. Thus, much care is needed in placing the chest leads. Feel for the bony points (see Fig. 1-14) to position V₂, V₃, and V₄. Small changes in electrode position can cause significant changes in the record obtained with these leads.

The most common error is the reversal of the left and right arm leads. The ECG records the following: the P wave is negative in lead I and upright in aVR; lead I is a mirror image of I, and therefore the entire complex that is usually positive becomes negative; there is reversal of lead aVR and aVL (aVR is aVL: aVL shows a negative P wave and a relatively negative complex because it is aVR); and there is reversal of leads II and III (lead II is III and lead III is II). Figure 2-40 shows the effect of the reversal of the arm leads. The P, QRS, and T waves are inverted in leads I and aVL; the precordial (V) leads remain normal, however, and thus rule out dextrocardia, in which the limb leads are similar but there is loss of R waves or poor R wave progression from V₂ through V₆ (see Fig. 2-36).
3

P Wave Abnormalities

CONTENTS

FEATURES OF THE NORMAL P WAVE
FEATURES OF ABNORMAL P WAVES
(See Figs. 2-21 and 2-22)

The P wave represents the spread of the electrical impulse through both atria (see Fig. 1-8). The electrical impulse begins in the SA node and depolarizes the right atrium and then the left atrium. Thus, the first part of the P wave reflects right atrial activity, and the late portion of the P wave represents electrical potential generated by the left atrium.

FEATURES OF THE NORMAL P WAVE

The following are some features of the normal P wave:

• It should be upright in leads I and II, as well as in the precordial leads V3 through V6 (Figs. 3-1 and 3-2).
• It is always inverted in aVR.
• It is usually upright in aVF and V3, but occasionally a diphasic or flat P wave may be seen.
• It is variable in leads III, aVL, V1, and V2: upright, inverted, or diphasic. (A P or T wave that is partly above the baseline and partly below it is referred to as diphasic.)

FEATURES OF ABNORMAL P WAVES
(See Figs. 2-21 and 2-22)

• Inverted in II, III, and aVF and upright in aVR: diagnostic of an atrioventricular (AV) junctional (see Fig. 3-2) or ectopic atrial rhythm. When there is abnormal propagation of the electrical impulse through the atria, the polarity or axis of the P wave is abnormal.

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Inverted in lead I and upright in aVR, with lead I being the mirror image of I: caused by reversed arm leads or dextrocardia, but in true dextrocardia there is a loss of R wave in V₄ through V₆ (see Figs. 2-36 and 2-40).

Duration ≥0.12 second (three small squares). Most prominent in leads II, III, and aVF; caused by left atrial enlargement (see Figs. 2-21 and 2-40). P waves are seen best in leads II and V₁; thus, these leads should be used for rhythm strips and arrhythmia detection.

Notching of a wide P wave in lead II, III, or aVF: a distance between peaks >0.04 second usually indicates left atrial enlargement (see Fig. 2-22).
Fig. 3-1. B, Same tracing as in (A) showing normal upright P waves in leads V₃ through V₆.

- Diphasic in V₁: the second half of the P wave is dominantly negative and wide (see Figs. 2-22, 2-23, and 3-3). The depth of the inversion multiplied by the width represents the P terminal force; if it is ≥−0.04 mm (i.e., a negative amplitude of 1 mm with duration of 0.04 second), consider left atrial enlargement (see Figs. 2-22, 2-23, and 3-3). In V₁, the negative deflection is normally <1 mm.
- Large diphasic in V₁: if the first half of the P wave is positive ≥1.5 mm and the second half is negative ≥1 mm and wide, consider biatrial enlargement (see Fig. 2-23).
- High amplitude, peaking (see Figs. 2-22 and 2-23): tall, pointed P waves, taller in lead III than in lead I; high amplitude (≥2.5 mm), particularly in lead II, III, or aVF, indicates right atrial enlargement. Consider the presence of right ventricular hypertrophy, cor pulmonale, pulmonary hypertension, or pulmonary and tricuspid stenosis. Positive amplitude of the first half of the P wave in V₁ or V₂ ≥1.5 mm indicates right atrial enlargement (Figs. 3-4 and 3-5).
Fig. 3-2. P wave is inverted in leads II, III, and aVF and is upright in aVR, indicating junctional rhythm.

Fig. 3-3. The second half of the P wave in V₁ is dominantly negative and wide, indicating left atrial enlargement.
Fig. 3-4. Tall pointed P waves, high amplitude (>2.5 mm), particularly in leads II, III and aVF. Note the P wave is much taller in lead III than in lead I; typical features of right atrial hypertrophy (enlargement).
Absent P waves: consider SA block and AV junctional rhythms. If the rhythm is irregular, consider atrial fibrillation (see Chapter 11).

Different morphologies: at least three different P wave morphologies in the same lead: consider multifocal atrial tachycardia (see Chapter 11).

---

**Fig. 3-5.** Right atrial enlargement in a 33-year-old man with pulmonary fibrosis, chronic cor pulmonale, and right ventricular failure. P pulmonale pattern is present in the limb leads with abnormally tall and peaked P waves also in leads V₁ through V₃. (From Chou TC: *Electrocardiography in Clinical Practice*, 4th ed., Philadelphia, 1996, WB Saunders, Elsevier Science.)
Bundle Branch Block

CONTENTS

RIGHT BUNDLE BRANCH BLOCK (RBBB)
LEFT BUNDLE BRANCH BLOCK (LBBB)

RIGHT BUNDLE BRANCH BLOCK (RBBB)

Diagnostic Criteria

- Wide QRS ≥0.12 second.
- A secondary R wave (R’) in V1 or V2 (i.e., an rSR’, rsR’, or rsr’ complex that often is M-shaped). The secondary R wave (R’) is usually taller than the initial R wave (Figs. 4-1, 4-2A, and 2-7A).
- A wide, slurred S wave in leads V5, V6, and I with duration >40 ms; the S wave is longer in duration (length) than the preceding R wave in leads V6 and I (see Figs. 2-7 and 4-2).
- The axis may be normal, right, or left. If left axis is present, consider left anterior fascicular block (hemiblock) (see Chapter 9).

Genesis of the QRS in RBBB

The typical M-shaped complex in V1 or V2 is derived from an alteration of the normal vector forces (see Figs. 1-17 and 4-1).

- The initial impulse depolarizes the septum normally from left to right. With RBBB, vector I remains intact; the electrical current traveling toward the electrode V1 positioned over the right ventricle registers an initial small R wave in leads V1 and V2 (see Fig. 4-1). Because the right bundle branch does not conduct the electrical impulse, vector II is directed leftward only, activates the left ventricle, and records an S wave in V1 and V2. Right ventricular activation occurs later (i.e., unopposed by left ventricular activation); the resultant force, vector III, causes a large R, termed R’ in V1 or V2. Thus, the rsR’ or rSR’ complex depicts an M shape. The deflection R’ is usually greater than the amplitude of the small R produced by vector I septal depolarization.

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• The unopposed late depolarization of the right ventricle, which causes the R′ in V1 or V2, is recorded as a wide, slurred S wave in leads V5, V6, and I, the electrodes overlying the left ventricle (see Figs. 2-7, 4-1, and 4-2).

• Because of delayed right ventricular activation, the QRS duration is increased to 0.12 second or more. Figure 4-2D shows typical features of RBBB.

• Brugada syndrome is a special form of incomplete or complete RBBB pattern. Although the condition is rare, it is a common cause of idiopathic ventricular fibrillation and sudden cardiac death in young adults, particularly of Asian origin, and notably in individuals without evidence of structural heart disease. Attention must be given to any condition that causes sudden death, particularly in young individuals. The typical ECG features are illustrated in Figs. 4-3 to 4-5. Note the RBBB pattern and persistent ST segment elevation in V1, V2, and V3 that has a typical pattern: coved or saddle-back shaped, a marker for sudden death in individuals without demonstrable structural heart disease. Note that this...
Fig. 4-2. A, An SR in V₁; M-shaped complex in V₁ and V₂; QRS duration ≥0.12 second; and wide, slurred S waves in V₅ and V₆ indicate right bundle branch block (RBBB). B, Same patient as in (A). Lead I, wide, slurred S wave indicates RBBB. C, Right bundle branch block.
Fig. 4-2.  Continued  D. Typical right bundle branch block: QRS, 0.14 second; rsR′ in V1; M-shaped complex in lead V2; and a slurred S wave in V5 and V6. The S wave in lead I and V6 is longer in duration (length) than the preceding R wave.
Fig. 4-3. Typical features of Brugada syndrome. Atypical, incomplete right bundle branch block with a curious (odd shape) ST segment elevation / deformity in V₁ to V₃, described as the coved (in V₁, V₂) and saddle-back patterns (V₃). Note there is no widened S wave in V₅ or in V₆, as seen in true incomplete or complete RBBB. Thus if you recognize an atypical RBBB think of Brugada syndrome and reassess for the characteristic features of this rare but important diagnosis. ECG from a 40 year old man with episodes of syncope / collapse. No recurrence of syncope over 3 years following ICD.
Fig. 4-4. Electrocardiograms from patient during sinus rhythm. ST segment elevation of the coved (lead V₁) and saddle-back types (lead V₂) can be seen. (From Miyazaki T et al.: J Am Coll Cardiol 27:1063, 1996.)

Fig. 4-5. Electrocardiograms from patient during sinus rhythm. Coved-type ST segment elevation can be seen in leads V₁ and V₂. (From Miyazaki T et al.: J Am Coll Cardiol 27:1063, 1996.)
is an atypical incomplete or complete RBBB pattern, because there are usually no widened S waves in V₅ and V₆ of true RBBB: the S wave in V₆ or lead I in RBBB is longer in duration than the preceding R wave. The ST segment elevation appears to be caused by an early high take-off (J wave) and mimics RBBB.

- Arrhythmogenic right ventricular dysplasia, another rare condition that shows an atypical RBBB pattern, is a marker for sudden cardiac death in younger individuals. With this condition, there is structural heart disease caused by a type of cardiomyopathy that involves the right ventricle and the left ventricle at a later stage. Fatty and fibro-fatty degeneration occurs in the right ventricular inflow and outflow tracts and in the apex. The typical ECG finding is shown in Fig. 4-6. Either incomplete or complete RBBB is observed. In approximately 40% of cases, a characteristic terminal notch is observed in the QRS of V₁ and V₂ (termed an epsilon wave) that is a result of slowed intraventricular conduction. Another feature is T wave inversion in V₁, V₂, and V₃. The echocardiogram may show an abnormal right ventricle as the disease progresses.

**Causes of RBBB**

- A normal finding in adults of all ages.
- Coronary artery disease and hypertensive and rheumatic heart disease.
- Congenital heart disease, often associated with ventricular septal defect (VSD) and Fallot tetralogy. With secundum atrial septal defect (ASD), more than 90% of individuals have incomplete RBBB.
- Coarctation of the aorta.
- Pericarditis and myocarditis including Chagas disease.
- Pulmonary embolism and cor pulmonale.
- Cardiomyopathy.
- Brugada syndrome and right ventricular dysplasia (atypical RBBB pattern).

**Incomplete RBBB**

**Diagnostic Criteria**

- The presence of an rSR' (i.e., RBBB pattern) in V₁ or V₂ and an S wave in leads I and V₆ should be confirmed.
- The QRS duration should be 0.08 to 0.11 second.

**Causes of Incomplete RBBB**

- Incomplete RBBB is a common ECG finding in normal individuals.
- More than 90% of patients with a secundum ASD show incomplete RBBB (**see** Figs. 2-34 and 4-3).
Fig. 4-6. A, Normal sinus rhythm in a patient with arrhythmogenic right ventricular dysplasia. The arrowheads point to late right ventricular activation called an epsilon wave. (From Braunwald E: *Heart Disease: A Textbook of Cardiovascular Medicine*, 6th ed., Philadelphia, 2001, WB Saunders, Elsevier Science.)
Fig. 4-6. B. Ventricular tachycardia in the same patient with right ventricular dysplasia. (From Braunwald E: Heart Disease: A Textbook of Cardiovascular Medicine, 6th ed., Philadelphia, 2001, WB Saunders, Elsevier Science.)
• WPW syndrome may mimic incomplete RBBB.
• Brugada syndrome and right ventricular dysplasia.

**RSr' Variant**

More than 5% of individuals without heart disease show an RSr' in $V_1$ or $V_2$. If the QRS duration is $\geq 0.08$ second and there is an S wave in $V_5$ or $V_6$ (see Fig. 4-3), the diagnosis of incomplete RBBB should be made. The diagnosis is strengthened if there is a slurred S wave in I, $V_5$, or $V_6$.

• If a slurred S wave is absent in leads I, $V_5$, or $V_6$ with QRS duration $<0.08$ second, the ECG is interpreted as an rSR', RSR', or RSr' variant, borderline ECG (see Fig. 4-4). An $R' < 6$ mm with an $R'/S$ ratio $<1$ suggests normality.

**Causes of rSR', RSR', and rSR' in $V_1$ or $V_2$: QRS Duration $\leq 0.11$ Second**

• Idiopathic; a normal finding in 5% of individuals without heart disease.
• Incomplete RBBB
• Straight back syndrome or pectus excavatum
• ASD
• Rarely, VSD and coarctation of the aorta
• Mitral stenosis and other acquired heart diseases
• Right ventricular hypertrophy
• Right ventricular volume overload
• Cor pulmonale or pulmonary embolism
• WPW syndrome (may mimic incomplete RBBB)
• Atrioventricular nodal reentrant tachycardia (see Chapter 11).
• Muscular dystrophy
• Late activation of the outflow tract of the right ventricle, the crista supraventricularis (may cause $r'$ wave in $V_1$)
• Incorrect placement of the $V_1$ electrode

The RSr' may appear if $V_1$ is placed in the third interspace and may disappear with the electrode in the fifth interspace, or incomplete RBBB may be recorded (see Figs. 2-34 and 4-8). The appearance at a higher intercostal space may be the only abnormality in some patients with a secundum ASD (see Figs. 2-34, 4-7, 4-9, and 10-1).

**RBBB and Myocardial Infarction**

• With acute anterior MI, pathologic Q waves occur in $V_1$, $V_2$, $V_3$, or $V_4$. A Q wave in $V_1$ and $V_2$ is not sufficient evidence for the diagnosis of MI.
Fig. 4-7. Sinus tachycardia, rate 147 beats/min. QRS duration 0.10 second and rSR′ in V₁ indicate incomplete right bundle branch block.

- Consider inferior MI only if pathologic Q waves are present in leads II, III, and aVF. Q waves in leads III and aVF are not diagnostic.
- The right bundle branch and the septum are supplied blood by the same artery; thus anteroseptal infarction commonly is associated with RBBB. In anteroseptal MI, the initial septal force, vector 1, is lost. Thus, a loss of the initial r wave occurs in V₁ with resultant q or Q wave in V₁, V₂. In addition, the normal small q wave in V₆ disappears (Fig. 4-10).
Fig. 4-8. RSr′ and rSR′ in V1; recorded in different intercostal spaces.

Fig. 4-9. Atrial septal defect. The patient is a 21-year-old woman with atrial septal defect proved by cardiac catheterization. The pulmonary arterial pressure was normal. The ECG shows a frontal plane QRS axis of 90 degrees and rSR′ pattern in lead V1. There also are ST and T changes in the right and middle precordial leads and inferior leads. (From Chou TC: Electrocardiography in Clinical Practice, 4th ed., Philadelphia, 1996, WB Saunders, Elsevier Science.)
Chapter 4 / Bundle Branch Block

**Fig. 4-10.** Mechanism of right bundle branch block. **A**, Note that the right ventricle is activated last and without any opposing forces, resulting in the late R’ in lead V1 and the S wave in lead V6. **B**, In anteroseptal infarction, a QR pattern develops in lead V1. Loss of anterior wall tissue causes an R/S pattern in lead V6. (From Wellens HJ et al.: The ECG in Emergency Decision Making, Philadelphia, 1992, WB Saunders, Elsevier Science.)

**LEFT BUNDLE BRANCH BLOCK (LBBB)

**Diagnostic Criteria**

- QRS duration ≥0.12 second.
- A broad monophasic R wave that is often notched or slurred in lead I, aVL, V5, or V6 (Figs. 4-11 to 4-13 and Fig. 2-8B).
- Late intrinsicoid deflection in leads I, V5, and V6 greater than 0.05 second.
- Leads V1 and V2 reveal QS or rS pattern with poor R wave progression in V2 and V3 (see Figs. 4-11 to 4-13). Figure 4-14A shows notching of lead I. Figure 4-14B shows all the typical features of LBBB.
- A presumptive diagnosis of incomplete LBBB may be made if the QRS duration is 0.10 to 0.11 second with notching of the R wave in V5 or V6.

**Genesis of the QRS Complex in LBBB**

- Depolarization of the left ventricle is delayed, and the QRS duration is prolonged to ≥0.12 second.
- The septum and left ventricle are activated by the electrical impulse from the right bundle.
- The normal direction of septal activation from left to right is reversed.
Vector I flows from right to left through the lower septum rather than from left to right. Thus, an electrode over the left ventricle records an R wave in V5, V6, and I and a QS or rS in V1 (see Figs. 4-11, 4-12, and Fig. 2-8B).

Vector II travels from left to right through the right ventricular mass and may cause a slur or notch in the R wave of leads I, aVL, V5, and V6 [marked V(II) in Fig. 4-11]. The notched R and R’ may result in an M-shaped complex in lead I, V4, V5, or V6 (see Figs. 4-11 to 4-14).

Vector III travels right to left and causes an R’ in V6 [marked V(III) in Fig. 4-11].

The marked derangement in depolarization of the left ventricle causes the ST segment in leads V1 through V4 to be abnormally elevated (see Fig. 4-13).

The direction of the ST segment and T waves is opposite the direction of the terminal QRS (see Figs. 4-11 to 4-13).

Because LBBB deranges normal vector forces, the diagnosis of left ventricular hypertrophy (LVH) cannot be made in the presence of LBBB. ST elevation, poor R wave progression in V1 through V3, and increased
Fig. 4-12. Sinus bradycardia, 40 beats/min; QRS duration >0.12 second. Note the broad, monophasic R wave, notched in V5 and V6, and poor R wave progression in V2 and V3, typical features of left bundle branch block.
Fig. 4-13. QRS duration ≥0.12 second; poor R wave progression; and notched R wave in $V_6$. Note ST segment elevation in $V_1$ through $V_5$, typical of left bundle branch block that mimics anterior myocardial infarction.
Fig. 4-14. A. Notching in lead I indicates left bundle branch block (two patients). (continued)
Fig. 4-14. Continued

B. All of the typical electrocardiographic features of LBBB. Note the absence of normal q waves in $V_5$, and $V_6$. 
voltage are common features of LBBB and do not indicate LVH, myocardial injury, or MI (see Figs. 4-12, 4-13, and Fig. 2-8B and Chapter 6).

**Causes of LBBB**

- Cardiomyopathies and degenerative diseases.
- Coronary artery disease (CAD); patients with CAD and LBBB have a high incidence of left ventricular dysfunction and congestive heart failure.

QRS $\geq 0.11$ second but not typical RBBB or LBBB configuration

1. Atypical RBBB or LBBB
   (Figure 11-27)
   - **Spot for** Delta wave + PR $\leq 0.12$
   - Present
   - **WPW Syndrome**

2. Atypical RBBB but WPW Excluded and no slurred S wave in V$_5$ and V$_6$
   (therefore not true RBBB)
   - **Spot for** ST elevation in V$_1$ and V$_2$
   - (Coved or Saddle-back)
   - Present
   - **Brugada Syndrome** (Figures 4-3, 4-4, 4-5)

3. Atypical RBBB 1 and 2 Excluded
   - **Spot for** A terminal notch in the QRS (Epsilon Wave)
   - + T $\downarrow$ V$_1$ V$_3$
   - Present
   - **Right Ventricular Dysplasia** (Figure 4-6)

4. 1 to 3 Excluded
   - **Diagnosis**
   - **IVCD**

*= In $\approx 20\%$, the QRS is $<0.11$ second
†= QRS duration may be 0.10 to 0.12 second

**Fig. 4-15.** Before making the diagnosis of nonspecific IVCD, consider steps 1, 2, and 3.
• Hypertensive heart disease.
• Advanced valvular heart disease.
• Congenital heart disease.
• Idiopathic in patients with structurally normal hearts. Although LBBB usually occurs in patients with underlying heart disease, the condition may occasionally occur in individuals with structurally normal hearts. These individuals fall primarily into a category of young, healthy adults with idiopathic LBBB or older subjects with primary disease of the conducting system. New LBBB that develops at age 45 or later is likely caused by a significant disease process. In the Framingham study, 55 individuals developed new LBBB at an average age of 62, and coronary heart disease was evident in 89% of these; 50% died within 10 years of onset of LBBB.

**Nonspecific Intraventricular Conduction Delay**

**Diagnostic Criteria**

- QRS duration >0.11 with QRS morphology that does not satisfy the criteria for either RBBB or LBBB.
- Figure 4-15 illustrates steps to consider before making the diagnosis of nonspecific IVCD. Notching of a QRS complex with a QRS duration <0.11 should not be classified as nonspecific IVCD.

**Causes of IVCD**

- Coronary heart disease
- Hypertensive heart disease
- Severe valvular heart disease
- Congenital heart disease
- Cardiomyopathies
- Heart failure (all causes)
- Antiarrhythmic agents
- Tricyclic antidepressants
The ST segment begins after the final deflection of the QRS complex and ends at the ascending limb of the T wave (see Fig. 2-1).

**WHY EMPHASIZE THE ST SEGMENT?**

Because important cardiac ECG diagnoses are made from observation of abnormalities of the ST segment, the interpreter should rapidly focus on the ST segment. This assessment is Step 4 in the method for accurate ECG interpretation (Fig. 5-1). This step is carried out before the assessment for loss of R waves or for the presence of pathologic Q waves, T wave abnormalities, hypertrophy, and axis determination. The diagnosis of acute MI, ischemia, and pericarditis depends on careful scrutiny of the ST segment.

**STEP 4**

Assess the ST segment for the following:
- Elevation
- Depression
- Nonspecific changes

The PR segment is usually used to assess the degree of ST segment elevation or depression. The commencement of the ST segment is
**Fig. 5-1.** Step-by-step method for accurate ECG interpretation. Step 4: Assess for ST segment elevation or depression. Exclude other causes of ST elevation:

- Normal variant: 1- to 2-mm ST elevation, mainly in leads V₂ through V₄, nonconvex, and with fishhook appearance. Common in African Americans: even 4-mm ST elevation (see Fig. 2-15).
- Coronary artery spasm: ST returns to normal with nitroglycerin or pain relief.
- Left bundle branch block: QRS $\geq$0.12 second and typical configuration (see Fig. 2-8B and Chapter 4).
- LV aneurysm: known old infarction with old Q waves (see Chapter 6).

usually located at the same horizontal level as the T-P (the isoelectric interval, see Fig. 2-1).

Abnormal ST elevation can be caused by the following:

- Acute MI
- Coronary artery spasm
• Acute pericarditis
• Left ventricular (LV) aneurysm
• Left bundle branch block (LBBB)
• Left ventricular hypertrophy (LVH)

ST ELEVATION MYOCARDIAL INFARCTION (STEMI)

The early diagnosis of acute MI is paramount to successful percutaneous coronary intervention (PCI) or for the timely administration of thrombolytic therapy. This early diagnosis depends on the observation of abnormalities of the ST segment and not on the presence of Q waves or on the results of cardiac enzymes. Reliance on the presence of pathologic Q waves stems from the proven electrocardiographic principles that were used appropriately from 1930 to the late 1980s.

• ST segment depression = ischemia
• ST segment elevation = injury current
• Q waves = necrosis = infarction

With the advent of thrombolytic therapy and PCI, it became necessary to diagnose acute MI within 1 hour of onset of symptoms, and the diagnosis has to be made without reliance on the presence of abnormal Q waves. Most patients with chest pain and abnormal ST segment elevation in two or more contiguous leads develop Q waves from 4 to 24 hours after the onset of symptoms.

Currently, two descriptive terms are used and recognized internationally:

• ST elevation MI (STEMI)
• Non–ST segment elevation MI (probable non–Q wave infarction)

The acute-injury current of infarction elevates the ST segment and deforms its shape. ST segment elevation patterns of infarction and a normal variant are illustrated in Fig. 5-2.

Diagnostic Criteria

Diagnostic criteria for ST elevation MI (probable Q wave infarction) are as follows:

• Abnormal ST elevation of $\geq 1$ mm in two or more contiguous limb leads.
• Elevation in leads II, III, and aVF indicates inferior infarction (Figs. 5-3, 5-4, and Fig. 2-13A). ST elevation in leads I, aVL, V5, and V6 indicates anterolateral infarction (see Fig. 2-18A).
Fig. 5-2. A, ST segment elevation, pattern of normal variant. Note fishhook appearance; the ST segment usually retains the normal concave shape; the T waves are often prominent and peaked. B, Abnormal ST elevation caused by acute myocardial infarction. (From Khan, M. Gabriel: *On Call Cardiology*, 3rd ed., Philadelphia, 2006, WB Saunders, Elsevier Science.)

Fig. 5-3. ST segment elevation in leads II, III, and aVF is diagnostic of acute inferior myocardial infarction. Note reciprocal depression in leads I and aVL, which strengthens the diagnosis.
Abnormal ST elevation of \( \geq 1 \) mm in two or more contiguous precordial leads indicates anterior infarction (see Fig. 5-1). ST elevation in V\(_1\) through V\(_3\) indicates anteroseptal infarction or anteroinferior MI. Recent studies indicated that the area of necrosis is more likely to be anteroinferior rather than anteroseptal (Fig. 5-5). Elevation in V\(_3\) through V\(_6\) (may involve V\(_2\) and V\(_1\)) indicates anterior infarction (Fig. 5-6 and Fig. 2-13B).

Extensive anterior infarction is denoted by ST elevation in eight or more leads (Fig. 5-7).

- ST elevation in V\(_3R\) and V\(_4R\) associated with inferior infarction indicates added right ventricular infarction (Fig. 5-8). It is advisable to record V\(_4R\) if the patient with an acute inferior infarct has hemodynamic deterioration.
- Tall R waves in V\(_1\) and V\(_2\) associated with ST elevation in II, III, aVF, or V\(_4R\) may indicate added posterior infarction. Posterior infarction
Fig. 5-5. Abnormal ST segment elevation in V₁ through V₃: consider acute antero-septal or anteroapical myocardial infarction (*see* text for discussion of anteroapical MI). This patient’s ECG showed reciprocal depression in leads II, III, and aVF, which strengthens the diagnosis of acute infarction.
Fig. 5-6. A, ST segment elevation in V₁ through V₅ and poor R wave progression in V₂ through V₄ typical of recent anterior infarction.

(continued)
**Fig. 5-6.** Continued B, Variation in shapes of ST elevation.
Fig. 5-7. (A) Marked ST segment elevation in eight leads, V₁ through V₆ and leads I and aVL (shown in 7B), indicate extensive anterior MI. The tracing also showed reciprocal depression in the inferior leads. (From Khan, M. Gabriel: *Heart Disease Diagnosis and Therapy*, Baltimore, 1996, Williams & Wilkins.)
Fig. 5-7. Continued  B. See legend on page 125. (From Khan, M. Gabriel: *Heart Disease Diagnosis and Therapy*, Baltimore, 1996, Williams & Wilkins.)

Fig. 5-8. A. Abnormal ST segment elevation in leads II, III, and aVF indicates recent inferior myocardial infarction. B. Same patient as in (A). Leads V₄ through V₆ as labeled were appropriately placed on the right side of the chest: leads V₄R and V₅R show abnormal ST segment elevation, which indicates acute inferior and right ventricular infarction. This tracing was read incorrectly by the computer and cardiologist as “widespread ST elevation, consider pericarditis; changes in V₄ through V₆ indicate lateral infarction.” (Note: V₄ through V₆ were right-sided chest leads and should have been labeled V₄R, 4V₅R, and V₆R. ST elevation in leads V₃R and V₄R is the main electrocardiographic feature of right ventricular infarction that may occur in association with inferior MI; see Fig. 6-18.)
Rapid ECG Interpretation

occurs virtually always in association with inferior or right ventricular infarction. Tall R waves in $V_1$ and $V_2$ and T wave upright with no other ECG evidence of MI requires cardiac enzyme confirmation.

Other signs that strongly support the diagnosis of acute MI include the following:

- The simultaneous presence of reciprocal depression is not diagnostic for MI but helps confirm the diagnosis. This is of particular diagnostic importance because ST elevation that may occur as a normal variant is not associated with reciprocal ST depression. With acute pericarditis, ST depression occurs only in lead aVR and sometimes in $V_1$ (see Fig. 2-33, Chapter 10, and further discussion in this chapter).
- Evolving Q waves. Q waves may become fully developed in 2 to 24 hours from onset of symptoms (Figs. 5-9 and 5-10). In many patients with acute ST elevation MI, Q waves may not develop, particularly

Fig. 5-9. ST segment elevation in $V_1$ through $V_4$ indicates acute anteroseptal infarction, anteroapical MI, or anterior MI.
when thrombolytic agents are used. After 12 hours, Q waves may be smaller or appear in fewer leads, and the reduced R wave amplitude (loss of R) is less pronounced in patients with reperfusion than in those without. Nonetheless, Q wave regression in patients with acute anterior MI who have been administered thrombolytic therapy does not correlate with improvement in left ventricular function and appears to have no prognostic significance.

- Diminution of R waves in V2 through V4 (i.e., poor R wave progression), especially if an R wave is present in V1 or V2 and disappears or becomes smaller in V3 or V4.

Fig. 5-10. Same patient as in Fig. 5-9. ECG taken 10 hours later shows evolutionary changes: Q waves are present in V1 through V4, which indicates anterior infarction; convex ST segment elevation is decreased; and T wave inversion has emerged. (From Khan, M. Gabriel: Heart Disease Diagnosis and Therapy, Baltimore, 1996, Williams & Wilkins.)
• Evolutionary ST-T wave changes that occur during the 10 to 30 hours after the onset of infarction (see Fig. 5-10).
• A decrease in ST segment elevation of 2 mm (0.2 mV) or more may be observed within 30 minutes after the beginning of thrombolytic treatment and may continue for 6 hours.

**Infarct Size**

An approximation of the size of the infarction can be gauged from the extent of ST elevation:

• Small MI: ST elevation in two or three leads
• Moderate MI: four or five leads
• Large MI: six or seven leads
• Extensive MI: eight or nine leads (see Fig. 5-7)

**Value of Lead aVR in Diagnosis of Acute Myocardial Infarction**

• aVR is a lead that is often ignored but recently has gained importance in the diagnosis of left main coronary artery (LMCA) occlusion.
• Figures 5-11 and 5-12 show ST elevation in aVR that is greater than the elevation in V₁, a marker of LMCA obstruction. This criterion is not specific: specificity is 80% and sensitivity is 81%. Circumflex branch occlusion also may cause ST elevation in aVR, but with no elevation in V₁. In addition, right ventricular overload may reveal ST elevation in aVR, but the clinical scenario is easily differentiated. Subendocardial infarction with marked ST segment depression in V₄ through V₆ that is not caused by left main coronary occlusion may reveal ST segment elevation in aVR, but the elevation may be less than that observed in V₁.
• Because LMCA occlusion is a highly serious condition, any noninvasive diagnostic clue represents a valuable addition to the diagnostic armamentarium.

Reciprocal ST segment depression in aVR with PR segment elevation in aVR with reciprocal PR segment depression in other leads is a feature of acute pericarditis (see Chapter 10).

**Mimics of ST Elevation Infarction**

• Normal variants: ST segment elevation is often observed as a normal variant in healthy African Americans, Hispanics, and some other ethnic
Fig. 5-11. Representative 12-lead ECG tracings at admission in a patient in the left main coronary artery (LMCA) group (A), the left anterior descending coronary artery (LAD) group (B), and the right coronary artery (RCA) group (C). In the patient in the LMCA group, ST segment elevation is apparent in lead aVR. In the patient in the LAD group, significant ST segment elevation in the precordial leads is seen, whereas ST segment shift in lead aVR is negligible. In the patient in the RCA group, ST segment elevation in the inferior leads is marked. (From Yamaji H et al.: J Am Coll Cardiol 38:1351, 2001.)
Patient with chest pain for 3 hours. Inferior myocardial infarction and ST elevation in aVR and V₁. Left main occlusion.
The ST elevation commonly seen in V2 through V5 often shows a notched J-point, fishhook appearance. This normal variant is inappropriately termed early repolarization changes. ST elevation may occur in leads II, III, and aVF, but reciprocal depression does not occur. The degree of ST elevation is variable, often 1 to 4 mm; the normal concave shape remains, but it may end in a prominent, peaked T wave (see Figs. 5-2, 5-13, and Fig. 2-15). Occasionally, ST elevation with T wave inversion is observed in one or two precordial leads in healthy athletes (Fig. 5-14).
Fig. 5-14. Benign ST and T wave changes in a healthy 24-year-old professional athlete. The changes, especially in V_4 and V_5, mimic myocardial injury and ischemia and remained the same 15 months later. (From Chou TC: Electrocardiography in Clinical Practice, 4th ed., Philadelphia, 1996, WB Saunders, Elsevier Science.)

- Acute pericarditis causes diffuse ST segment elevation that is not confined to an anatomic coronary blood supply; thus ST elevation is observed in leads I through III, lead aVF, and most precordial leads. The ST segment retains a normal concave shape. Reciprocal depression may be observed in aVR and sometimes in V_1 (see Fig. 2-33 and discussion under “Pericarditis” in Chapter 10).
- MI age indeterminate (see Fig. 2-18B) in the absence of LV aneurysm may exhibit mild ST elevation, and the differentiation from acute infarction requires clinical correlation and comparison with previous ECGs.
- Coronary artery spasm, Prinzmetal angina, causes ST elevation during the brief period of chest pain (Fig. 5-15).
- LV aneurysm: ST elevation can persist 3 days to 4 weeks after acute infarction; persistence beyond 4 weeks suggests LV aneurysm (Figs. 5-16 and 5-17).
Fig. 5-15. Variant angina. The patient had severe coronary artery disease involving all three major vessels, especially the anterior descending branch, as demonstrated by coronary arteriogram. A, Tracing recorded during angina at rest. B, Tracing recorded 20 minutes later, after the pain had subsided. The latter tracing is representative of the patient’s baseline ECG. During angina (A), marked ST segment elevation is present in leads II, III, aVF, and V₃ through V₆, with reciprocal ST segment depression in leads aVR and aVL. There is also an increase of the amplitude of the R wave in leads II, III, and aVF, with the disappearance of the S waves in leads showing significant ST segment elevation. The resulting complexes resemble the monophasic transmembrane potential. Many similar episodes were observed in this patient. (From Chou TC: *Electrocardiography in Clinical Practice*, 4th ed., Philadelphia, 1996, WB Saunders, Elsevier Science.)
Fig. 5-16. Ventricular aneurysm. The patient is a 52-year-old man who had an acute extensive anterior myocardial infarction 5 months before the recording of this ECG. Note the persistent ST segment elevation in the precordial leads and in leads I and aVL. Left anterior hemiblock also is present. (From Chou TC: Electrocardiography in Clinical Practice, 4th ed., Philadelphia, 1996, WB Saunders, Elsevier Science.)

- LBBB nearly always causes abnormal ST elevation in leads V_1 through V_6 and can mimic acute or old infarction (see Fig. 4-13).
- LVH may cause poor R wave progression in V_1 through V_3, and occasionally ST elevation is observed (see Figs. 2-23 and 2-25).
- Hypertrophic cardiomyopathy causes Q waves, but, occasionally, persistent ST segment elevation is present (see Chapter 6).
- Acute myocarditis in persons with acquired immunodeficiency syndrome may cause nonspecific ST-T changes; ST elevation and Q waves may occur (see Chapter 6).
- Cocaine abuse may cause ST elevation and, in some individuals, frank infarction (see Chapter 6).
- Pulmonary embolism may cause ST elevation, albeit rarely (see Chapter 10 and Fig. 10-20).
Fig. 5-17. V leads of a patient who sustained an anterior infarction 6 months earlier. Pathologic Q waves are present from V₁ through V₆, and the ST segment is elevated in V₁ through V₅. The tracing is in keeping with an old anterior myocardial infarction with left ventricular aneurysm.

NON–ST SEGMENT ELEVATION MI

- ST segment depression $\geq 1$ mm in two or more leads in a patient with chest discomfort and an abnormal troponin or CK-MB is diagnostic of non–ST segment elevation MI (non–Q wave MI) (see Fig. 5-18 and Fig. 2-14A).
Fig. 5-18. A, Non–Q wave infarction (acute subendocardial infarction) in a patient with a clinical picture of infarction and elevated CK-MB. Note widespread ST-T depression in the limb and chest leads but no associated Q waves. B, The same patient’s ECG tracing 18 hours earlier than depicted in (A). (From Khan, M. Gabriel: On Call Cardiology, 3rd ed., Philadelphia, 2006, WB Saunders, Elsevier Science.)
• Transient ST segment changes >0.05 mm (mV), associated with angina at rest and positive troponin or CK-MB indicate non–ST segment elevation MI. In patients with negative cardiac enzymes within 6 hours of onset of pain, another sample should be drawn between 6 and 12 hours. Patients with acute coronary syndrome, particularly those with rest pain ≥20 minutes accompanied by ECG changes are further risk-stratified depending on troponin levels as follows:

1. High risk: troponins (TnT or TnI) >0.1 ng/mL (elevated troponin levels indicate myocardial necrosis, MI).
2. Intermediate risk: troponin slightly elevated >0.01 but <0.1 ng/mL.
3. Low risk: troponin normal.

**ISCHEMIA**

ST segment depression indicative of definite myocardial ischemia should fulfill the following criteria:

• Greater than 1 mm depression.
• Present in two or more leads.
• Present in two or more consecutive QRS complexes.
• Flat (horizontal) or down-sloping with or without T wave inversion (these patterns of ischemia are all shown in Figs. 5-19, 5-20, and Fig. 2-14B).
• Abnormal convex coving of the ST segment in V₁ through V₃ or V₂ through V₄ associated with T wave inversion.

The terminal portion of the abnormal ST segment may show a typical hitched-up pattern (Fig. 5-21); this pattern is often caused by a tight obstruction in the proximal left anterior descending artery.

**NONSPECIFIC ST CHANGE**

Minor ST segment depression ≤1 mm is not an uncommon finding in normal individuals. Consider ST segment changes to be nonspecific if the following prevails:

• ST depression ≤1 mm in the absence of typical symptoms of unstable angina, including rest pain ≥20 minutes (Fig. 5-22)
• Accompanied by baseline drift
• With or without T wave inversion
• Commonly associated with low, flat, or slightly inverted T waves

T waves normally should be ≥0.5 mm in height in leads I and II (see “T Waves” in Chapter 8). Figure 5-23 depicts nonspecific ST-T changes.
Fig. 5-19. Flat (horizontal) and down-sloping ST segment depression greater than 1 mm in a patient with proven angina and obstructive coronary artery disease. A, Limb leads.
Causes of Nonspecific ST-T Wave Changes

Nonspecific ST-T wave changes can be caused by a number of conditions, such as the following:

- Improper electrode contact
- Ischemia (must be considered; the ECG must be interpreted in regard to the clinical findings)
- Electrolyte abnormalities
- Arrhythmias
- Myocarditis
- Pericarditis, constrictive pericarditis
- Intraventricular conduction defects
- Cardiomyopathy
- Pulmonary embolism

(text continues on pg. 135)
Fig. 5-20. V leads of a patient with severe angina and left ventricular hypertrophy. Note increased voltage and marked ST segment depression in V₄ through V₆.
Fig. 5-21. V leads in a patient with unstable angina. ST-T segment abnormalities seen in V₁ through V₄. The tracing was taken when the patient was pain free. Note the “hitched up” ST segment in V₂ and V₃ with deep T inversion: The pattern is typical of significant proximal left anterior descending coronary artery stenosis.
Fig. 5-22. V leads in a patient with no history of heart disease. ST segment is flat in V₄ through V₆ with minimal T wave inversion; similar findings were observed in leads I and aVL: The anterolateral ST-T wave abnormalities are nonspecific; note that ischemia cannot be excluded. Abnormal ECG.
Fig. 5-23. The ST segment is borderline flat but not depressed and is associated with minimal T wave inversion in leads V₃ through V₆; similar findings were present in leads I and aVL: nonspecific ST-T wave changes. Borderline ECG.

- Drink of cold water
- Hyperventilation
- Drug use, including ethanol abuse
- Digoxin
- Subarachnoid hemorrhage or cerebral hemorrhage (see Fig. 8-14)
Q Wave Abnormalities

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CRITERIA FOR NORMAL AND ABNORMAL Q WAVES

The QRS complex should be assessed for the presence of normal and abnormal Q waves and for normal or abnormal R wave progression as outlined in Step 5 of the method for accurate ECG interpretation (Fig. 6-1).

The assessment of pathologic Q or normal q waves should take into account the following:

• Their width
• Their depth
• The leads in which they are observed
• The age of the individual
• Relevant clinical findings

Normal Parameters

• In general, a Q wave that is wider than 0.03 second is considered abnormal, except in leads III, aVR, and V_1, in which Q waves may be wide and deep in normal individuals (see Fig. 6-2A and Table 2-1).
• Lead aVR normally records a negative QRS, QS, or QR complex (see Fig. 6-2 and Fig. 2-2).
• Normal QS complexes occasionally are found in leads III and V_1 and rarely in V_2.

From: Contemporary Cardiology: Rapid ECG Interpretation, 3e
by: M. Gabriel Khan © Humana Press Inc., Totowa, NJ
a. Assess for Q waves, leads I, II, III, aVF, and aVL.

Normal if <0.04 second (one millimeter square = 0.04 second) and ≤3 mm deep, except lead III normal up to 0.04 second and up to 7 mm deep in III and aVL; lead I ≤1.5 mm deep (see Table 2-1)

If abnormal Q II, III, aVF consider inferior MI

I, aVL, V₅, V₆ determine age of infarct

anterolateral MI (see Figure 6-15) recent (see Figures 2-17, A, and 6-12) old (see Figure 2-17, B) exclude mimics

hypertrophic cardiomyopathy (see Figure 6-23, A and B) WPW syndrome bordering Qs, II, III, aVF (see Figure 6-13)

b. Assess for R wave progression in V₁ through V₆ or pathologic Q waves.

R should be

0 to 6 mm in V₁*

>0.2 mm in V₂ (normal 0.3 to 12 mm)

≥1 mm in V₃ (normal 1 to 24 mm)

If poor R progression, consider

late transition (see Figure 6-7) normal variant (see Figure 6-5)

anterolateral MI LVH (see Figures 2-25 and 6-27 and Chapter 7) LBBB (QRS ≥0.12) (see Figures 2-8, B, and 6-6 and Chapter 4) emphysema (see Figures 6-8 and 6-28) Exclude mimics of MI (see Chapter 5)

V₁ through V₄, consider V₅ and V₆, consider

lateral MI (see Figures 2-19 and 6-15) hypertrophic cardiomyopathy (see Figure 6-23, B)

recent (see Figures 2-13, B, 2-18, A, and 6-10)

indeterminate† (see Figures 2-18, B, and 6-15)

old (see Figures 2-18, C, and 6-9)

*Age >30; see text and Table 2-1 for exceptions and normal parameters.
†Compare old ECGs.

Fig. 6-1. Step-by-step method for accurate ECG interpretation. Step 5: Assess for Q waves and R wave progression.
• A narrow Q wave may occur as a normal finding in lead III; this should be ≤0.04 second in duration (1 square), <10 mm deep, and not accompanied by abnormal Q waves in leads II and aVF (see Fig. 6-2 and Fig. 2-2D). The depth of the Q wave is not as important as the width.

• Lead aVL may record a Q wave <0.04 second and up to 7 mm deep in individuals older than age 30 years and up to 10 mm deep in children.

Fig. 6-2. A, Isolated, deep, narrow Q wave in lead III is ≤0.04 second as part of a normal ECG. Note the absence of abnormal Q waves in leads II and aVF.

B, Note small, normal q waves <0.04 second and <2 mm deep in duration in leads II and aVF. Normal ECG. (continued)
A negative P wave followed by a QS or QR deflection with a negative T wave may be recorded in a normal vertical heart.

- In leads II and aVF, small, narrow Q waves may occur but should be ≤0.03 second in duration and <4 mm deep (see Fig. 6-2). Occasionally, the Q in leads II, III, and aVF are borderline width and the ECG is interpreted as follows: inferior Qs noted; clinical correlation required; borderline ECG.
- In lead I, the depth of a Q wave should not exceed 1.5 mm in adults older than age 30 years (see Fig. 6-2A).
• A small q wave in $V_6 \leq 0.03$ second is present in more than 75% of normal individuals. In leads $V_5$ and $V_6$ and rarely in $V_4$, normal q waves $\leq 0.03$ second and $< 3$ mm deep may occur (Fig. 6-3). Normal Q waves should be $< 3$ mm in adults older than age 40 years; they should not exceed a depth of 4 mm in those younger than age 30 years. Rarely, an amplitude $> 4$ mm may be seen in healthy teenagers.

• In contrast, Figure 6-23B shows abnormal Q waves in $V_4$ through $V_6$; they are 0.04 second wide and 3 mm deep in a 52-year-old woman with

![ECG Diagram]

Fig. 6-3. Normal q wave in $V_4$ through $V_6 \leq 0.03$ second, $< 4$ mm deep.
proven hypertrophic cardiomyopathy. Small Q waves may occur in V2 through V6 with extreme counterclockwise rotation (Fig. 6-4).

- A Q wave of less than 0.03 second and greater than 2 mm deep in V2 through V4 is abnormal if V1 shows an initial R and there is no significant shift of the transitional zone to the left or right.
- Poor R wave progression may simulate infarction patterns (pseudoinfarction).
- Poor R wave progression in V2 through V4 with minute R waves may mimic QS complexes and lead to incorrect diagnoses and uncertainties for cardiologists, the attending physician trainee, or the family physician. Causes of poor R wave progression include the following:
  - In women, and rarely in men younger than age 30 years, minute R waves may be present in leads V1, V2, and sometimes, V3; this poor R wave progression is not uncommon and may lead to an erroneous diagnosis of anteroseptal infarction (Fig. 6-5).

![Diagram of ECG configurations](Fig. 6-4. Variations in normal QRS configuration and correlation with abnormals. (From Khan, M. Gabriel: *On Call Cardiology*, 3rd ed., Philadelphia, 2006, WB Saunders, Elsevier Science.))
Fig. 6-5. ECG from a healthy 74 year old female. Poor wave progression $V_2-V_3$ is a not uncommon finding in females and may mimic old anteroseptal MI. Caution is needed in positioning leads $V_1$ and $V_2$ in both females and males.
- Improper lead placement of $V_2$ and $V_3$, particularly in women, may cause decreased R wave amplitude, often incorrectly assessed as anteroseptal infarction.
- Old anteroseptal and anterior infarction.
- Left bundle branch block (LBBB) (Fig. 6-6).
- Left ventricular hypertrophy (LVH) (see Fig. 7-6).
- Severe chronic obstructive pulmonary disease (COPD), particularly emphysema (Fig. 6-8).
- Late transition (Fig. 6-7).
- Left anterior fascicular block.

Fig. 6-6. Poor R wave progression in $V_1$ through $V_4$ and QRS duration $>$0.12 second indicate left bundle branch block.
Fig. 6-7. Poor R wave progression in V2 through V4 with normal QRS duration. Note that the transition, instead of occurring normally in lead V3, occurs in V5, as indicated by a negative QRS in V5. Tracing from a normal 48-year-old woman. Normal ECG.

A net negative QRS complex in V5 or V6 in the absence of right ventricular hypertrophy (RVH) indicates late transition (Fig. 6-7). Severe COPD is considered if the P wave amplitude is greater than 2.5 mm in any of lead II, III, or aVF (Fig. 6-8). Severe COPD, particularly that caused by emphysema, may reveal poor R wave progression in V1 through V4, or Q waves may indicate pseudoinfarction (see Fig. 6-8). Note the P waves of right atrial hypertrophy, characteristic of P pulmonale, which strengthens the diagnosis of COPD pseudoinfarction.
Chronic obstructive pulmonary disease with pulmonary hypertension. The patient is a 58-year-old man with a pulmonary arterial pressure of 42/25 mm Hg at rest. The ECG shows P pulmonale with a vertical P axis. The QRS complexes in lead I are small, and the frontal plane QRS axis is +90 degrees. There is poor progression of the R wave in the precordial leads, with an R/S ratio in leads V5 and V6 of less than 1. The amplitude of the QRS complexes in V6 also is small. (From Chou TC: *Electrocardiography in Clinical Practice*, 4th ed., Philadelphia, 1996, WB Saunders, Elsevier Science.)

- Note that poor R wave progression in V2 through V4 in the absence of late transition, LVH, or COPD suggests a diagnosis of anterior infarction (Fig. 6-9), but consideration of lead placement error and clinical correlation is required always.

**Q WAVE MYOCARDIAL INFARCTION**

Abnormal Q waves caused by myocardial necrosis occur as early as 2 hours and as late as 24 hours after the onset of clinical symptoms of acute MI. Q waves of acute infarction are always associated with abnormal ST elevation.

**Diagnostic Criteria**

- The presence of ST segment elevation ≥1 mm with or without Q waves in two or more contiguous leads in a patient with acute chest discomfort is diagnostic of ST elevation MI (STEMI), probable Q wave MI (Fig. 6-10A and B, and Fig. 2-13B).
Fig. 6-9. Poor R wave progression in V2 through V4 with abnormality of the ST segment: consider old anteroseptal myocardial infarction. Note the transition zone is not late, in that the QRS complex is not negative in V5 or V6. Comparison with old ECGs and clinical correlation is required. Abnormal ECG.

- From 6 to 12 hours after onset of symptoms, ST segment elevation recedes, but Q waves become more prominent (Fig. 6-11).
- Pathologic Q waves and ST elevation in leads II, III, and aVF indicate inferior infarction (Fig. 6-12).
- The Q waves in leads II, III, and aVF are >0.03 second in duration; the Q wave in lead III is >0.04 second wide. Figure 6-13 shows acute inferior MI with ST-T wave abnormalities and evolutionary changes.
- An abnormal Q wave in lead III (≤0.04 second in duration) not associated with pathologic Q waves in lead II or aVF should be considered a normal variant.
Fig. 6-10. A, Q waves in leads V₁ and V₂ with marked abnormal ST segment elevation in V₁ through V₄ indicates acute anterior myocardial infarction. B, Pathologic Q waves in leads V₁ through V₅ and abnormal ST segment elevation in V₁ through V₅ indicate large acute anterior myocardial infarction. C, Acute anterior myocardial infarction. D, Same patient as in (C). Tracing made 1 hour later.
Fig. 6-10. Continued
Fig. 6-11. **A**, A patient with chest pain and ST elevation in V₁ through V₄, acute anterior myocardial infarction. (From Khan MG: *Heart Disease Diagnosis and Therapy*, Totowa, NJ, 2005 Humana Press.) **B**, Same patient as in (A). Tracing taken 10 hours later indicates evolutionary changes: Q waves have developed in V₁ through V₄, and T wave inversion has emerged. Acute anterior infarction confirmed. (From Khan, M. Gabriel: *Heart Disease Diagnosis and Therapy*, Totowa, NJ, 2005 Humana Press.)
Fig. 6-12. A, Deep pathologic Q waves in II, III, and aVF with marked ST segment elevation indicate acute inferior myocardial infarction. B, Same patient as in (A). ECG taken 1 hour later shows decrease in ST segment after thrombolytic therapy; ST-T wave abnormality indicative of evolutionary changes.
Most of the erroneous diagnoses of infarction are made based on findings of nondiagnostic Q waves in leads III and aVF (Fig. 6-14 and 6-11).

Figure 6-10 shows features of acute anterior MI.

Pathologic Q waves of infarction may diminish in amplitude and duration during the years after acute infarction. Pathologic Q waves persist in more than 80% of patients 4 to 5 years after acute MI. In some patients, abnormal coving of the ST segment and T wave inversion persist for several years and may be difficult to differentiate from a recent infarct. The tracing is often interpreted as infarction age indeterminate (Fig. 6-15 and Fig. 2-18B). In 10% of patients, Q waves become nondiagnostic but still suspicious; in the remaining 10%, Q waves disappear. In approximately 5% of patients with Q wave infarction, the ECG returns to normal.
**Fig. 6-13.** Tracing from a 49-year-old man with no evidence of heart disease; note narrow small Q waves in leads II, III, and aVF <0.04 second. Diagnosis: inferior Q waves noted, nondiagnostic, clinical correlation required: borderline ECG.

**Fig. 6-14.** Acute anteroseptal infarction in a patient presenting to the hospital 2 hours after the onset of chest pain. Note the presence of Q waves in leads V2 to V4 (left panel). After successful thrombolytic therapy, these Q waves disappear (right panel), indicating that the tissue was still salvageable. (From Wellens JJ, Conover MB: *The ECG in Emergency Decision Making*, Philadelphia, 1992, WB Saunders, Elsevier Science.)
Nonatheromatous Cause of MI

Rarely, infarction may occur in the absence of atherosclerotic coronary artery disease and can be caused by:

- Severe coronary artery spasm.
- Cocaine abuse (Fig. 6-16).
Fig. 6-16. Acute anteroseptal and inferior MI related to cocaine abuse. The patient is a 30-year-old woman known to be a cocaine user. She developed severe chest pain, and the ECG recorded 90 minutes after the onset of pain revealed ST segment elevation in the anteroseptal and inferior leads (not shown here). Coronary arteriogram revealed complete thrombotic occlusion of the proximal left anterior descending artery. Percutaneous transluminal angioplasty was performed with satisfactory result. The visualized artery was long and wrapped around the apex of the heart to supply a substantial part of the inferior wall. The ECG recorded on the next day shows signs of acute anteroseptal and inferior MI. There is a QS deflection in lead V1, and the R waves in leads V2 and V3 are very small. The ST segment in leads V1 through V3 is elevated, and the T waves are inverted in all the precordial leads and in lead I. In the inferior leads, Q waves are present with ST segment elevation and T wave inversion. (From Chou TC: Electrocardiography in Clinical Practice, 4th ed., Philadelphia, 1996, WB Saunders, Elsevier Science.)
Fig. 6-17. Acute inferior MI in a 16-year-old girl with Kawasaki disease and coronary artery aneurysm. The aneurysm was demonstrated by coronary arteriogram before the development of MI. The ECG on 5-12-80 (A) was obtained before and that on 11-8-80 (B), after the infarction occurred. Note the appearance of Q waves with ST segment elevation and T wave inversion in the inferior leads on the tracing of 11-8-80. (Courtesy Dr. Samuel Kaplan. From Chou TC: *Electrocardiography in Clinical Practice*, 4th ed., Philadelphia, 1996, WB Saunders, Elsevier Science.)

- Kawasaki disease may cause coronary artery aneurysm and MI (Fig. 6-17). A Q wave duration >0.03 second, depth >4 mm in children with symptoms suggestive of an angina can be caused by Kawasaki disease or anomalous left coronary artery arising from the pulmonary artery.
Location of Infarction

It is important to emphasize that localization of the infarcted area from the electrocardiographic findings is far from precise, particularly for anterolateral, anteroseptal, and posterior infarctions.

Inferior Infarction

• Pathologic Q waves in leads II, III, and aVF
  • Acute infarction: ST segment elevation in II, III, and aVF; diagnostic specificity is enhanced if there is reciprocal depression in leads I, aVL, V1, and V2 (see Figs. 2-13A, 5-3, 5-4, 5-8, and 6-12) during the early hours of infarction. Only ST segment elevation caused by the current of injury may be observed with no Q waves or only small emerging Q waves visible.
  • Old infarction: Pathologic Q waves may be associated with nonspecific ST-T wave changes in leads II, III, and aVF (see Fig. 2-17B), but these changes may be minor, being present in only two of the three leads; lead III is the most unreliable lead. The ST segment may be normal; the T wave inversion of acute infarction may persist indefinitely, but the ST segment and T waves may both normalize. The specificity of a Q wave of >0.03 second in leads II and aVF is 96% and the sensitivity is approximately 50%.

Anterior Infarction

• Pathologic Q, QS, or QR waves in leads V2 through V4 or V5, or V1 through V6, with extensive anterior infarction.
  • Acute infarction: ST segment elevation in leads V2 through V4 or V5 (see Figs. 5-6, 5-9, 6-10, and 6-11). Also, V1 through V6 may show ST elevation with extensive anterior infarction (see Fig. 5-7). Reciprocal depression may develop in leads II, III, and aVF during the early hours of infarction. Only ST segment elevation caused by the current of injury may be observed with no Q waves or small emerging Q waves visible.
  • Old infarction: the following: pathologic Q waves, QS complexes in V2 through V4 or V5; the ST segment is usually isoelectric but some deformity of the segment often remains to raise suspicion of an old infarct (see Figs. 2-18B-D, 6-9, and 6-15). In most patients, the ST segment is not elevated, but if it persists more than 1 month after infarction and is >1 mm in one or more leads, this suggests the presence of a left ventricular aneurysm (see Fig. 5-17). T wave inversions may partially normalize but may persist indefinitely (see Figs. 2-19, 2-18B and D, and 6-15).
**Anteroseptal or Anteroapical Infarction**

- Pathologic Q waves, QS deflection in leads V₁ through V₃ in the absence of lead misplacement, and rotational changes that may occur in some conditions, including severe emphysema (see Figs. 6-8, 6-9, and 6-14).
- Acute infarction: ST segment elevation in V₁ through V₃ in patients with acute onset of chest pain (see Fig. 5-5). This abnormality has long been attributed to anteroseptal infarction. Recent echocardiographic and angiographic findings in patients whose disorders were classified as acute anteroseptal infarction showed that 92% of patients with ST segment elevation in leads V₁ through V₃ had an anteroapical infarct with a normal septum.
- Old infarction: QS in V₁ through V₃ with deformity of the ST segment, which may be isoelectric with abnormal or normal T waves (see Fig. 6-9).

**Anterolateral Infarction**

- Pathologic Q waves in leads V₅, V₆, I, and aVL may reflect anterolateral infarction, but the pattern has low specificity. There are many conditions that can produce this electrocardiographic finding. The ECG pattern often reflects an anteroapical infarct; it may be found in patients with septal fibrosis and hypertrophic cardiomyopathy (see Fig. 6-23).
  - Acute infarction: ST segment elevation in leads I, aVL, V₅, and V₆.
  - Old infarction: Pathologic Q waves associated with an ST segment abnormality. The ST segment may be isoelectric with or without T wave inversion.
  - A QS pattern in V₄ increases the specificity (Figs. 2-19 and 6-15).

**Right Ventricular Infarction**

- Right ventricular infarction is usually associated with inferior infarction. Diagnostic ECG features are ST elevation in V₄R and V₃R in association with ST elevation and emerging Q waves in leads II, III, and aVF (Figs. 6-18 and 6-19). The ST elevation in V₄R recedes within 8 hours of onset of symptoms (see Fig. 6-18 and Fig. 5-8B).

**Posterior Infarction**

- True posterior infarction often occurs in association with inferior MI, and this association increases the specificity and makes other possibilities for tall R waves in V₁ and V₂ less likely. The possibilities must always be examined, however (see Table 2-3).
Fig. 6-18. A, Serial tracings from a patient with acute inferoposterior and right ventricular infarction. B, Note that the diagnostic changes of right ventricular infarction seen in lead V₄R have disappeared 7½ hours after the onset of pain. (From Wellens JJH, Conover MB: The ECG in Emergency Decision Making, Philadelphia, 1992, WB Saunders, Elsevier Science.)
Fig. 6-19. Example of complete atrioventricular nodal block in a patient with an acute inferoposterior myocardial infarction and right ventricular involvement. (From Wellens JJH, Conover MB: The ECG in Emergency Decision Making, Philadelphia, 1992, WB Saunders, Elsevier Science.)
Fig. 6-20. Note tall R waves in V₁ and V₂ in the absence of right ventricular hypertrophy, Wolff-Parkinson-White syndrome, or right bundle branch block, in keeping with posterior infarction. Note upright T wave in V₁ and V₂; limb leads showed inferior myocardial infarction. (From Khan, M. Gabriel: *Heart Disease Diagnosis and Therapy*, Baltimore, 1996, Williams & Wilkins.)
Fig. 6-21. V leads in a normal 26-year-old woman. Note the R wave is tall in V1; QRS is positive in V2, indicating early transition; there is no posterior infarction; R/S ratio in V1 is less than 1; the limb leads showed no pathologic Q waves in II, III, and aVF; and there is no indication of inferior or right ventricular infarction. Normal ECG.

- In lead V1, the R wave is >S, and the R/S is >1. The T wave that is often negative is upright and may be peaked (Fig. 6-20); the R wave should be >0.04 second.
- In lead V2, the R wave is tall, and the usual positive T wave may be peaked.
- Specificity is increased if the ST segment is elevated in leads V7 to V9 in the presence of the ECG pattern of acute inferior infarction. The following must receive consideration:
  - A tall R wave in V2 and occasionally in V1 is not an uncommon normal variant, particularly if transition is early (see Fig. 2-2 and Fig. 6-21).
  - Right ventricular hypertrophy, Wolff-Parkinson-White syndrome, and other causes of tall T waves in V1 and V2 must be excluded (see Table 2-3).
Fig. 6-22. Acquired immunodeficiency syndrome myocarditis simulating an anteroseptal myocardial infarction. (From Braunwald E: Heart Disease: A Textbook of Cardiovascular Medicine, 5th ed., Philadelphia, 2001, WB Saunders, Elsevier Science.)

- Presence of the diagnostic right ventricular infarction changes in V_{3R} and V_{4R} outlined previously increase the specificity.

**MIMICS OF Q WAVE MYOCARDIAL INFARCTION**

- Myocarditis, including Chagas disease and acquired immunodeficiency syndrome (Fig. 6-22), may cause pathologic Q waves.
- Pathologic Q waves may occur in patients with hypertrophic cardiomyopathy (Fig. 6-23).
- Pseudo-Q waves in leads II, III, and aVF in Wolff-Parkinson-White syndrome may mimic inferior MI (Figs. 6-24 to 6-26).
- In LVH, QS may occur in lead V_{1}, V_{2}, or V_{3} and simulate MI (Fig. 6-27).
Fig. 6-23. A, Deep, wide pathologic Q waves in leads II, III, and aVF in a 52-year-old woman with known hypertrophic cardiomyopathy. B, Same patient as in (A). Wide, deep pathologic Q waves in leads V₄ through V₆.
Fig. 6-24. Tracing from a 42-year-old woman with Wolff-Parkinson-White syndrome; note pseudo–Q waves in leads II, III, and aVF, which can mimic inferior myocardial infarction.
Fig. 6-25. Wolff-Parkinson-White syndrome mimics inferior myocardial infarction.
Fig. 6-26. Wolff-Parkinson-White syndrome; deep Q waves in leads II, III, and aVF. Mimics inferior MI. See Step 3 of the step-by-step method for accurate ECG interpretation. See also discussion in Chapter 2 regarding the necessity to include WPW syndrome early in the interpretive sequence, at the same time as assessment for blocks.
FIG. 6-27. Left ventricular hypertrophy: QS complexes in V₁ and V₂ and a minute R wave in V₃. The tracing can mimic old anteroseptal myocardial infarction, a common error of interpretation.

• Typically in LBBB, R waves are absent or minute in V₁ through V₃. LBBB can simulate anteroseptal infarction (see Figs. 2-8B and 6-6). In addition, Q waves may occur in leads II, III, and aVF in the absence of infarction.
• In some patients with emphysema, a QS pattern may be recorded in leads V₁ through V₄ and mimic anterior MI. The precordial leads should be placed one intercostal space lower than usual (Fig. 6-28).
• A left-sided pneumothorax may cause a QS pattern in leads V₁ through V₄.
• Massive pulmonary embolism may cause a QS pattern in leads V₁ through V₄ (see Chapter 10).
• Nonpenetrating chest trauma may cause Q waves simulating MI. Conditions that may cause pseudoinfarction patterns are given in Table 6-1.
Fig. 6-28. QS deflections in leads V₁ through V₄ mimic anterior myocardial infarction. Additional precordial leads were recorded one intercostal space below the levels of the routine electrode locations. ECG of a 58-year-old man with severe emphysema. The QRS complexes are partially normalized. (From Chou TC: Cardiovasc Clin 5:199, 1973.)
LEFT BUNDLE BRANCH BLOCK AND INFARCTION

The diagnosis of MI in the presence of LBBB is difficult but often can be made. The usual finding in LBBB is a QS pattern in $V_1$ and $V_3$ or very small R waves in $V_1$ through $V_3$ simulating infarction or LVH.

**ECG Features Suggestive of Left Bundle Branch Block with Infarction**

- Figure 6-29 shows the replacement of the constantly present R wave in $V_5$, $V_6$, with a Q wave of infarction in LBBB caused by anteroseptal MI. Figure 6-30 shows LBBB with inferior infarction.
- Q waves in leads I, aVL, $V_5$, and $V_6$ indicate an anterior (anterolateral or anteroseptal) infarction (Figs. 6-31 and 6-32), but a false-positive diagnosis is common. These findings often occur in the absence of infarction and may occur in patients with severe LVH or nonspecific fibrosis.
Fig. 6-29. A, With uncomplicated LBBB, early septal forces are directed to the left. Therefore, no Q waves will be seen in V₅ and V₆ (right panel). B, With LBBB complicated by anteroseptal infarction, early septal forces may be directed posteriorly and rightward (left panel). Therefore, prominent Q waves may appear in V₅ and V₆ as a paradoxical marker of septal infarction (right panel). (Adapted from Dunn MI, Lipman BS: Lipman-Massie Clinical Electrocardiography, 8th ed., Chicago, 1989, Year Book.) C, Anterior wall infarction (involving septum) with LBBB. Note the presence of QR complexes in leads I, aVL, V₅, and V₆.
Fig. 6-30. Complete left bundle branch block with acute inferior myocardial infarction. Note the prominent ST segment elevation in leads II, III, and aVF, with reciprocal ST segment depression in I and aVL superimposed on secondary ST-T changes. The underlying rhythm is atrial fibrillation. (From Braunwald E: *Heart Disease: A Textbook of Cardiovascular Medicine*, 6th ed., Philadelphia, 2001, WB Saunders, Elsevier Science.)
Fig. 6-31. Complete left bundle branch block with MI proved by autopsy. The ECG diagnosis of MI is based on the Q waves in leads I, aVL, V₅, and V₆. An autopsy performed 10 days later showed severe generalized atherosclerosis with total occlusion of the left circumflex artery. There was an extensive recent lateral wall MI in addition to a previous one. Left ventricular hypertrophy also was present. (From Chou TC: Electrocardiography in Clinical Practice, 4th ed., Philadelphia, 1996, WB Saunders, Elsevier Science.)

Fig. 6-32. Left bundle branch block and myocardial infarction. The patient was a 57-year-old man with a history of hypertension and coronary artery disease. The tracing at top was recorded after he developed congestive heart failure and experienced more frequent attacks of angina pectoris. It shows first-degree atrioventricular block, complete LBBB with a QRS duration of 0.18 second, and a digitalis effect. Six weeks later, he developed severe substernal chest pain and episodes of ventricular tachycardia. The ECG at bottom shows the loss of R waves in leads V₃ and V₄ and the development of a small Q wave in lead V₅. The patient died 5 days later. At autopsy, the heart weighed 1,200 g with notable biventricular hypertrophy. Severe coronary artery disease was present, with a massive acute anterior MI. An old inferior MI and fatty degeneration and infiltration of the interventricular septum also were observed.
• A reversal of R wave progression in the right and midprecordial leads (R waves in V₁ and V₂ that decrease in amplitude in V₃ and V₄) may indicate anterior infarction, but false-positive diagnoses are common.

**New Diagnostic Information**

Abnormal ST segment deviations occur during infarction and ischemia in patients with LBBB, and these deviations having been documented during percutaneous interventions in patients with LBBB.

• Discordant ST segment deviations are an exaggeration of normal ST segment elevation in leads V₁ to V₄ that possess a dominant S wave. Figures 6-33 and 6-34 show extensive ST segment elevation in leads V₂, V₃, and V₄ indicating acute injury pattern manifested by discordant ST segment elevation, which is equal to or exceeds the QRS amplitude in leads V₂ through V₄.

• In patients with LBBB and acute onset of chest pain, electrocardiographic features of inferior infarction are reflected by ST segment eleva-

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**Fig. 6-33.** Precordial leads of an 84-year-old man with acute myocardial infarction of the anterior wall. **A**, LBBB with acute injury pattern causes discordant ST segment elevation, which in leads V₂ and V₃ exceeds 1 mV. **B**, One day later, there is evolution of the infarction pattern without LBBB. Cardiac catheterization revealed severe three-vessel coronary artery disease with apical akinesis and a left ventricular ejection fraction of 30%. (From Chou TC: *Electrocardiography in Clinical Practice*, 5th ed., Philadelphia, 2001, WB Saunders, Elsevier Science.)
Fig. 6-34. Precordial leads of a 45-year-old man with LBBB and acute myocardial infarction of the anterior wall. **A**, Acute injury pattern manifested by discordant ST segment elevation, which is equal to or exceeds the QRS amplitude in leads V₂ through V₄. **B**, One day later, there is evolution of the infarction pattern with decreasing ST segment elevation and beginning terminal T wave inversion in leads V₂ through V₄. Note the Cabrera sign (notch on the S ascent in leads V₂ through V₄). Coronary angiography revealed a high-grade complex stenotic lesion in the proximal left anterior descending coronary artery. There was anterior apical akinesis with an estimated left ventricular ejection fraction of 40%. (From Chou TC: *Electrocardiography in Clinical Practice*, 5th ed., Philadelphia, 2001, WB Saunders, Elsevier Science.)

- Right bundle branch block (RBBB) occurs in approximately 15% of patients with acute MI.
- RBBB may be associated with deep Q waves in leads III and aVF without infarction. MI is likely only if there is an added Q wave in lead II.
- RBBB with infarction is often accompanied by left anterior fascicular block (*see* Fig. 9-8).
Rapid ECG Interpretation

**Fig. 6-35.** Right bundle branch block with acute anterior infarction. Loss of anterior depolarization forces results in QR-type complexes in the right precordial to midprecordial leads, with ST elevations and evolving T wave inversions (V₁ through V₆). (From Braunwald E: *Heart Disease: A Textbook of Cardiovascular Medicine*, 6th ed., Philadelphia, 2001, WB Saunders, Elsevier Science.)

- RBBB can be associated with Q waves in leads V₁ and V₂ without infarction. Added Q waves in V₃ and beyond suggests infarction. Figure 4-10 indicates vector forces; Fig. 6-35 shows RBBB and acute anterior MI.

**LOW-VOLTAGE QRS**

ECGs should be recorded with the graph paper moving at 25 mm/s. At this speed, a 1-mm square on the horizontal plane equals 0.04 second. The voltage of the P wave, QRS complex, and T wave are measured vertically with reference to the calibration or standardization, which should be set at 1 mV = 10 mm. With this universal standardization, a 1-mm square in a vertical direction measures 0.1 mV.

**Criteria for Low-Voltage QRS**
- In all limb leads, the amplitude of the entire QRS complex (R + S) is <5 mm.
- In each of the precordial leads, the amplitude of the entire QRS complex (R + S) is <10 mm.
Fig. 6-36. Myxedema heart disease. The patient is a 70-year-old woman with a 15-year history of myxedema. Symptoms and signs of myxedema recurred 1 year after she stopped taking her medication. She has no symptoms of coronary artery disease. The heart is not enlarged on radiographic examination. The ECG shows first-degree atrioventricular block. There is low voltage of the P waves and the QRS complexes, with abnormal left-axis deviation. The T waves are inverted in leads V1 through V3. (From Chou TC: Electrocardiography in Clinical Practice, 4th ed., Philadelphia, 1996, WB Saunders, Elsevier Science.)

**Causes of Low-Voltage QRS**

- Obesity
- Pericardial effusion
- Constrictive pericarditis
- Myxedema (Fig. 6-36)
- Amyloidosis and other restrictive cardiomyopathy and diffuse myocardial diseases
- Pleural effusion
- COPD
Atrial and Ventricular Hypertrophy

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Atrial Hypertrophy
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ATRIAL HYPERTROPHY
Left Atrial Hypertrophy

Diagnostic Criteria
- The P wave duration is ≥0.12 second (3 small squares) in leads II, III, or aVF. The P wave may be widely notched. These features are most apparent in lead II.
- The terminal deflection of the P waves in V₁ is downward and its duration is prolonged ≥0.04 second (Fig. 7-1).
- The depth of the terminal negative deflection in V₁ is ≥1 mm.
- The product of the depth of the terminal negative deflection in V₁ (in millimeters) and the duration in seconds (the P terminal force) is ≥−0.04 mm s.
- The P terminal force (PTF-V₁) is determined rapidly by observation of the P wave in V₁. The P wave terminal negative duration equal to 1 small square (0.04 second) and a depth of 1 small square (1 mm) yield a P terminal force of −0.04 mm s (Figs. 7-1 and 7-2).

Reliability of Criteria
The diagnostic changes are observed in patients with left atrial enlargement or hypertrophy without significant enlargement and in patients with an increase in left atrial pressure and volume.
- The combined sensitivity of PTF-V₁ ≥−0.04 mm s and P wave duration in II, III, or aVF >0.10 second (100 ms) has been shown to be 82%.
- A PTF-V₁ of ≥−0.06 has been shown to correctly predict left atrial enlargement in 80% of cases.

From: Contemporary Cardiology: Rapid ECG Interpretation, 3e
by: M. Gabriel Khan © Humana Press Inc., Totowa, NJ
• P wave duration >0.10 second appears to have a specificity of approximately 85% but with a low sensitivity of <33%.

Importantly, these relationships relate to echocardiographic left atrial enlargement >4 cm; left atrial hypertrophy may occasionally occur without significant echocardiographic enlargement, and the electrocar-
diographic findings may be the only sign of left ventricular disease caused by hypertension and other cardiac diseases. Thus, the term left atrial abnormality is preferred to cover enlargement, hypertrophy, or increase in atrial volume or pressure.

**Causes**

It is most important to assess for the electrocardiographic features of left atrial hypertrophy, because this may be the only abnormality in the electrocardiogram in patients with various forms of heart disease. It may be the only clue to the diagnosis of underlying left ventricular hypertrophy, valvular heart disease, congenital heart disease, ischemic heart disease that has caused heart failure, left ventricular dysfunction, the various cardiomyopathies, or constrictive pericarditis. Left atrial hypertrophy is a common ECG finding. Causes of left atrial abnormality include the following:

- Mitral stenosis.
- Mitral regurgitation.
- Left ventricular failure, particularly acute pulmonary edema in which the abnormality may decrease or disappear after approximately 1 week of successful therapy.
- Left ventricular hypertrophy (LVH). (The atrium hypertrophies in response to altered left ventricular compliance).
- Aortic valve disease.

**Right Atrial Hypertrophy**

**Diagnostic Criteria**

- The P wave is tall and peaked with a height ≥2.5 mm in lead II, III, or aVF and is of normal duration (Fig. 7-3).
- The positive component of the P wave in lead V1, V2, or V3 is tall and peaked with a height ≥1.5 mm.
- The P wave frontal axis is greater than 75 degrees.

**Helpful Guides**

- An abnormally tall P wave in leads V1, V2, and sometimes V3 ≥1.5 mm is a more specific electrocardiographic sign for right atrial hypertrophy than the usual diagnostic criteria based on peaked P waves in leads II, III, or aVF (P pulmonale).
- An initial positive component of the P wave in V1 or V2 ≥0.04 second is an indication of right atrial hypertrophy. The findings in the right chest leads are not common with cor pulmonale, however; it is more
commonly seen in children with significant congenital heart disease and in individuals with right ventricular hypertrophy.

• The typical pattern of P pulmonale seen in leads II, II, and aVF is less specific for right atrial hypertrophy than the findings in the right chest leads, but because cor pulmonale and some conditions may show no abnormalities in leads V1 to V3, it is absolutely necessary to pay attention to findings in leads II, III, and aVF.

• Right atrial hypertrophy is not a common disorder, but it is important to search diligently for electrocardiographic signs of right atrial hypertrophy because the finding may lead to the discovery of significant underlying congenital heart disease and is an important clue to the presence of right ventricular hypertrophy.

CAUSES

The causes of right atrial hypertrophy include the following:

• Congenital heart disease (some forms)
• Cor pulmonale

Fig. 7-3. Leads II, III, and aVF show tall, peaked P waves >2.5 mm right atrial hypertrophy.
Fig. 7-4. Sinus tachycardia 120 beats/min, ventricular premature beats, RSR', V₁, V₂. Lead V₁ shows characteristics of right and left atrial hypertrophy. Voltage increased V leads that satisfies the criteria for LVH; ST-T changes V₄ and V₅, which can be caused by LVH or ischemia.
• Pulmonary stenosis
• Pulmonary hypertension
• Tricuspid stenosis
• Tricuspid regurgitation
• Right ventricular hypertrophy (RVH)

**Bilateral Atrial Hypertrophy**

Bilateral enlargement is indicated by the following:

- A large biphasic P wave in lead V₁, with the positive component >1.5 mm and the initial terminal negative deflection reaching 1 mm in depth and with a duration of 0.04 second (see Figs. 7-1, 7-2, 7-4, 7-9, and Fig. 3-3).
- Peaked P waves ≥1.5 mm in V₁, V₂, and V₃.
- Notched P waves in V₄ to V₆.
- P wave amplitude ≥2.5 mm and duration ≥0.12 second in the limb leads.

**VENTRICULAR HYPERTROPHY**

**Left Ventricular Hypertrophy**

The genesis of the normal QRS complex is described in Chapter 1, and the genesis of the QRS complex in LVH is shown in Fig. 7-5. In LVH, the left atrium becomes hypertrophied to compensate for decreased compliance of the compromised left ventricle. Left atrial hypertrophy is an early ECG manifestation of LVH.

**Diagnostic Criteria for Patients Older Than 35 Years of Age**

The electrocardiographic voltage criteria for the diagnosis of left ventricular hypertrophy are imprecise and unreliable and have a low sensitivity of approximately 50%; the specificity approaches 94%.

**Supporting Evidence**

Asymmetric ST segment depression and T wave inversion in V₅ and V₆: left ventricular strain pattern; the proximal descending limb of the inverted T wave has a slow descent, and the ascending limb rises steeply. These changes should be maximal in V₅ and V₆ and minimal in V₄ (see Figs. 7-6, 2-25, and 6-27). Note that ST-T changes that are prominent in V₃ and V₄ most likely reflect ischemia. Changes in V₅ and V₆ with no changes in V₃ and V₄ may be caused by LVH, ischemia, or both (see Fig. 8-12A and B).
The QRS duration must be <0.12 second.

1. Sokolow-Lyon Voltage Criteria
   - R wave in lead I + S wave in lead III > 25 mm (2.5 mV)
   - R wave in aVL > 11 mm (1.1 mV)
   - R wave in V₆ > 26 mm (2.6 mV)
   - R wave in V₆ + S wave in V₁ > 35 mm (3.5 mV) (Figs. 7-6 and 2-25)

   The criteria reportedly have a 49% sensitivity and a specificity of approximately 90%.
   - Left axis is supportive of LVH but is not necessary for the diagnosis.
   - Onset of intrinsicoid deflection in V₅ or V₆ > 0.05 second.

2. Cornell Voltage Criteria
   - S wave in V₃ + R wave in aVL > 28 mm (2.8 mV) in men or > 20 mm (2.0 mV) in women; the sensitivity is approximately 49% and the specificity is approximately 90%.

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**Fig. 7-5.** The contribution of vector II to the ECG features of left ventricular hypertrophy. The thicker the left ventricular muscle, the greater the magnitude of vector II; thus, the deep S wave in lead V₁ and tall R wave in leads V₅ and V₆. Note that the T wave has a gradual descending and a steep ascending limb “strain pattern.” (From Khan, M. Gabriel: *On Call Cardiology*, 2nd ed., Philadelphia, 2001, WB Saunders, Elsevier Science.)
3. Author’s Criteria for Patients Older Than 35 Years of Age: 3 points = probable LVH; 4 points = significant LVH
   - S wave in V₁ + R wave in V₆ ≥ 35 mm = 2 points
   - R wave in aVL + S wave in V₃ > 28 mm in men or > 20 mm in women = 3 points
   - Left atrial enlargement with P terminal force ≥ −0.04 mm s or P wave duration ≥ 0.12 in leads II, III, or aVF = 2 points
   - Asymmetric ST depression in V₅ and V₆ = 2 points

Fig. 7-6. Significant increase in voltage of R wave in V₅ or V₆ and S wave in V₁ or V₂ > 35 mm. There is asymmetric ST segment depression and T wave inversion in V₅ and V₆, features typical of left ventricular hypertrophy; note the lack of ST-T changes in V₄.
4. Romhilt-Estes Scoring System
   - R wave in the limb leads ≥20 mm, S wave in lead V₁ or V₂ ≥30 mm, or R wave in lead V₅ or V₆ ≥30 mm = 3 points
   - Negativity of P wave in V₁ >1 mm in depth with duration >0.03 second = 3 points
   - ST-T wave changes (if patient is not taking digoxin) = 3 points (if patient is taking digoxin = 1 point)
   - Presence of left axis = 2 points
   - A score of 4 points indicates probable LVH, and a score of 5 or more points indicates LVH. The sensitivity is approximately 30% and the specificity is approximately 90%.

**Pitfalls in Diagnosis of Left Ventricular Hypertrophy**

The preceding criteria do not apply in subjects younger than age 35 years, because QRS voltage can be notably increased in healthy young individuals (Fig. 7-7; see Table 2-1).

- QRS voltage appears to be increased by left anterior fascicular block.
- QRS voltage is higher in African Americans than in Caucasians.
- Conditions that decrease QRS voltage and that may mask the ECG signs of LVH include severe chronic obstructive pulmonary disease; pericardial effusion; large, old anterior infarctions; myxedema (see Fig. 6-34); and heart muscle diseases such as dilated cardiomyopathy, amyloidosis, and scleroderma.

**Right Ventricular Hypertrophy**

The QRS duration must be <0.12 second because the diagnosis of RVH cannot be made accurately in the presence of right bundle branch block (RBBB) or Wolff-Parkinson-White (WPW) syndrome, posterior MI, dextroposition.

**Diagnostic Criteria for Patients Older Than 30 Years of Age**

Two or more of the following criteria are required for the diagnosis of RVH:

- Right-axis deviation greater than +110 degrees
- Tall R wave in V₁ ≥7 mm (can be a normal variant), S wave in V₁ ≤2 mm, R/S ratio in V₁ >1, R/S ratio in V₅ or V₆ ≤1 (Fig. 7-8), patient older than age 30 years (see Table 2-1)
- S wave in V₅ or V₆ >2 mm
- qR pattern in V₁ (not commonly observed but increases specificity) (Figs. 7-9 and 7-10)
Fig. 7-7. High QRS voltage: S in V₁ + R in V₅ or V₆ = 54 mm (5.4 mV) in a normal 24 year old male. Caution with voltage criteria in individuals younger than age 30 and think of LVH only if additional features are present (left atrial hypertrophy, and/or ST-T changes, the strain pattern).
Fig. 7-8. Severe mitral stenosis. The patient is a 47-year-old man with severe mitral stenosis proved at surgery. In the ECG, the P waves are consistent with biatrial enlargement. The abnormal right axis deviation with an R/S ratio greater than 1 in lead V1 and the T wave inversion in the right precordial leads are consistent with right ventricular hypertrophy. (From Chou TC: *Electrocardiography in Clinical Practice*, 4th ed., Philadelphia, 1996, WB Saunders, Elsevier Science.)

Fig. 7-9. Right ventricular hypertrophy. Q wave in V1 suggests that the right ventricular pressure exceeds the left ventricular pressure. The left axis of the P wave and the prominent negative P wave in lead V1 are unusual and most likely are a result of a markedly enlarged right atrium projecting to the left and posteriorly. The prominent P waves in leads V2 and V3 are diagnostic of right atrial enlargement. (From Braunwald E: *Heart Disease: A Textbook of Cardiovascular Medicine*, 5th ed., Philadelphia, 2001, WB Saunders, Elsevier Science.)
Fig. 7-10. Right ventricular hypertrophy pattern most consistent with severe pressure overload. Note the combination of findings, including (1) a tall R wave in V1 (as part of the qR complex), (2) right-axis deviation, (3) T wave inversion in V1 through V3, (4) delayed precordial transition zone (rS in V6), and (5) right atrial abnormality. An S1Q3 pattern is also present and can occur with acute or chronic right ventricular overload syndrome. (From Braunwald E: *Heart Disease: A Textbook of Cardiovascular Medicine*, 5th ed., Philadelphia, 2001, WB Saunders, Elsevier Science.)
• Most important is scrutiny for right atrial hypertrophy: peaked P waves with an amplitude in V₁, V₂, or V₃ ≥1.5 mm (see Fig. 7-9) or ≥2.5 mm in II, III, or aVF (this increases specificity)

**Supporting Evidence**

• Onset of intrinsicoid deflection in V₁ = 0.035 to 0.055 second
• ST-T strain pattern in V₁ through V₃ (see Fig. 7-9)
• Right atrial enlargement

**Pitfalls in Diagnosis of Right Ventricular Hypertrophy**

RVH and right atrial enlargement are uncommon ECG diagnoses; cardiologists should refrain from making an ECG diagnosis of RVH in the presence of the following conditions:

• RBBB
• WPW syndrome
• True posterior myocardial infarction
• In children (the preceding ECG findings can be a normal variant)
• Early transition (the R wave is increased in V₁ and V₂, but the R/S ratio in V₅ or V₆ is greater than 1)
• Dextroposition (see Table 2-3)
• Hypertrophic cardiomyopathy (a tall R wave in V₁ with an R/S ratio greater than 1 may be observed)
The T wave represents repolarization, the recovery period of the ventricles. As emphasized in Chapter 5, the skillful interpreter focuses on the ST segment and does not try to make diagnoses based on T wave changes. T wave changes often are nonspecific and always should be interpreted in light of associated abnormalities of the ST segment and clinical findings. An algorithmic approach for the interpretation of T wave changes is depicted in Fig. 8-1.

NORMAL DIRECTION OF T WAVE

• The T wave is always upright (positive) in leads I (1), II (2), and V₄ through V₆ (Figs. 8-2 and 8-3).
• The T wave is normally upright in lead aVF if the QRS complex is <5 mm tall, but the T wave can be flat or inverted.
• The T wave is variable in leads III and aVL.
• The T wave is always inverted in aVR (see Fig. 8-2).
• The T wave in V₁ is inverted in approximately 50% of women and in <33% of men (see Fig. 8-2).
• In women with a persistent juvenile pattern, the T wave is inverted in V₁ and V₂ and sometimes in V₃ (Fig. 8-4). This finding is common in African-American women.
• Diagnoses based solely on the appearance of abnormal-looking T waves are fraught with danger. Figures 8-4 to 8-6 indicate errors that can be made in the interpretation of T or ST-T wave changes.
Assess the pattern of T wave changes.

- **Peaked**
  - V1 to V6
  - Hyperkalemia, ↑K (see Chapter 10)
  - Consider posterior MI (see Chapter 6)
- **Flat**
  - V1 and V2
  - Nonspecific ST-T changes (see Figures 2-8, 8-9, and 8-11 and Chapter 5)
- **Inverted**
  - Q waves or ST elevation or depression
  - Definite ischemia (see Figure 8-8)
  - Probable ischemia (see Figure 8-7)
  - Digitalis effect
  - LVH (see Figure 2-25)
  - V1 to V3 consider pulmonary embolism (see Figure 10-24) or RVH (see Figures 2-26 and 7-8)

**If abnormal, assess if associated with ≥1 mm ST depression, ST elevation, or an abnormally shaped ST segment.**

**Upright:** Leads I, II, and V3 to V6
- Associated with
  - V1 and V2 associated with Q waves or ST elevation or depression
  - Definite ischemia (see Figure 8-8)
  - Probable ischemia (see Figure 8-7)
  - Digitalis effect
  - LVH (see Figure 2-25)
  - V1 to V3 consider pulmonary embolism (see Figure 10-24) or RVH (see Figures 2-26 and 7-8)

**Deep T inversion >5 mm**
- a. Localized V2 to V3: likely ischemia or post-MI
- b. Localized II, III, aVF: likely ischemia or post-MI
- c. Diffuse: cardiomyopathy or other nonspecific

**Definite ischemia** (see Figure 8-8)
- a. V5 and V6, less in V4: LVH (see Figure 2-25)
- b. V1 to V3: RVH or embolism (see Figures 2-26, 7-8, and 10-24)
- c. Diffuse: cardiomyopathy or other nonspecific

**Peaked**
- High K⁺ level (see Chapter 10)
- Normal variant

**Minor inversion <5 mm**
- Nonspecific ST-T changes (see Figures 8-9, 8-11, and 2-28)
- Consider
  - Ischemia (see Chapter 6)
  - Electrolyte depletion
  - Alcohol
  - Cardiomyopathy
  - Myocarditis
  - Normal variant
  - Other

---

**Fig. 8-1.** Step-by-step method for accurate ECG interpretation. **A**, Step 8: Assessment of T wave changes. LVH, Left ventricular hypertrophy; MI, myocardial infarction; RVH, right ventricular hypertrophy. **B**, Step 8: Alternative methods for the assessment of T wave changes.
Chapter 8 / T Wave Abnormalities

T wave always inverted
aVR

Variable lead 3
aVL*
aVF*
V₁**
V₂***
V₃****

* = usually upright; can be inverted if R wave < 5 mm.
** = inverted in >50% of women and <20% of men who are >30 years of age.
*** = usually upright; can be inverted with juvenile pattern.
**** = usually upright; rarely flat or biphasic in women or with juvenile pattern.

Fig. 8-2. Normally occurring negative T waves in aVR and V₁.

Fig. 8-3. T wave, normal variability. (From Khan, M. Gabriel: On Call Cardiology, 3rd ed., Philadelphia, 2006, WB Saunders, Elsevier Science.)
Fig. 8-4. Sinus rhythm 80/ min; ST segment coving with juvenile T wave inversion in leads V₁–V₃ : normal variant in this 10 year old male. Also, normal variant ST elevation in II, III, aVF, V₅, V₆ (so called early repolarization changes) The diffuse ST segment elevation and PR elevation in aVR might suggest pericarditis. But, in pericarditis the J point level almost equals the height of the T wave in V₆. Clinical correlation required.
Fig. 8-5. Prominent T waves in a healthy 31-year-old man. ST segment elevation also is present in the precordial leads. They may be mistaken as signs of myocardial injury and ischemia or acute pericarditis. (From Chou TC: Electrocardiography In Clinical Practice, 4th ed., Philadelphia, 1996, WB Saunders, Elsevier Science.)

Fig. 8-6. Benign ST and T wave changes in a healthy 24-year-old professional athlete. The changes, especially in leads V₄ and V₅, mimic myocardial injury and ischemia and remained the same 15 months later. (From Chou TC: Electrocardiography in Clinical Practice, 4th ed., Philadelphia, 1996, WB Saunders, Elsevier Science.)
ABNORMALITIES OF T WAVE

Inverted T Wave

• T wave inversion in leads I, II, and V3 through V6 is abnormal.
• If T wave inversion is accompanied by abnormal coving of the ST segment (horizontal or down-sloping ST segment depression >1 mm; Figs. 8-7 and 8-8), a diagnosis of ischemia can be made with confidence.

Fig. 8-7. T wave inversion in V2 through V5 associated with abnormal curvature of the ST segment; likely caused by ischemia.
Fig. 8-8. Tracing from a 53-year-old woman with a 1-week history of unstable angina. Tracing taken in the absence of pain. Deep T wave inversion in V_2 through V_4. Note the abnormal coving of the ST segment and “hitched-up” ST segment in V_1 and V_2. Definite ischemia is indicated.

- If T wave inversion is associated with <1-mm ST depression or an up-sloping depression, the finding is nonspecific (Fig. 8-9) and can be caused by a host of cardiac and noncardiac conditions.
- Isolated T wave inversion is nonspecific (Figs. 8-10 and 8-11) but ischemia cannot be excluded.
- The 5-year mortality rate in patients with moderate T wave inversion associated with the presence of heart disease reportedly is 21% versus 3% when heart disease is absent.

Diffuse, deep T wave inversion in the absence of ST segment elevation or significant depression is not diagnostic (see Fig. 8-10) and can be associated with the following:

- Ischemia
- Post-MI evolutionary changes
• Left ventricular hypertrophy with or without ischemia (Fig. 8-12)
• Post–Stokes-Adams attack
• Post–supraventricular tachycardia or ventricular tachycardia
• Myocarditis
• Pericarditis
• Apical cardiomyopathy (causes giant T wave inversion) (Fig. 8-13)
• Pulmonary embolism
• Cardiomyopathies
• Primary or secondary cardiac tumors
• Cocaine abuse

**Fig. 8-9.** Minimal T wave inversion in V₂ through V₄ with <1-mm ST depression: nonspecific ST-T changes; cannot exclude ischemia.
Alcohol abuse
Electrolyte imbalance
Subarachnoid hemorrhage (see Figs. 8-13 and 8-14)
Acute pancreatitis and gallbladder disease
Pheochromocytoma
Other causes

Symmetric T wave inversion is four times more common in women than it is in men. Interpreting symmetric, deep T wave inversion as a sign of ischemia without considering other diagnoses is a common error.

Minor T wave inversion not associated with significant ST segment changes can be caused by all of the aforementioned conditions, as well as by the following:

Fig. 8-10. T wave inversion in V₄ through V₆; similar changes were observed in leads aVL, II, III, and aVF: nonspecific ST-T changes; cannot exclude ischemia.
Fig. 8-11. Tracing from a 50-year-old man with no history of heart disease; non-specific ST-T wave changes as seen from V1 through V3; the limb leads show no abnormality. Abnormal ECG.

- Hyperventilation
- Postprandial (after the patient has a meal or a cold drink, the tracing normalizes in the fasting state)
- Mitral valve prolapse
- Intraventricular conduction defects
- Pneumothorax
- Ventricular hypertrophy (see Chapter 7)

Minor T wave inversion not associated with significant ST segment changes also can be a normal variant. T wave inversion occurs in V1 through V3 in some young adults as a persistent juvenile pattern; this is more common in women (see Fig. 8-4). Benign T wave inversion in V4 through V6 may be observed in healthy young adults and may be associated with ST elevation as a normal variant (see Chapter 5).
Fig. 8-12. A, Voltage increase: probable left ventricular hypertrophy; compare with (B). B, Same patient 12 hours later: The ST-T in V₄ through V₆, which was caused by ischemia, may have been interpreted incorrectly as left ventricular hypertrophy with “strain” pattern if no comparison was available.

(continued)
Fig. 8-12. Continued
Deep T Wave Inversions: Selected Examples

Ischemia

CVA

Apical HCM

Fig. 8-13. Deep T wave inversion can result from a variety of causes. Note the significant QR prolongation in conjunction with the cerebrovascular accident (CVA) T wave pattern caused here by subarachnoid hemorrhage. Apical hypertrophic cardiomyopathy (HCM) is another cause of deep T wave inversion that can be mistaken for coronary disease. (From Goldberger AL: ACC Curr J Rev Nov/Dec:28, 1996.)

Fig. 8-14. Patient with subarachnoid hemorrhage. ECG shows many of the findings commonly associated with central nervous system lesion. Careful autopsy examination revealed mild left ventricular hypertrophy and dilation but no myocardial damage. (From Chou TC, Susilavorn B: J Electrocardiol 2:193, 1969. By permission.)
**Tall T Waves**

The height of the normal T wave is usually <5 mm in the limb leads and <10 mm in any precordial lead. T waves that are >6 mm in the limb leads or >10 mm in the precordial leads may occur as follows:

- In V2 through V5 in some normal individuals. Note the base of normal peaked T waves is not narrow, as it is with hyperkalemia (see Fig. 8-5 and Fig. 10-8). Peaked T waves occasionally may be associated with ST elevation occurring as a normal variant; the ST elevation is commonly and inappropriately interpreted as repolarization changes (see Chapter 5).
- In patients with severe myocardial ischemia or acute MI (hyperacute T waves may occur).
- In patients with hyperkalemia (see Fig. 10-8 and Chapter 10).
- In patients with left ventricular overload, as in severe mitral regurgitation.
- Occasionally, in patients with cerebrovascular accidents.

**U WAVES**

Figure 8-15 shows a variety of U waves.

![Fig. 8-15](image-url)  
*Fig. 8-15. U waves. Upper row. Upright U waves: A, Normal. B, C, and D, Prominent U waves in hypokalemia. Bottom row. Inverted U waves: A, Tracing from which this was taken showed no abnormalities except for U wave inversion in several leads. This situation is referred to as isolated U wave inversion. B, From a patient with hypertension whose tracing showed left ventricular strain including inverted U waves. C, From a patient with coronary insufficiency but without hypertension. D, Note marked inversion of T wave and U wave; from a patient with hypertension. (From Marriott JHL: Practical Electrocardiography, Baltimore, 1988, Williams & Wilkins.)*
Normal U Waves

- The U wave is a very small wave that follows the T wave and is observed only in some individuals. In the normal subject, the U wave is virtually always upright if the T wave is upright.
- The U wave is best visible in leads V3 and V2 (looks like the hump on a camel’s back; see Fig. 8-15 and Fig. 10-7). It is barely visible in other leads, and its electrophysiologic source remains uncertain.
- The U wave may merge with the T wave, and the QU interval may be measured, causing a falsely lengthened QT interval.
- The U wave coincides with the phase of supernormal excitability during ventricular recovery, and most ventricular premature beats occur around the time of the U wave.

Causes

U waves are considered large when the amplitude is ≥1.5 mm. The causes of prominent U waves include the following:

- Hypokalemia
- Digitalis use
- Quinidine use
- Hypercalcemia
- Intracranial hemorrhage
- Thyrotoxicosis

Abnormal U Waves

A negative U wave is rarely recorded in normal individuals.

- The most common cause of U wave inversion is severe hypertension, systolic or diastolic overload.
- Rarely, U wave inversion may be the earliest ECG sign of acute coronary syndrome.
- U wave inversion may be the only ECG finding in acute ischemia.
- Exercise-induced transient U wave inversion has been correlated with left anterior descending artery stenosis.
Electrical Axis and Fascicular Block

CONTENTS
Electrical Axis
Fascicular Block

ELECTRICAL AXIS

The electrical axis is discussed early and extensively in most books on ECG interpretation. Although the electrical axis is an important parameter that should be documented, it provides little or no assistance in the diagnosis of most cardiac conditions, particularly those that require specific therapy.

Determination of the electrical axis is useful mainly in the supporting diagnosis of 4 of the 100 or more diagnoses made from ECG tracings:

1. Left anterior fascicular block (LAFB) (hemiblock).
2. Right ventricular hypertrophy (RVH). Right-axis deviation (RAD) is usually a feature. The electrical axis is of minor assistance in the diagnosis of left ventricular hypertrophy (LVH); left axis is not necessary for the diagnosis of LVH.
3. Ventricular tachycardia (VT). Some forms of VT are associated with left-axis deviation (LAD) or an axis in “no man’s land,” but RAD may occur in some.
4. Left posterior fascicular block (LPFB). Criteria for the diagnosis of LPFB are imprecise and unreliable.

Figure 9-1 shows the vectorial genesis of the QRS complex and axis. The addition of all the vectors of ventricular depolarization produces one large mean QRS vector. The QRS axis represents the direction of the mean QRS vector in the frontal plane. The electrical axis is determined by using the hexaxial reference system, which was derived from Einthoven’s equilateral triangle (Fig. 9-2).

From: Contemporary Cardiology: Rapid ECG Interpretation, 3e
by: M. Gabriel Khan © Humana Press Inc., Totowa, NJ

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Fig. 9-1. The mean QRS vector.

Because of the minor contribution of the electrical axis to clinical cardiologic diagnosis, this topic is discussed late in this text and is relegated to Step 9 in the 11-step method for accurate ECG diagnosis. See Fig. 9-3, Table 9-1, and instructions given in Chapter 2 for the determination of the electrical axis.

**STEP 9**

[Diagram of the electrical axis determination process]

**Fig. 9-3.** A, Step-by-step method for accurate ECG interpretation. Step 9: Detection of the electrical axis. Leads are indicated in parentheses. See Table 9-1.  
(continued)
**Axis**

<table>
<thead>
<tr>
<th>Normal</th>
<th>Lead I</th>
<th>aVF</th>
<th>II</th>
<th>III</th>
<th>aVR</th>
<th>aVL</th>
</tr>
</thead>
<tbody>
<tr>
<td>+45°</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>+60°</td>
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<tr>
<td>-45°</td>
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<td></td>
</tr>
<tr>
<td>-60°</td>
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<td></td>
</tr>
<tr>
<td>+150°</td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
</tbody>
</table>

**STEP 9 Continued**

*Most equiphasic lead.*

**Fig. 9-3. Continued B,** See Table 9-1 and Figs. 9-4 and 9-5.

**Table 9-1**

**Electrical Axis**

<table>
<thead>
<tr>
<th>Most equiphasic lead</th>
<th>Lead perpendicular*</th>
<th>Axis</th>
</tr>
</thead>
<tbody>
<tr>
<td>III</td>
<td>aVR</td>
<td>Normal = +30 degrees</td>
</tr>
<tr>
<td>aVL</td>
<td>II</td>
<td>Normal = +60 degrees</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Lead I positive and aVF negative = left axis</td>
</tr>
<tr>
<td>II</td>
<td>aVL (QRS positive)</td>
<td>Left = −30 degrees</td>
</tr>
<tr>
<td>aVR</td>
<td>III (QRS negative)</td>
<td>Left = −60 degrees</td>
</tr>
<tr>
<td>I</td>
<td>aVF (QRS negative)</td>
<td>Left = −90 degrees</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Lead I negative and aVF positive = right axis</td>
</tr>
<tr>
<td>aVR</td>
<td>III (QRS positive)</td>
<td>Right = +120 degrees</td>
</tr>
<tr>
<td>II</td>
<td>aVL (QRS negative)</td>
<td>Right = +150 degrees</td>
</tr>
</tbody>
</table>

*Lead perpendicular (at right angle) to the most equiphasic (isoelectric) lead usually has the tallest R or deepest S wave.
• The range of the electrical axis in the majority of normal adults older than the age of 40 years is −30 degrees to +90 degrees (see Fig. 9-3); for those younger than age 40, the range is 0 degrees to +105 degrees. Normal children may have an axis of up to +110 degrees. Most normal individuals have values between +30 degrees and +75 degrees.

**Left- Axis Deviation**

An axis of −15 degrees to −30 degrees, which is relatively normal for individuals older than the age of 40 years, is sometimes termed leftward axis to distinguish it from LAD. An axis of −30 degrees to −90 degrees is termed marked LAD (Fig. 9-4).

![Fig. 9-4. Left-axis deviation −45 degrees; lead II is the most equaphasic QRS; aVL is perpendicular and lies at −30 degrees; aVR is the next most equaphasic; lead III is perpendicular at −60 degrees; therefore, the axis that lies between = −45 degrees (see Fig. 9-3B).](image-url)
**Causes**
- Normal variation
- LAFB (hemiblock)
- Left bundle branch block
- LVH
- Mechanical shifts causing a horizontal heart; high diaphragm; pregnancy, ascites
- Some forms of VT
- Endocardial cushion defects and other congenital heart disease

**Right-Axis Deviation**
- Criteria for RAD in adults include an axis of +100 degrees to +180 degrees (Fig. 9-5)

---

**Fig. 9-5.** A, Right-axis deviation; QRS axis +110 degrees; lead I is the most equiphasic; aVF is perpendicular at +90 degrees; aVR is the next most equiphasic, with lead III being perpendicular at +120 degrees. The exact axis lies somewhere between +90 degrees and +120 degrees (i.e., at +110 degrees). Patient is older than 40 years of age. B, Right-axis deviation. ECG from a 26-year-old healthy man.


**Causes**

- Normal variation
- RVH
- LPFB
- Lateral MI
- Pulmonary embolism
- Dextrocardia
- Normal variants: mechanical shifts or emphysema causing a vertical heart

\[\text{Fig. 9-5. Continued}\]
**FASCICULAR BLOCK**

*Left Anterior Fascicular Block (Hemiblock)*

Figure 9-6 illustrates the division of the left bundle branch into anterior and posterior fascicles. The anterior fascicle traverses an antero-superior course and ends at the base of the anterior papillary muscle. The anterior fascicle is thin and long, has a single blood supply, and is commonly damaged by ischemic-disease fibrosis and other pathologic processes, resulting in LAFB. Rosenbaum initially called the block left anterior hemiblock, and because this term is easy to write, many cardiologists use it rather than LAFB.

![Diagram of hemiblock patterns in the limb leads: left anterior hemiblock (LAH) and left posterior hemiblock (LPH). The “anterior” papillary muscle is above and lateral to the “posterior” papillary muscle, and the two divisions of the left bundle branch course toward their respective papillary muscles. Thus, if the anterior division is blocked, initial electromotive forces are directed downward and to the right, inscribing a small Q wave in leads I and aVL and an S wave in leads II, III, and aVF. The subsequent forces are directed mainly upward and to the left, writing an R wave in I and aVL and an S in II, III, and aVF, to produce a left-axis deviation. In LPH, the initial forces spread upward and to the left to write an R in I and aVL and a small Q in II, III, and aVF while subsequent forces are directed downward and to the right to produce right-axis deviation. (From Marriott JLH: *Practical Electrocardiography*, 8th ed., Baltimore, 1988, Williams & Wilkins.)*

*Fig. 9-6.*
**Diagnostic Criteria**

- LAD, −45 degrees to −90 degrees (preferably −60 degrees to −90 degrees)
- A small q wave, 0.5 to less than 2 mm deep in lead I, qR in I (Fig. 9-7)

**Fig. 9-7.** Left-axis deviation −60 degrees. There is a small q wave in lead I and a small r wave in lead III. These are the criteria for the diagnosis of left anterior fascicular block: borderline ECG.
• A small r wave, 1 to 4 mm tall in lead III, rS in III
• A normal QRS duration, provided right bundle branch block (RBBB) or other conduction defects are absent (other fascicles conduct normally; thus depolarization of the ventricles is not delayed)

Figure 9-6 shows how the ECG configuration of LAFB is derived. With block of the anterior fascicle, depolarization starts at the posterior papillary muscle and inferior wall and proceeds upward, superiorly and to the left to activate the left ventricular muscle mass that lies above the papillary muscle. Thus, the electrical axis is directed strongly to the left at −60 degrees to −90 degrees. Because the electrical impulse originating from the posterior papillary muscle travels initially downward from endocardium to epicardium, it registers a small r wave of <4 mm in lead III. The small impulse is directed away from lead I and thus causes a small q wave in lead I. The current then travels upward to the left and causes an r wave in lead I and an s wave in lead III, because electrical impulse travels away from the inferior leads (see Figs. 9-4 and 9-6).

CAUSES
• Acute or chronic ischemic heart disease
• Cardiomyopathy and specific heart muscle disease
• Chagas disease
• Myocarditis

LAFB is a normal finding in approximately 1% of men older than age 40 years.

PITFALLS IN DIAGNOSIS OF LEFT ANTERIOR FASCICULAR BLOCK
• Acute or old MI: When LAFB occurs during acute inferior MI, the initial small r wave caused by LAFB in leads II, III, and aVF masks the Q wave of infarction.
• Hypertensive heart disease: LAFB lowers the QRS voltage in the precordial leads and may mask LVH; conversely, LAFB increases QRS voltage in the limb leads and may mimic LVH.

Left Posterior Fascicular Block

LPFB or posterior hemiblock occurs rarely, because the posterior bundle is thick and short and has a double blood supply. The fascicle runs to the base of the posterior papillary muscle (see Fig. 9-6). The
diagnosis can be made only after excluding RVH and chronic obstruc-
tive pulmonary disease (COPD). The following are criteria for the
diagnosis of LPFB:

- RAD, +120 degrees to +180 degrees
- A small r wave <4 mm in leads I and aVL and an S wave in lead I
- A small q wave in lead II or III
- A normal QRS duration
- Absence of RVH or cor pulmonale, COPD, a vertical heart, and other
causes of RAD (thus, a confident diagnosis of LPFB is not often made)

**Bifascicular Block**

- The combination of LAFB and RBBB occurs commonly (Fig. 9-8) but
rarely progresses to serious block; thus pacing is rarely required.
- The combination of LPFB and RBBB occurs rarely (Figs. 9-9 and
9-10).

---

Fig. 9-8. A, V leads from a patient with right bundle branch block (RBBB). B,
Features of left anterior fascicular block (LAFB). The QRS axis is −75 degrees
with a small Q wave in lead I and a small R wave in lead III, which is in keeping
with LAFB. Diagnosis: bifascicular block: RBBB and LAFB.

(continued)
Fig. 9-8. Continued

Fig. 9-9. A, Tracing of a patient with right bundle branch block and left posterior fascicular block: bifascicular block. B, Left posterior fascicular block.
Fig. 9-9. Continued
Fig. 9-10. Left posterior hemiblock. Note the right-axis deviation, the small r in leads I and aVL, and the small q in lead II. Complete right bundle branch block is also present in this patient. (From Wellens JJH, Conover MB: The ECG in Emergency Decision Making, Philadelphia, 1992, WB Saunders, Elsevier Science.)
After a methodical assessment of the P waves, the QRS duration for bundle branch blocks (left and right), the ST segment, Q waves, hypertrophy, and the electrical axis has been completed, an assessment for miscellaneous conditions is appropriate. A search for miscellaneous conditions is logically performed at Step 10 (Fig. 10-1).

The ECG may reveal clues to the diagnosis of 12 or more miscellaneous conditions:

- Atrial septal defect
- Acute pericarditis
- Long QT interval
- Hypokalemia
- Hyperkalemia
- Digitalis toxicity
- Dextrocardia
- Electrical alternans
• Electronic pacing
• Pulmonary embolism (PE) (the ECG is not diagnostic)
• Hypothermia and hyperthermia
• Hypercalcemia and hypocalcemia

**ATRIAL SEPTAL DEFECT**

Incomplete right bundle branch block (RBBB) is a common and well-known finding in atrial septal defect (see Fig. 2-34).

**A New Diagnostic Sign**

A characteristic “crochetage” on the R wave of leads II, III, and aVF has been reported (Fig. 10-2).
PERICARDITIS

Diagnostic Criteria

• Stage 1: Widespread ST segment elevation, generally upwardly concave in all leads except aVR and occasionally V₁. ST segment elevation may persist for a few days (Figs. 10-3, 10-4, and Fig. 2-33). Reciprocal ST segment depression occurs in aVR and sometimes in V₁ (see Figs. 10-3 and 10-4). The PR segment is elevated in aVR, and PR segment depression generally occurs in all leads except occasionally V₁.

• Stage 2: A few days later, the ST and PR segments become normal (isoelectric); the T wave remains normal or may be decreased in amplitude and may become flattened.

• Stage 3: After normalization of the ST segment, diffuse T wave inversion occurs.

• Stage 4: This stage lasts from days to weeks. The T waves normalize; rarely do they remain inverted.

Other Clues Suggestive of Acute Pericarditis

• Early PR segment depression, particularly in leads II, aVF, and V₄ through V₆, is suggestive of pericarditis.

• Sinus tachycardia may be the only finding if ST segment elevation has resolved and the T waves remain normal.
Fig. 10-3. Widespread ST segment elevation, generally upwardly concave in all leads except aVR and $V_1$: acute pericarditis. (From Khan, M. Gabriel: *Heart Disease Diagnosis and Therapy*, Baltimore, 1996, Williams & Wilkins.)
Fig. 10-4. Characteristic features of acute pericarditis: ST segment elevation in most leads: I, II, aVL, aVF, V5, and V6, with reciprocal ST depression and PR segment elevation in aVR. In addition, note sinus tachycardia commonly seen with acute pericarditis.
• Electrical alternans usually involves the QRS complex. Total alternans with involvement of the P, QRS, and T waves may occur with cardiac tamponade (see Fig. 10-14).
• Low-voltage QRS may occur when pericardial fluid accumulates.

**LONG QT INTERVAL**

The QT interval indicates the total duration of ventricular systole. A prolonged QT interval represents delayed repolarization of the ventricles and predisposes to reentrant arrhythmias, such as torsades de pointes (see Fig. 11-46).

**Diagnostic Criteria**

• A rough guideline to remember is that the QT interval should be less than half the preceding RR interval at heart rates of 60 to 100 beats/min.
• The QT interval varies with heart rate, and several formulas have been used to provide a corrected QT interval (QTc).
• The QTc also has limitations because of difficulties with obtaining exact measurements. Because it is difficult sometimes to define the end of the T wave, the measurement is often inaccurate, particularly when a U wave merges with the T. Thus in clinical practice, the QT interval should be assessed mainly for excessive prolongation, using a lead that does not show a U wave (Figs. 10-5 and 10-6).
• See Table 2-5 for a clinically useful approximation of QT intervals.

**Causes**

A prolonged QT interval may be caused by the following:

• Drugs
  • Class 1 antiarrhythmics (e.g., disopyramide, procainamide, quinidine)
  • Class 3 antiarrhythmics (e.g., amiodarone, sotalol)
• Tricyclic antidepressants
  • Phenothiazines
  • Astemizole
  • Terfenadine
  • Adenosine
  • Antibiotics (e.g., erythromycin and other macrolides)
  • Antifungal agents
  • Pentamidine, chloroquine
• Ischemic heart disease
• Cerebrovascular disease
Fig. 10-5. QT interval 0.46 second; the heart rate is 67 bpm. The normal range of a QT interval at a heart rate of 67 to 100 bpm is 0.33 to 0.42 second (see Table 2-5).

Fig. 10-6. The QT interval is prolonged, measuring approximately 600 milliseconds, with T wave alternans. The tracing was recorded in a patient with chronic renal disease shortly after dialysis. (From Braunwald E: Heart Disease: A Textbook of Cardiovascular Medicine, 5th ed., Philadelphia, 1997, WB Saunders, Elsevier Science.)
• Rheumatic fever
• Myocarditis
• Mitral valve prolapse
• Electrolyte abnormalities
• Hypocalcemia
• Hypothyroidism
• Liquid protein diets
• Organophosphate insecticides
• Congenital prolonged QT syndrome

A short QT interval is not of great concern and occurs rarely with the following:
• Hypercalcemia, a feature of malignancy and hyperparathyroidism
• Digitalis intoxication

HYPOKALEMIA

Diagnostic Criteria

• Progressive ST segment depression: A small U wave normally has the same polarity as the T wave; when the serum potassium level falls to <3.5 mEq/L, the amplitude of the T wave decreases.
• A marked increase in U wave amplitude with potassium <3 mEq/L: the U wave becomes taller than the T wave: with serum potassium <1.5 mEq/L, the T and the U wave may become fused. The changes are seen best in leads V2 through V5 (Figs. 10-7 and 10-8C).

![ECG images showing hypokalemia](image-url)

Fig. 10-7. Hypokalemia produced by diuretics. The serum potassium level was 2.7 mEq/L, sodium was 124 mEq/L, and calcium was 9.2 mg/dL. The ECG shows diffuse ST segment depression, T wave flattening, and prominent U waves. The prominent U waves may be mistaken for T waves. (From Chou TC: Electrocardiography in Clinical Practice, 4th ed., Philadelphia, 1996, WB Saunders, Elsevier Science.)
**Fig. 10-8.** A, ECG signs of hyperkalemia.

- $K^+ > 5.7$ mEq/L: Earliest signs are T wave peaked and narrow base (“tented”); PR interval may be prolonged.
- $K^+ > 7$ mEq/L: P wave flat or absent; QRS widens; prominent S wave.
- $K^+ > 8$ mEq/L: S wave becomes wider and deeper and moves steeply into the T wave; there is virtually no isoelectric ST segment; occasionally ST segment elevation.

**B**, ECG changes in hyperkalemia. On day 1, at a $K^+$ level of 8.6 mEq/L, the P wave is no longer recognizable and the QRS complex is diffusely prolonged. Initial and terminal QRS delay is characteristic of $K^+$-induced intraventricular conduction and is best illustrated in leads V$_2$ and V$_6$. On day 2, at a $K^+$ level of 5.8 mEq/L, the P wave is recognizable with a PR interval of 0.24 second, the duration of the QRS complex is approximately 0.10 second, and the T waves are characteristically tented. **C**, ECG changes in hypokalemia. On day 1, at a $K^+$ level of 1.5 mEq/L, the T and U waves are merged. The U wave is prominent and the QU interval is prolonged. On day 4, at a $K^+$ level of 3.7 mEq/L, the tracing is normal. (A from Khan, M. Gabriel: *Medical Diagnosis and Therapy*, Philadelphia, 1994, Lea & Febiger; B and C from Braunwald E: *Heart Disease: A Textbook of Cardiovascular Medicine*, 5th ed., Philadelphia, 1997, WB Saunders, Elsevier Science.)
• ST segment depression.
• An increase in the QRS duration.
• Slight prolongation of the PR interval.

**HYPERKALEMIA**

**Diagnostic Criteria**

• Mild hyperkalemia: At a serum potassium level of approximately 5.7 to 6.5 mEq/L, the P wave widens; tall, peaked, narrow-based, “tented” T waves appear in many leads; and first-degree atrioventricular (AV) block occurs (see Fig. 10-8A and B).
• Severe hyperkalemia: At a serum potassium level >6.5 mEq/L, the second portion of the QRS complex shows significant widening, which may show notching or slurring, and thus the wide QRS merges with the tall, tented T waves. The ST segment may be elevated (see Fig. 10-8).
• High-degree AV block: P waves disappear.
• Ventricular tachycardia (VT), ventricular fibrillation (VF), or idioventricular rhythm.

**DIGITALIS**

**Digitalis Effect**

• Digitalis effect revealed by the ECG does not imply toxicity and is suggested by the following:
  • Sagging ST segment depression with upward concavity
  • Decreased amplitude of T wave, which may be biphasic
  • Shortening of the QT interval
  • Prolonged PR interval; first-degree AV block
  • Increased amplitude of the U wave

**Digitalis Toxicity**

• Digitalis toxicity is suggested by the occurrence of almost any type of arrhythmia or conduction defect, with the exception of bundle branch block.
• Common arrhythmias include the following:
  • Excitant disturbances such as ventricular premature beats (VPBs), especially bigeminy and multifocal VPBs; atrial tachycardia; AV junctional tachycardia; accelerated junctional rhythm; VT; bidirectional tachycardia; and VF.
  • Suppressant disturbances such as sinus bradycardia, first-degree AV block, second-degree AV block, Mobitz type I (Wenckebach) block, and complete AV block.
• Combined disturbances such as atrial tachycardia with AV block (i.e., paroxysmal atrial tachycardia [PAT] with block) and regular, accelerated junctional rhythm in the presence of atrial fibrillation (Fig. 10-9).

![ECG tracing]

**Fig. 10-9.** Atrial fibrillation with junctional tachycardia (rate, 95 bpm) resulting from digitalis toxicity. Note the absolute regularity of the rhythm. (From Wellens JH, Conover MB: *The ECG in Emergency Decision Making*, Philadelphia, 1992, WB Saunders, Elsevier Science.)
DEXTROCARDIA: TRUE DEXTROCARDIA (WITH SITUS INVERSUS)

Diagnostic Criteria

- In lead I, the P, QRS, and T waves are inverted or upside down (Fig. 10-10).
- Leads aVR and aVL are reversed (aVL is now aVR); thus prominent negative deflections are recorded in aVL with positive deflections in aVR.
- Lead II represents the usual lead III and vice versa.
- Lead aVF is unaffected.
- There is decreasing R wave amplitude from leads V₁ through V₆. V₁ is the equivalent of the usual V₂ and vice versa.

Diagnostic Confirmation

The ECG should be repeated with the right and left arm leads reversed. Placing the V leads in the equivalent positions on the right side of the chest is necessary for accurate interpretation of the ECG.

Diagnostic Pitfalls

- Incorrect arm lead placement: Reversal of the arm leads can produce similar recordings in leads I, aVR, and aVL as observed with true dextrocardia but not in V₁ through V₆. A tip off to this error is the normal R wave progression in V₂ to V₆ and the marked dissimilarity of the record in leads I (negative complex) and V₆ with a normal amplitude R wave.
- Isolated dextrocardia without situs inversus invariably is associated with complicated cardiac malformations and is rarely seen in adults.
- In dextroposition, the heart is displaced to the right by lung disease. The ECG is normal, but R waves are prominent in V₁ through V₃ and decrease in amplitude from V₂ through V₆ (Fig. 10-11).
Fig. 10-10. Mirror-image dextrocardia with situs inversus. The patient is a 15-year-old girl. There is no evidence of organic heart disease. A, Tracing recorded with the conventional electrode placement. B, Tracing obtained with the left and right arm electrodes reversed. The precordial lead electrodes also were relocated in the respective mirror-image positions on the chest. The tracing is within normal limits. (From Chou TC: *Electrocardiography in Clinical Practice*, 4th ed., Philadelphia, 1996, WB Saunders, Elsevier Science.)
Fig. 10-11. Dextroposition. The patient is a 40-year-old woman with hypoplastic right pulmonary artery and right lung, probably of congenital origin. The heart and mediastinum are displaced to the right side of the chest. In the ECG, the limb leads are normal except for a relatively large R wave in lead II and S wave in lead aVR. The precordial leads show tall R waves in leads V1 through V3; the amplitude of the R waves decreases from V2 through V6. (From Chou TC: Electrocardiography in Clinical Practice, 4th ed., Philadelphia, 1996, WB Saunders, Elsevier Science.)

ELECTRICAL ALTERNANS

- Electrical alternans refers to regular alternation in the amplitude, direction, or configuration of the QRS complexes in any or all leads (Figs. 10-12 to 10-14). The RR intervals remain unchanged (regular).
- Total electrical alternans refers to involvement of the P, QRS, and T waves and occasionally the U wave.

Causes

- Alternans of the QRS complex is rare in patients with cardiac tamponade and occurs in some patients with a large pericardial effusion, particularly with malignancy.
- Total electrical alternans is almost diagnostic of cardiac tamponade, although it occurs in fewer than 10% of patients with tamponade and may be associated with a “swinging heart” on echocardiography.
Fig. 10-12. Electrical alternans in postpericardiotomy syndrome. The patient is a 30-year-old man who developed pericarditis and pericardial effusion 3 weeks after aortic valve surgery. In the ECG, in addition to the alternation of the QRS complex, T wave alternans can be seen in lead III. (From Chou TC: *Electrocardiography in Clinical Practice*, 4th ed., Philadelphia, 1996, WB Saunders, Elsevier Science.)

Fig. 10-13. Electrical alternans during supraventricular tachycardia (orthodromic atrioventricular reentrant tachycardia). The patient is a 23-year-old woman with type A Wolff-Parkinson-White syndrome without other evidence of organic heart disease. Alternation of the QRS complex is seen during the tachycardia. (From Chou TC: *Electrocardiography in Clinical Practice*, 4th ed., Philadelphia, 1996, WB Saunders, Elsevier Science.)
Fig. 10-14. Total electrical alternans (P-QRS-T) caused by pericardial effusion with tamponade. This finding, particularly in concert with sinus tachycardia and relatively low voltage, is a highly specific, although not sensitive, marker of cardiac tamponade. (From Braunwald E: *Heart Disease: A Textbook of Cardiovascular Medicine*, 5th ed., Philadelphia, 1997, WB Saunders, Elsevier Science.)
• Severe coronary artery and hypertrophic heart disease is a rare cause of electrical alternans.
• Supraventricular tachycardia with a very rapid ventricular rate, mainly occurring in patients with Wolff-Parkinson-White syndrome (orthodromic) reentrant tachycardia (see Fig. 10-13), is another cause.

**ELECTRONIC PACING**

**Ventricular Pacing**

• The pacemaker impulse, a sharp narrow spike, is followed by a QRS complex of different morphology than the intrinsic QRS. With right ventricular pacing, the QRS complex is similar to that of left bundle branch block (Figs. 10-15 and 10-16).
• With left ventricular epicardial myocardial pacing, the QRS shows a RBBB morphology.
• Pacemakers in a “unipolar pacing mode” cause a larger amplitude spike than that of bipolar pacing.

**Ventricular Demand Pacing (VVI)**

Pacing output is inhibited by sensed ventricular signal (see Fig. 10-20).

**Atrial Pacing**

Pacemaker spike is followed by P wave and narrow paced QRS complexes in response to paced atrial beats (Fig. 10-17).

**Atrial Demand Pacing (AAI)**

Pacing output is inhibited by sensed atrial signal.

**Atrioventricular Sequential Pacing**

• Atrial followed by ventricular pacing (Fig. 10-18).
• Could be pacing in both atrium and ventricle; senses R waves only (DVI pacing mode).
• Pacing in and senses both atrium and ventricle (DDD mode); synchronizes with atrial activity and paces ventricle after preset AV interval (Fig. 10-19).
• Output inhibited by sensed atrial signal (AAI) and by sensed ventricular signal (VVI), but tracking of atrial rate by ventricular sensing does not occur (DDI pacing mode).
• Fixed-rate (asynchronous) atrial and ventricular pacing at specific AV interval (DOO pacing mode).
Fig. 10-15. Electronic pacing; capture rate = 60 bpm.
Fig. 10-16. Electronic pacemaker, ventricular capture; rate = 60 bpm. No further analysis is attempted because of pacemaker rhythm.
Fig. 10-17. Electronic pacemaker, atrial pacing; rate = 70 bpm.
Pacemaker Malfunction

Undersensing Malfunction

For a pacemaker in the inhibited mode, undersensing is diagnosed on ECG by a pacemaker spike at an inappropriately short interval after a spontaneous (intrinsic) event (i.e., a failure of the pacemaker to be inhibited by an appropriate intrinsic atrial or ventricular depolarization [QRS]). Figure 10-20 shows accurate sensing. With sensing malfunction, the pacemaker operates like a fixed-rate pacemaker; the spontaneous intrinsic QRS complexes are not sensed (Fig. 10-21).

Pacing Malfunction

• Not firing: failure of appropriate pacemaker output, which may be caused by failure of the pacemaker impulse to depolarize the ventricle because of inadequate voltage output from the pulse generator, a broken lead wire, or electrode displacement (Fig. 10-22).

Fig. 10-19. Different modes of pacemaker function are shown. A, AOO, fixed rate atrial pacing. Note narrow, paced QRS complexes in response to paced atrial beats. B, VDD, the pacemaker senses the atrium and the ventricle and paces the ventricle. Each spontaneous P wave is followed by a paced ventricular complex. C, DDD, the pacemaker senses and paces in the atrium and the ventricle. The sixth complex of this strip represents a spontaneous P wave that conducts to the ventricle, resulting in a narrow QRS complex with the pacing spike occurring in the ventricular refractory period. Arrows indicate pacing stimulus artifacts. (From Saksena S: In Khan, M. Gabriel: Heart Disease Diagnosis and Therapy, Baltimore, 1996, Williams & Wilkins.)

Fig. 10-18. Electronic pacemaker, ventricular capture; rate = 72 bpm; atrioventricular sequential pacemaker.
Fig. 10-20. Electronic pacemaker, demand mode; ventricular capture rate = 75 bpm. Note the spontaneous beat in V4 is followed by sensing and pacemaker capture at the appropriate interval, which is equal to that shown in lead I. The pacemaker output is inhibited appropriately in response to the intrinsic QRS complex (the first beat in lead V4).

Fig. 10-21. Ventricular demand pacemaker (VVI) with sensing malfunction. The pacemaker operates like a fixed-rate pacemaker. The spontaneous ventricular beats are not sensed. The spontaneous rhythm is atrial fibrillation. (From Chou TC: Electrocardiography in Clinical Practice, 4th ed., Philadelphia, 1996, WB Saunders, Elsevier Science.)
Fig. 10-22. Right and left ventricular pacemakers and pacemaker malfunction. These tracings were obtained from the same patient. A, Tracing recorded when the patient had a transvenous right ventricular demand pacemaker that was functioning properly. Because there are no spontaneous beats, the demand function of the pacemaker cannot be demonstrated. B, Tracing showing intermittent pacing failure. The first QRS complex is a spontaneous beat. The second complex is pacemaker induced. The next pacemaker spike appears prematurely and is not followed by ventricular depolarization. A pseudofusion beat follows. None of the last three pacemaker stimuli captures the ventricle. The sensing function of the pacemaker appears intact. The pacing malfunction was found to be the result of a broken lead. (From Chou TC: Electrocardiography in Clinical Practice, 4th ed., Philadelphia, 1996, WB Saunders, Elsevier Science.) C, Tracing recorded after the patient received an epicardial left ventricular pacemaker. (From Chou TC: Electrocardiography in Clinical Practice, 4th ed., Philadelphia, 1996, WB Saunders, Elsevier Science.)
Fig. 10-23. Battery failure. The battery power failure is indicated by a decrease of the pacing rate from 70 to 47 bpm. (From Chou TC: Electrocardiography in Clinical Practice, 4th ed., Philadelphia, 1996, WB Saunders, Elsevier Science.)

- Battery power failure: indicated by a decrease of the pacing rate (Fig. 10-23).

**PULMONARY EMBOLISM**

The ECG findings are nonspecific; more importantly, the transient occurrence of the following should heighten clinical suspicion of PE:

- Sinus tachycardia
- Symmetrical T wave inversion; strain pattern in leads V₁ through V₃ or V₄
- ST depression in leads I, II, and V₃ through V₆
- S₁Q₃ or S₁Q₃T₃ pattern (Fig. 10-24)
- Incomplete or complete RBBB pattern
- Q waves in leads V₁, III, and aVF but not in lead II
- QR in V₁
- ST segment elevation in leads V₁ through V₂ or V₃, aVR, and III (see Fig. 10-24)
- ST segment depression in V₃ through V₅ or V₆ because of associated myocardial ischemia
Fig. 10-24. Acute massive pulmonary embolus with the characteristic S1Q3 pattern and the more common but nonspecific changes, including incomplete right bundle branch block and ST segment elevation in leads V1 through V3 with terminal T wave inversion. (From Braunwald E: Heart Disease: A Textbook of Cardiovascular Medicine, 5th ed., Philadelphia, 1997, WB Saunders, Elsevier Science.)

Fig. 10-25. Hypothermia. Note marked elevation of the J deflection is maximal in midprecordial leads. (From Marriott JLH: Practical Electrocardiography, 8th ed., Baltimore, 1988, Williams & Wilkins.)
• S₁, S₂, S₃ pattern
• Arrhythmias that include premature beats, atrial flutter or atrial fibrillation, and VF
• Right atrial enlargement
• Right-axis deviation

In the presence of submassive PE, the ECG may show no significant abnormality. With massive PE causing syncope, cardiogenic shock, or acute right-sided heart failure, at least two of the preceding ECG changes usually occur.

HYPOTHERMIA

• All intervals (PR, RR, QRS, and QT) may lengthen.
• Elevated “T waves” (Osborn waves) appear (Fig. 10-25); the start of the ST segment, especially in V₃ and V₄, is elevated, hitched-up, representing distortion of early repolarization.
• Atrial fibrillation occurs often with body temperatures below 32°C.

HYPERCALCEMIA

• Typically causes a shortened QT interval, reflected by and abrupt ascending slope and a gradual downslope of the descending limb of the T wave. This causes a virtual absence of the ST segment.

HYPOCALCEMIA

• Typically causes a prolonged QT interval. The T wave remains normal.
ARRHYTHMIAS

CONTENTS

- ATRIAL PREMATURE BEATS
- JUNCTIONAL OR NODAL PREMATURE BEATS
- VENTRICULAR PREMATURE BEATS
- BRADYARRHYTHMIAS
- NARROW QRS TACHYCARDIAS
- WIDE QRS TACHYCARDIA
- REGULAR WIDE QRS TACHYCARDIA

ATRIAL PREMATURE BEATS

ECG Diagnostic Points

- The morphology of an atrial premature P wave is different from that of the sinus P wave (Fig. 11-1).
- The premature P wave is usually followed by a QRS complex similar to that with the normally conducted sinus beat. The premature P wave may be unrecognizable because it is hidden in the preceding T wave, hence the admonition “search the T for the P” (Figs. 11-2 and 11-3).
- The PR interval of an atrial premature beat (APB) is more than 0.11 second; if the P wave is inverted in leads II, III, and aVF, the PR should be more than 0.11 second to distinguish an APB from a junctional premature beat.
- Early occurring APBs may trigger atrial tachycardia (see Fig. 11-2), atrial flutter, or atrial fibrillation.
- APBs that follow every sinus beat cause atrial bigeminy (see Fig. 11-1).
- The atrial premature P wave may not be conducted, resulting in a pause (see Fig. 11-3).

Nonconducted APBs are the most common cause of pauses. If the premature P waves are not identified, the rhythm may be misinterpreted as sinus bradycardia.
Fig. 11-1. Atrial trigeminy: each sinus beat is followed by a pair of atrial premature beats (APBs): this is true atrial trigeminy. If every third beat is an APB but not a pair of APBs the condition should not be termed atrial trigeminy. Non diagnostic inferior Q waves noted.
**Fig. 11-2.** A. The third and fifth beats are atrial premature beats (APBs). Note that the shorter RP of the second APB is complemented by a much prolonged PR interval. B. Atrial bigeminy in which the PR of the APBs is much prolonged compared with the normal PR of the sinus beats. C. The fourth beat is an APB with right bundle branch block aberration. Note the deformed T wave and the less than compensatory postectopic cycle. D. When the APB is premature enough to make the PP interval (40) less than half the preceding PP interval (88), an atrial tachyarrhythmia is triggered. (From Marriott JLH: *Practical Electrocardiography*, 8th ed., Baltimore, 1988, Williams & Wilkins.)

**Fig. 11-3.** After four sinus beats in (A) and after three sinus beats in (B), a run of nonconducted atrial bigeminy develops. Note in each strip the subtle deformity of the T wave compared with the preceding T waves, resulting from superimposed P’ waves. In (C), the T waves look a little too pointed for natural T waves; however, when no previous T waves are available for comparison, it is impossible to diagnose the atrial bigeminy. (From Marriott JLH: *Practical Electrocardiography*, 8th ed., Baltimore, 1988, Williams & Wilkins.)
• If the APB traverses the atrioventricular (AV) junction at a time when one of the bundle branches is still refractory, aberrant ventricular conduction may occur. The QRS is wide and resembles a ventricular premature beat (VPB) (see Fig. 11-2C). Examination of the preceding T wave may reveal a deformity caused by a P wave stuck on the T wave, as shown in Fig. 11-2C. In addition, a postectopic cycle that is less than compensatory points to atrial ectopy with aberration.

• Multiple APBs may cause an irregularly irregular pulse.

JUNCTIONAL OR NODAL PREMATURE BEATS

ECG Diagnostic Points

• Junctional P waves may activate the atria retrogradely, and the retrograde P wave may precede the QRS complex (Fig. 11-4). Retrograde conduction may not be observed, and the P wave may become lost in the QRS complex (Fig. 11-5). Occasionally, the P wave follows the QRS complex.

• The P wave, when visible, is inverted in leads II, III, aVF, V₁, V₅, and V₆ and is upright in leads I, aVR, and aVL.

• The P wave may precede the QRS complex by less than 0.11 second.

• The terms upper-, mid-, or lower-nodal rhythm have been replaced by the term junctional rhythm.

Fig. 11-4. Junctional premature beats with antecedent P’ waves. In each lead, the first beat is a sinus beat and the second is a junctional premature beat with a short PR interval and typical P wave polarity. (From Marriott JLH: Practical Electrocardiography, 8th ed., Baltimore, 1988, Williams & Wilkins.)
VENTRICULAR PREMATURE BEATS

**ECG Diagnostic Points**

- There is a wide, bizarre, premature QRST complex, with ST segment sloping off in the direction opposite the abnormal QRS complex (Figs. 11-6 to 11-8).
- There have been no preceding premature P waves. Retrograde conduction of ectopic ventricular impulses occurs often. The retrograde P wave is usually hidden in the ventricular complex but occasionally can cause retrograde capture of the atria, and the inverted P wave may be observed following the VPB (see Fig. 11-7).
- A VPB usually is followed by a fully compensatory pause, but this rule is often broken, and pauses may be less than compensatory.
- VPB duration generally is greater than 0.11 second, but occasionally, VPBs can be as short as 0.1 second in duration.
- If in V1 the abnormal-looking QRS shows a left “rabbit ear” larger than the right “rabbit ear” (see Fig. 11-6), a diagnosis of VPB is certain. If the left “rabbit ear” is smaller than the right, no firm conclusion can be made from the morphology alone.
- Figure 11-8 shows ventricular bigeminy.
- A run of two beats is called a couplet; of three consecutive beats, a triplet or a salvo of three (Fig. 11-9); more than three consecutive VPBs is called ventricular tachycardia (VT) (Fig. 11-10).

With multifocal VPBs, the coupling intervals vary; with unifocal VPBs, the coupling intervals are equal. Unifocal VPBs are of little consequence. VPBs that occur early, close to the T wave or R on T, and
Fig. 11-6. A ventricular premature beat: The ST segment slopes in the direction opposite the slope of the abnormal QRS complex; V₁ shows a left “rabbit ear” larger than the right “rabbit ear.”
Fig. 11-7. A, The fourth beat is an interpolated ventricular premature beat (VPB). Note that it lengthens the PR interval of the next sinus beat (from 0.15 to 0.22 second)—evidence of “concealed” (retrograde) conduction into the atrioventricular junction. B, The two ventricular extrasystoles are followed by retrograde (inverted) P waves at normal RP intervals of 0.17 and 0.19 second. C, Both VPBs are followed by retrograde P waves at abnormally prolonged RP intervals (0.28 and 0.22 second). (From Marriott JLH: Practical Electrocardiography, 8th ed., Baltimore, 1988, Williams & Wilkins.)

Fig. 11-8. Ventricular bigeminy.
Fig. 11-9. Ventricular premature beats occur in pairs (couplets) and in salvos of three (triplets).

Fig. 11-10. Short run of nonsustained ventricular tachycardia.
that are multifocal or multiform, occurring as couplets or triplets, may trigger VT (Fig. 11-11).

VPBs occur commonly in normal and abnormal hearts. The word *beat* denotes an electrical and mechanical event and is preferred to the word *contraction*, which implies a mechanical event. This text uses the terms VPBs and APBs, not VPCs and APCs.

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**Fig. 11-11.** Holter monitor showing multifocal ventricular premature beats: couplet, salvos of three, nonsustained ventricular tachycardia. (From Khan, M. Gabriel: *On Call Cardiology*, 3rd ed., Philadelphia, 2006, WB Saunders, Elsevier Science.)
BRADYARRHYTHMIAS

First-Degree Atrioventricular Block

ECG Diagnostic Points

- PR interval is longer than 0.2 second; usually 0.22 to 0.48 second, but can be as long as 0.8 second (Fig. 11-12). Some normal individuals have intervals up to 0.22 second.
- The PR interval should be constant.
- Each P wave should be followed by a QRS complex.

Fig. 11-12. Sinus tachycardia: rate, 118 bpm; PR is prolonged to 0.28 second: first-degree atrioventricular block.
Second-Degree Atrioventricular Block: Mobitz Type I (Wenckebach) Block

**ECG Diagnostic Criteria**

- There is progressive prolongation of the PR interval until the P wave is blocked, the impulse fails to conduct to the ventricles, and the QRS beat is dropped.
- After the dropped QRS beat, the PR interval reverts to near normal; the PR interval that follows the blocked P wave is always short.
- The RR interval containing the nonconducted P wave is shorter than two of the shorter cycles (i.e., shorter than the sum of two PP intervals) (Fig. 11-13).
- Because there is usually a progressive shortening of the RR interval before a P wave is blocked, beats often are grouped in pairs (bigeminy) or trios (trigeminy). This group pattern is a hallmark of Wenckebach but is not necessary for the diagnosis (Fig. 11-14).

Second-Degree Atrioventricular Block: Mobitz Type II Block

**ECG Diagnostic Criteria**

- At least two regular and consecutive atrial impulses are conducted with the same PR interval before the dropped beat (Fig. 11-15).

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Fig. 11-13. Mobitz type I second-degree atrioventricular block without the typical Wenckebach phenomenon. Note the random variation of the PR and RR intervals except that the PR interval that follows the blocked P waves is always short. The QRS complexes that terminate the pauses may be junctional escape beats. (From Chou TC: *Electrocardiography in Clinical Practice*, 4th ed., Philadelphia, 1996, WB Saunders, Elsevier Science.)
Fig. 11-14. Mobitz type I second-degree atrioventricular block without the typical Wenckebach phenomenon. There is no progressive shortening of the RR interval before the block. (From Chou TC: Electrocardiography in Clinical Practice, 4th ed., Philadelphia, 1996, WB Saunders, Elsevier Science.)

Fig. 11-15. A, Mobitz type II second-degree atrioventricular (AV) block. Two consecutive PR intervals are unchanged before the dropped beat. The conducted beats have a normal PR interval and show right bundle branch block. The fourth beat is a right ventricular premature beat. B, High-grade second-degree AV block. Sinus rhythm with 2:1 and 3:1 AV block. (From Marriott JLH: Practical Electrocardiography, 8th ed., Baltimore, 1988, Williams & Wilkins.)
• With type II, high-grade second-degree AV block (Fig. 11-15B), two or more consecutive atrial impulses fail to be conducted because of the block itself. The diagnosis is strengthened if the atrial rate is slow (less than 135 beats/min [bpm]) in the absence of interference by an escaping subsidiary pacemaker that may prevent conduction.
• Intermittent nonconducted P waves are observed, but with no evidence for atrial prematurity.
• The RR interval containing the nonconducted P wave is equal to two PP intervals.
• The PR interval remains constant and is normal or slightly prolonged.
• The ventricular rhythm is irregular because of nonconducted beats.
• If the conduction problem is in the bundle of His, the QRS complex remains narrow, but it will be longer than 0.12 second if the lesion is below the bundle of His.

**Complete (Third-Degree) Atrioventricular Block**

**ECG Diagnostic Criteria**

• P waves are sinus and plentiful with few QRS complexes.
• There is AV dissociation: no relationship between P waves and QRS complexes: complete absence of AV conduction (Figs. 11-16 and 11-17).

**Fig. 11-16.** Congenital complete atrioventricular block. The narrow QRS complexes suggest that the escape pacemaker is junctional in origin. (From Chou TC: *Electrocardiography in Clinical Practice*, 4th ed., Philadelphia, 1996, WB Saunders, Elsevier Science.)
Rapid ECG Interpretation

Note that A-V dissociation may occur in the absence of third-degree A-V block; the clue here is that the ventricular rate is faster than the atrial rate.

- The RR intervals are regular. The QRS complex is narrow if the site of block is in the A-V node with an escape rhythm originating in the A-V junction. The QRS is wide if the escaped rhythm originates from the ventricle or in the A-V junction in the presence of bundle branch block.
- The atrial rate is faster than the ventricular rate.
- The ventricular rate usually is very slow (less than 45 bpm), but with congenital A-V block, rates may be 40 to 60 bpm (see Fig. 11-16).
- With complete A-V block, anterograde conduction never occurs, but in less than 20% of complete A-V blocks, retrograde conduction to the atria occurs.

Note that A-V dissociation may occur in the absence of third-degree A-V block.

**NARROW QRS TACHYCARDIAS**

Tachycardias should be differentiated as narrow QRS or wide QRS, then as regular or irregular (Fig. 11-18).

*Fig. 11-17.* Complete atrioventricular block with idioventricular rhythm. The QRS complexes are abnormally wide and are different from those seen during sinus rhythm. The ventricular rate is 36 bpm. (From Chou TC: *Electrocardiography in Clinical Practice*, 4th ed., Philadelphia, 1996, WB Saunders, Elsevier Science.)
A. Narrow QRS tachycardia

Regular
- Sinus tachycardia
- Atrioventricular nodal reentrant tachycardia (AVNRT)
- Atrial flutter (with fixed AV conduction)
- Atrial tachycardia (paroxysmal and nonparoxysmal)
- WPW syndrome (orthodromic circus movement tachycardia)

Irregular
- Atrial fibrillation
- Atrial flutter (with variable AV conduction)
- Atrial tachycardia (variable AV block or Wenckebach)
- Multifocal atrial tachycardia

B. Wide QRS tachycardia

Regular
- Ventricular tachycardia
- Supraventricular tachycardia (with preexisting or functional bundle branch block)
  - AV NRT
  - WPW syndrome (orthodromic)
  - Sinus tachycardia
  - Atrial tachycardia
  - Atrial flutter with fixed AV conduction
- WPW syndrome (antidromic, preexcited tachycardia)

Irregular
- Atrial fibrillation (with bundle branch block or with WPW syndrome [antidromic])
- Atrial flutter (varying AV conduction, with bundle branch block or WPW syndrome [antidromic])
- Torsades de pointes

Fig. 11-18. Step-by-step method for accurate ECG interpretation. Step 11: Assess arrhythmias: differential diagnosis of narrow QRS tachycardia (A) and wide QRS tachycardia (B). AV, atrioventricular; WPW, Wolff-Parkinson-White. (Modified from Khan, M. Gabriel: Heart Disease Diagnosis and Therapy, New Jersey, 2006, Humana Press.)
A comparison of the entire 12-lead ECG during tachycardia with the ECG in sinus rhythm is most helpful for clarifying the diagnosis. Careful assessment of leads II, III, aVF, V₁, and V₆ should reveal clues to the diagnosis.

**Sinus Tachycardia**

Sinus tachycardia always should be considered if the rate is 100 to 130 bpm because it is the most common cause of narrow QRS tachycardia (Fig. 11-19). With faster rates, the sinus P wave may be hidden in the T waves and mimic supraventricular tachycardia (SVT) or atrial flutter. The sinus P wave can be revealed by carotid massage.

**Atrioventricular Nodal Reentrant Tachycardia**

Atrioventricular nodal reentrant tachycardia (AVNRT) is the most common cause of a paroxysmal, narrow, regular QRS tachycardia. The

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**Fig. 11-19.** Sinus tachycardia; rate, 165 bpm.
ventricles are activated from the anterograde path of the circuit, with activation of the atrium by the retrograde path (Fig. 11-20).

**ECG Diagnostic Points**

- A rapid, regular rhythm, usually 150 to 225 bpm, is present. A rate greater than 230 bpm should prompt the search for Wolff-Parkinson-White (WPW) syndrome.
- QRS is less than 0.12 second.
- In more than 50% of cases, P waves are hidden within the QRS complex and are not visible; the QRS complex is identical to that of a tracing during sinus rhythm (see Fig. 11-20).
- In approximately 45% of cases, P waves appear hidden, but on careful scrutiny they are visible at the end of the QRS in leads II, III, and aVF.

---

**Fig. 11-20.** A representation of the sites of origin and mechanism of paroxysmal supraventricular tachycardia as determined by the position and polarity of the P waves in relation to the QRS complexes. In atrial tachycardia, the P wave precedes the QRS; its polarity in lead III depends on its location. In AV nodal reentry tachycardia, the P wave is buried within the QRS or may distort the end of the QRS; that portion of the QRS is then negative in lead III. In circus movement tachycardia, the P wave follows the QRS. (From Wellens JJH, Conover MB: *The ECG in Emergency Decision Making*, Philadelphia, 1992, WB Saunders, Elsevier Science.)
as they distort the terminal forces of the QRS complex, resulting in pseudo–S waves in leads II, III, and aVF (see Figs. 11-20 and 11-21). The distortion causes a pseudo r′ wave in lead V1 that mimics RSr′ or incomplete right bundle branch block (RBBB) (see Figs. 11-20, 11-21B, and 11-22).

**Fig. 11-21.** A, The limb leads of a patient with supraventricular tachycardia: rate, 184 bpm. Note the distortion of the terminal QRS in lead III, a pseudo–S wave: typical features of the common form of atrioventricular nodal reentrant tachycardia (AVNRT). B, The V1 lead in the same patient. Note the distortion of the terminal QRS resulting in a pseudo–r′ wave: typical feature of the common form of AVNRT.
Fig. 11-22. Atrioventricular nodal reentrant tachycardia (AVNRT): rate, 140 bpm. Note pseudo–R′ wave in V1, typical of the common type of AVNRT.

- In fewer than 5% of cases, P waves are discernible at the beginning of the QRS and cause pseudo–Q waves in leads II, III, and aVF.
- In a rare form of AVNRT, P waves are negative in leads II, III, and aVF but follow the QRS after a prolonged duration such that the RP interval is greater than the PR interval. It is impossible to distinguish this rare form of AVNRT from the rare type of WPW circus movement tachycardia using the retrograde, slow accessory pathway to activate the atria (see subsequent section on Wolff-Parkinson-White syndrome).
Paroxysmal Atrial Tachycardia with or without Atrioventricular Block

ECG Diagnostic Points

- The P wave precedes the QRS, and its contour is different from that of the sinus P wave. P waves are often small, are not easily identified, and may be hidden in the T wave or QRS; the arrhythmia may be mistaken for sinus tachycardia or AV junctional tachycardia. The PR interval is normal or prolonged.
- The morphology of the P waves depends on the location of the ectopic atrial pacemaker (Fig. 11-23). The RR intervals are equal except for a warm-up period in the automatic type.

![ECG Diagram](image)

**Fig. 11-23.** Atrial tachycardia. The P waves are barely discernible in lead I and are inverted in II, III, and aVF. There is 2:1 atrioventricular block. The atrial rate is 264 bpm; ventricular rate is 132 bpm. Note the isoelectric baseline between the P wave and the QRS complex.
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• The atrial rate may range from 110 to 260 bpm. If the atrial rate is not rapid and AV conduction is not depressed, each P wave may conduct to the ventricle. With digitalis excess, AV conduction may be delayed, resulting in paroxysmal atrial tachycardia with block, but digitalis toxicity is not the only cause of this arrhythmia (Fig. 11-24).

• Variable AV conduction occurs: a 2:1 conduction is common; a 3:1 conduction or Wenckebach phenomenon may occur, causing an irregular rhythm (Fig. 11-25). At times, the varying AV block may result in an irregular ventricular rhythm that may be mistaken for atrial fibrillation.

• An isoelectric baseline exists between the P wave and the QRS complex (see Figs. 11-23 to 11-25).

• The differentiation of atrial tachycardia and atrial flutter may be difficult if the atrial rate is rapid; carotid sinus massage or adenosine brings out the flutter waves if atrial flutter is present.

• Atrial tachycardia persists despite the development of AV block, and this feature excludes WPW syndrome.

**Persistent (Incessant) Atrial Tachycardia**

The incessant nature of atrial tachycardia, a rare tachycardia, may cause dilated (congestive) cardiomyopathy.

**ECG Diagnostic Points**

- The rhythm is regular.
- The P wave precedes the QRS complex. The P wave polarity depends on the site of origin in the atrium.

**Lead II**

![ECG Waveform](image-url)

**Continuous**

![ECG Waveform](image-url)

**Fig. 11-24.** Paroxysmal atrial tachycardia with block. There is 3:1 atrioventricular conduction. The patient was not receiving digitalis. (From Chou TC: Electrocardiography in Clinical Practice, 4th ed., Philadelphia, 1996, WB Saunders, Elsevier Science.)
There is variable AV conduction of 1:1 and 2:1, including Wenckebach phenomenon.
Carotid sinus massage or adenosine increases AV block and facilitates the diagnosis.

**Multifocal Atrial Tachycardia (Chaotic Atrial Tachycardia)**

**ECG Diagnostic Points**

- Atrial rate is 100 to 140 bpm.
- There are frequent multifocal premature beats, at least three different P wave morphologies with changing PR intervals in one lead (Fig. 11-26), and isoelectric baseline between P waves.
- The rhythm is completely irregular; PR, RR, and RP intervals are variable.
- One dominant atrial pacemaker, such as sinus rhythm, is absent, and multifocal APBs are present.
• Causes of multifocal atrial tachycardia include chronic obstructive pulmonary disease, theophylline, and digitalis (rarely).

**Wolff-Parkinson-White Syndrome**

**ECG Diagnostic Criteria**

• The QRS complex duration is ≥0.11 second; in approximately 20% of individuals, the QRS complex may not be >0.1 second. PR is <0.12 second.
• A delta wave is prominent, often in V₁ through V₆, and is a subtle finding in some leads (Fig. 11-27).
• In type A WPW syndrome, a tall R wave present in V₁ and V₂ can mimic right ventricular hypertrophy, RBBB, or posterior infarction (see Fig. 11-27 and Table 2-3).
In type B WPW pattern, the QRS complex is predominately negative in V₁ through V₃ and upright in V₅ and V₆. The pattern may resemble left bundle branch block (Fig. 11-28).

Pseudo–Q waves in inferior leads may mimic inferior myocardial infarction (see Figs. 11-27 and 11-29).

Clues During Tachycardia

- There is narrow QRS complex tachycardia, with regular rhythm.
- P waves follow the QRS at a distance; shape depends on the location of the accessory pathway. With a left lateral accessory pathway, the P wave is negative in lead I. If the location is posteroseptal, P waves are negative in leads II, III, and aVF and positive in aVR and aVL.
- In the common orthodromic circus movement tachycardia, the RP interval is shorter than the PR interval because of retrograde use of the
Fig. 11-27. Continued

Fig. 11-28. Type B Wolff-Parkinson-White pattern in a healthy 35-year-old man. The tracing resembles closely that of complete left bundle branch block. (From Chou TC: Electrocardiography in Clinical Practice, 4th ed., Philadelphia, 1996, WB Saunders, Elsevier Science.)
fast accessory pathway to activate the atria (see Figs. 11-20 and 11-30).

- In a rare form of orthodromic circus movement tachycardia with retrograde activation of the atria through a slowly conducting accessory pathway, the P wave occurs late retrogradely. Thus, the RP interval is longer than or equal to the PR interval and the P wave is negative in leads II, III, aVF, and V₄ through V₆. This arrhythmia pattern is similar to the rare form of AVNRT described previously. This rare type of WPW syndrome may manifest as a persistent (incessant) orthodromic circus movement tachycardia and cause a dilated cardiomyopathy with congestive heart failure (Fig. 11-31).
• Electrical alternans sometimes occurs during orthodromic circus movement tachycardia but rarely occurs with other narrow QRS tachycardias (see Fig. 10-13).

Summary Differential Diagnosis of Narrow QRS Regular Tachycardia

• If AV block is present or can be produced by carotid sinus massage or adenosine, WPW syndrome can be ruled out; atrial flutter and atrial tachycardia persist despite AV block.
• If the P wave is hidden within the QRS or is distorting the terminal QRS, causing a pseudo-S in leads II, III, and aVF or a pseudo r’ in V1, the diagnosis is the common form of AVNRT (see Figs. 11-20 to 11-22).
A negative P wave in lead I suggests WPW syndrome or left atrial tachycardia. A P wave following the QRS complex distinctly with an RP interval shorter than the PR interval is diagnostic of the most common type of WPW syndrome, orthodromic circus movement tachycardia (see Fig. 11-30). An RP interval that is greater than the PR interval indicates the rare WPW orthodromic type, the rare type of AVNRT, or atrial tachycardia (see Fig. 11-31).
• Positive P waves in leads II, III, and aVF with atrial tachycardia rule out AVNRT or WPW syndrome tachycardia.
• P waves negative in leads II, III, and aVF suggest AVNRT or WPW syndrome.
• A ventricular rate greater than 220 bpm with QRS alternans usually indicates WPW syndrome (see Fig. 10-13).
• A ventricular rate greater than 250 bpm with RR intervals less than 240 milliseconds (six small squares) suggests WPW syndrome.

**FOUR TYPES OF Wolff-Parkinson-White Syndrome Tachycardia**

1. Orthodromic circus movement tachycardia: This is the most common tachycardia. The ventricles are activated via the AV node and bundle of His, and the impulse retrogradely uses the fast accessory tract to activate the atria; thus the P wave is close to the preceding QRS complex, and the RP interval is shorter than the PR interval.
2. Rare orthodromic tachycardia: The activation of the ventricles via the AV node and His bundle is similar to that in circus movement tachycardia, but the impulse returns to the atria via the slow accessory tract. Therefore, the P wave follows the QRS at a distance, making the RP interval greater than the PR interval. This arrhythmia mimics the rare form of AVNRT.
3. Rare antidromic tachycardia: The ventricle is activated by anterograde (preexcited tachycardia) use of the bypass tract, causing tachycardia similar to VT, atrial flutter, or atrial fibrillation with a wide QRS complex (see discussion of wide QRS tachycardia).
4. Rare antidromic anterograde conduction: The ventricle is activated by two or more accessory pathways, resulting in wide QRS tachycardia.

**Diagnosis Based on Carotid Sinus Massage or Intravenous Adenosine**

• AVNRT converts to sinus rhythm or no effect.
• Circus movement tachycardia reverts to sinus rhythm or no effect.
• Persistent atrial tachycardia: Increased AV block facilitates recognition of the atrial origin, temporary slowing of heart rate with AV block, or no effect.
• Atrial flutter: Temporary slowing of the ventricular rate reveals flutter waves if not previously visible, or no effect.

*Atrial Flutter*

• A sawtooth pattern is seen in leads II, III, and aVF (Figs. 11-32 and 11-33). The downward deflection of the F waves has a gradual slope
Fig. 11-32. Atrial flutter. (From Khan, M. Gabriel: On Call Cardiology, 3rd ed., Philadelphia, 2006, WB Saunders, Elsevier Science.)
followed by an abrupt upward deflection. This results in the typical sharp spikes of the sawtooth pattern: There are positive, “spiky” P-like waves in lead V1 and negative P-like waves in leads V5 and V6. There is almost no atrial activity in lead I, and leads V5 and V6 often show negligible atrial activity (see Fig. 11-32).

- A ventricular response of 150 bpm is typical of atrial flutter. The atrial rate is often 300 bpm. With 2:1 AV conduction, the ventricular response is 150 bpm. This 2:1 ratio may not be apparent because an F wave may be partially obscured by the QRS complex and the second F wave is hidden in the T wave (Fig. 11-34). This pattern mimics sinus tachycardia or reentrant junctional tachycardia. Carotid sinus massage should reveal sinus P waves or F waves. Conduction ratios of 2:1 and 4:1 may occur. The ventricular rate may vary from 100 to 230 bpm. A ventricular response of greater than 250 bpm suggests WPW syndrome.
Fig. 11-34. Atrial flutter: atrial rate, 270 bpm; ventricular rate, 135 bpm. Note the downward deflection of F waves in leads II, III, and aVF has a gradual slope followed by an abrupt upward deflection. This causes the sawtooth pattern. Alternate F waves coincide with the QRS complex, and the diagnosis may be missed.
• The rhythm is regular but becomes irregular when there is variable AV conduction.
• The atrial rate varies from 250 to 400 bpm but can be less than 200 bpm in patients taking quinidine.

**Atrial Fibrillation**

**ECG Diagnostic Points**

• RR intervals are completely irregular (irregularly irregular) (Figs. 11-35 and 11-36).
• Irregular undulations of the baseline are usually most prominent in V₁; these may be gross (Fig. 11-37) or barely perceptible, described as coarse and fine fibrillation, respectively. Occasionally, there may be no recognizable undulations of the baseline, and careful measurement of the RR interval is necessary to detect slight irregularities.
• The atrial rate ranges from 400 to 700 bpm; there is variable AV conduction, resulting in a chaotic ventricular response.
• QRS complexes often vary in amplitude.

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Fig. 11-35. A, Atrial fibrillation with a ventricular response of 156 bpm. B, [See legend on opposite page]. B, Atrial fibrillation with a rapid ventricular rate of 175 bpm. C, Atrial fibrillation, same patient as in (B); controlled ventricular rate of 108 bpm.

*(continued)*
Fig. 11-35. Continued
Fig. 11-36. Atrial fibrillation with slow ventricular response; rate, 70 bpm. Patient is 80 years old and is not taking digoxin or a β-blocker. Rates <70 bpm commonly seen in older adults who are not taking cardiac medications should raise suspicion of sick sinus syndrome.
Fig. 11-37. Atrial fibrillation with a ventricular response rate of 104 bpm. Note the coarse atrial fibrillation in V1.

- The heart rate is commonly 100 to 180 bpm but can accelerate to more than 200 bpm. Rates of >240 bpm with the QRS complex ≥0.10 second should suggest WPW syndrome. With WPW antidromic tachycardia, wide QRS tachycardia with rates of 250 to 320 bpm may occur.

Figure 11-38 gives clues that assist with the diagnosis of supraventricular arrhythmias.
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AVNRT

Pseudo r' wave V1 in ~ 45% cases
pseudo S wave 11,111, or aVF ~ 45%
P wave hidden in QRS in ~ 50%
P waves before QRS forming
pseudo q waves in 11, 111, or aVF < 5%#

AVRT [ WPW ] *

Discrete P waves follow the QRS
at a distance: RP < PR **
P negative in I = lateral AP;
P negative in 11111, aVF, positive in
aVR, aVL = posteroseptal AP
QRS alternans: a clue to circus movement tachycardia rarely occurs with AVNRT

Atrial tachycardia

P waves precede the QRS; if stuck on the T wave, mimics sinus tachycardia.
Polarity depends on site of origin.
Variable AV conduction 1:1, 2:1, Wenckebach.
Carotid sinus massage increases the block, reveals atrial origin

Atrial flutter

Regular rhythm if fixed AV conduction; rate 150/min, highly suspect flutter.
Prominent flutter waves in lead II, III, aVF, often absent in V5, V6. Spiky P-like waves in V1

Atrial fibrillation

Completely irregular rhythm; absent P waves.
Coarse or fine undulations in V1
The most common significant arrhythmia

* common circus movement tachycardia uses the fast accessory pathway [AP]; rare form uses the slow AP and the RP is > PR.
** May occur with atrial tachycardia
# rare form: negative Ps follow the QRS with RP > PR

Fig. 11-38. Supraventricular arrhythmias: Key Diagnostic Clues.
WIDE QRS TACHYCARDIA

ECG Diagnostic Steps

- Define the QRS duration as ≥0.12 second.
- Define the tachycardia as regular or irregular (see Fig. 11-18B).

REGULAR WIDE QRS TACHYCARDIA

Regular wide QRS tachycardia includes the following:

- Ventricular tachycardia: Consider all wide QRS regular tachycardias as VT until proven otherwise.
- SVT with preexisting or functional bundle branch block: These tachycardias include AVNRT, orthodromic circus movement tachycardia (WPW), atrial tachycardia, and atrial flutter with fixed AV conduction.
- Antidromic circus movement tachycardia or preexcited tachycardia (WPW) is usually exhibited by a very rapid rate >250 beats/min and may be irregular due to atrial fibrillation.

Ventricular Tachycardia

The ECG diagnosis of VT requires the assessment of all 12 ECG leads. The precordial leads are more diagnostic than lead 11 or other limb leads.

The diagnosis of VT can be confidently made by careful scrutiny of the morphologic pattern of the QRS complexes in V_1 through V_6.

- If the QRS complexes are all negative in V_1 through V_6 (i.e., negative precordial concordance), the diagnosis of VT is certain (see Figs. 11-39, 11-40, and 11-41). Negative precordial concordance excludes WPW regular wide complex tachycardia during anterograde conduction over an accessory pathway.
- Finding of predominately negative QRS complexes in V_4 through V_6 is diagnostic of VT (see Fig. 11-39).
- The presence of a QR complex in one or more of precordial leads V_2 through V_6 is diagnostic of VT (Figs. 11-39 and 11-40).
- Note that negative precordial concordance is diagnostic of VT but that positive concordance (all complexes positive in V_1 through V_6) can result from VT or circus movement antidromic tachycardia WPW syndrome.

Findings in V_6 are most useful clues:

- A QS or rS in V_6 (net negative complex) (see Figs. 11-39 and 11-40).
Negative concordance precordial leads # = VT
Predominantly negative QRS V4–V6 # = VT
q R in one or more of V2–V6 = VT

V6: q R = VT
QS = VT
r S [predominantly – ve QRS] = HP

V1: Steeper R upstroke than downstroke
[taller left” rabbit ear”] = VT
q R = HP
broad, fat r wave > 0.03 sec = HP

QRS very wide > 0.14 * not diagnostic = HP
AV dissociation** not diagnostic = HP
Fusion beats *** not diagnostic : = HP

Fig. 11-39. Ventricular tachycardia: Key Diagnostic Clues.
Rapid ECG Interpretation

<table>
<thead>
<tr>
<th>Predominantly -ve QRS $V_4$ to $V_6$</th>
<th>QR complex in one or more of $V_2$ to $V_6$</th>
<th>Morphology $V_1$ and $V_6$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$V_1^{**}$</td>
<td></td>
<td>Left “rabbit ear”</td>
</tr>
<tr>
<td>$V_2$</td>
<td></td>
<td>or net negative in $V_6$</td>
</tr>
<tr>
<td>$V_3$</td>
<td></td>
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<tr>
<td>$V_4$</td>
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<td>$V_5$</td>
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<td></td>
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<tr>
<td>$V_6$</td>
<td></td>
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</tr>
</tbody>
</table>

* = or concordant negativity in leads $V_1$ through $V_4$. Positive concordance in leads $V_1$ through $V_6$ can be caused by VT or Wolff-Parkinson-White antidromic (preexcited) tachycardia.

** = it is necessary to study the entire 12-lead tracing with particular emphasis on leads $V_1$ through $V_6$; lead II may be useful for assessment of P waves and AV dissociation.

**Fig. 11-40.** Electrocardiographic hallmarks of ventricular tachycardia (VT). (From Khan, M. Gabriel: *On Call Cardiology*, 3rd ed., Philadelphia, 2006, WB Saunders, Elsevier Science.)
Fig. 11-41. Onset of a tachycardia with negative precordial concordance. Negative precordial concordance indicates ventricular tachycardia, because such a pattern does not occur during anterograde conduction over an accessory pathway. (From Wellens J.H., Conover M.B.: The ECG in Emergency Decision Making, Philadelphia, 1992, WB Saunders, Elsevier Science.)
Findings in V₁ are useful clues:

- A wide small r wave >0.03 second (see Fig. 11-39).
- An RS interval longer than 0.1 second, measured from the R wave to the nadir of the S wave in any precordial lead.
- A steeper R wave upstroke than downstroke in V₁ (taller left “rabbit ear”). The morphology in V₁ may be helpful: If the left “rabbit ear” is taller than the right in lead V₁, VT is the most likely diagnosis (Figs. 11-39 and 11-42). Note that the rabbit ear may be subtle. Figure 11-42 shows other characteristic features in V₁, V₂ and V₆.

Other helpful features include:

- AV dissociation: The presence of more QRS complexes than P waves supports the diagnosis of VT, but P wave identification may be difficult.
The terminal portion of the T wave or initial parts of the QRS may resemble P waves, leading to an incorrect diagnosis of SVT. In addition, in some cases of VT, 1:1 ventricular/atrial conduction may be observed because retrograde impulse conduction to the atria from the ventricular focus often occurs. AV dissociation is not a reliable diagnostic point and is observed in <45% of VTs.

- With VT, the axis is commonly −90 degrees to ±180 degrees. However, the axis may be normal in patients with idiopathic VT and other varieties of VT.
- Positive concordance: A positive QRS complex in V₁ through V₆ is suggestive of VT, but this pattern can be seen with WPW syndrome (Fig. 11-43). Negative precordial concordance is diagnostic of VT because
this pattern does not occur during antidromic circus movement tachycardia (WPW syndrome) in which conduction is anterograde over the bypass tract.

**IRREGULAR WIDE QRS TACHYCARDIA**

Irregular wide QRS tachycardias include the following:

- Torsades de pointes.
- Atrial fibrillation with bundle branch block or with the antidromic variety of WPW, anterograde conduction over the bypass tract (Fig. 11-44).
- Atrial flutter with varying AV conduction and bundle branch block or atrial flutter and varying AV conduction in the WPW syndrome with anterograde conduction (antidromic) over the bypass tract (Fig. 11-45).

![Fig. 11-44. Atrial fibrillation with wide QRS tachycardia in a patient with Wolff-Parkinson-White syndrome: antidromic tachycardia.](image-url)
Fig. 11-45. A 12-lead ECG from a patient with antidromic circus movement tachycardia. (From Wellens JJH, Conover MB: *The ECG in Emergency Decision Making*, Philadelphia, 1992, WB Saunders, Elsevier Science.)

**Torsades de Pointes**

- Torsades is a polymorphic VT that usually occurs in the presence of a prolonged QT interval.
- The RR interval is irregular; the QRS complexes show a typical twisting of the points.
- The amplitudes of the complexes vary and appear alternately above and below the baseline (Fig. 11-46).
- The ventricular rate varies from 200 to 300 bpm, but can reach 400 bpm, and is usually not sustained (lasting 30 seconds to 1 minute).
- Longer episodes degenerate into ventricular fibrillation.
- Drugs and conditions that may precipitate torsades include the following:
Fig. 11-46. Torsades de pointes. A, Continuous recording monitor lead. A demand ventricular pacemaker (VVI) has been implanted because of Mobitz type II second-degree AV block. After treatment with amiodarone for recurrent ventricular tachycardia (VT), the QT interval became prolonged (approximately 640 milliseconds during paced beats), and the patient developed episodes of torsades de pointes. In this recording, the tachycardia spontaneously terminates, and a paced ventricular rhythm is restored. Motion artifact is noted at the end of the recording as the patient lost consciousness. B, Tracing from a young boy with a congenital long QT syndrome. The QTU interval in the sinus beats is at least 600 milliseconds. Note TU wave alternans in the first and second complexes. A late premature complex occurring in the downslope of the TU wave initiates an episode of VT. (From Braunwald E: Heart Disease: A Textbook of Cardiovascular Medicine, 5th ed., Philadelphia, 1997, WB Saunders, Elsevier Science.)
• Antiarrhythmics known to increase the QT interval (e.g., quinidine, procainamide, amiodarone, disopyramide, sotalol)
• Tricyclic antidepressants and phenothiazines
• Histamine (H₁) antagonists (e.g., astemizole, terfenadine)
• Antiviral and antifungal agents and antibiotics
• Hypokalemia, hypomagnesemia
• Insecticide poisoning
• Bradyarrhythmias
• Congenital long QT syndrome
• Subarachnoid hemorrhage
• Chloroquine, pentamidine
• Cocaine abuse
ECG Board Self-Assessment Quiz

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ECG Board Self-Assessment Quiz
Answers to ECG Board Self-Assessment Quiz

(Note: See pages 391–399 for answers.)
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Rapid ECG Interpretation

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Fig. 12-87A.
Rapid ECG Interpretation

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Fig. 12-89.
Fig. 12-90.
ANSWERS TO ECG SELF-ASSESSMENT QUIZ

**Fig. 12-1.** Acute pericarditis: sinus tachycardia 126/min; typical features: widespread ST elevation and PR segment elevation in aVR, which shows ST segment depression. Note: The J-point level almost equals the height of the T wave in V6.

Causes of ST segment elevation include:

- Normal variant
- Acute ST segment elevation MI (STEMI)
- Coronary artery spasm: Prinzmetal angina
- Left ventricular aneurysm
- Acute pericarditis
- Left ventricular hypertrophy
- Left bundle branch block
- Acute myocarditis
- Hyperkalemia
- Brugada syndrome; ST elevation V1–V3

**Fig. 12-2.** Atrial fibrillation with rapid ventricular response 152 beats/min; marked ST segment depression in V2 to V6, in keeping with subendocardial ischemia, probable non–ST segment elevation MI.

**Fig. 12-3.** Acute extensive anterior infarct. Marked ST elevation and pathologic Q-waves in V1 to V6. Inferior MI, age indeterminate, sinus tachycardia 115/min.

**Fig. 12-4.** WPW syndrome: prominent Delta wave, short PR interval, tall R wave in V2. Note the features in II, III, and aVF mimic inferior MI, and the pattern in V1 mimics IRBBB.

- Thus, the assessment for WPW syndrome is done early in the interpretive sequence (Step 3) soon after the assessment for bundle branch block.

**Fig. 12-5.** Hyperkalemia; note the tall tented T waves in V2–V4; serum potassium 5.9 mmol/L.

**Fig. 12-6.** Left ventricular hypertrophy; note also, left atrial hypertrophy. The ST segment depression in lead V3 suggests the presence of underlying ischemia.

**Fig. 12-7.** Supraventricular tachycardia; rate 230/min; orthodromic circus movement tachycardia. Patient with WPW syndrome.

**Fig. 12-8.** A, Note tall R wave in V1–V2, small R and deep S in V6 interpreted by the computer as right ventricular hypertrophy. Findings are caused by incorrect placement of the precordial leads; this is a rare error for technicians. B, Normal ECG; same patient as in (A), correct placement of V leads.
**Fig. 12-9.** A 2:1 AV block, probably Mobitz type I in view of a normal narrow QRS; approximately 30% of Mobitz II exhibits a narrow QRS complex. Thus type II block cannot be excluded.

The diagnosis of Mobitz type II block is certain when at least two regular and consecutive atrial impulses (P waves) are conducted with a constant PR interval before the occurrence of the dropped beat. Two or more consecutive PR intervals are unchanged before the dropped QRS beat. Note that Wenckebach did not have the use of electrocardiography when he cleverly deduced two forms of blocks from studies of the jugular venous waves in 1906. Mobitz in 1924 using the ECG described type I and type II blocks. Mobitz type I includes Wenckebach phenomenon (with each successive beat, the PR interval gradually lengthens and a beat is dropped). Importantly, not all Mobitz type I AV block reveal Wenckebach phenomenon (see Figs. 11-13, 11-14, and 11-15 for Mobitz type II block).

**Fig. 12-10.** Supraventricular tachycardia (SVT). Rate 155/min. Note the diagnostic pseudo–r’ wave in V1, and the small distortion of the terminal QRS complex (pseudo–S wave) in leads II, III, aVF. These are typical features of AV nodal reentrant tachycardia (AVNRT) observed in ~45% of SVTs.

**Fig. 12-11.** Atrial fibrillation and ventricular premature beats (VPBs); note the couplets; multiform VPBs. Also left axis. Abnormal ECG.

**Fig. 12-12.** ST segment elevation V1, V2, V3; normal variant in a 46-year-old man. Note the fishhook pattern in V2.

**Fig. 12-13.** Anterior MI, probably in recent past: age indeterminate. Consider LV aneurysm if ST elevation V2–V4 has persisted beyond 6 months.

**Fig. 12-14.** Female age 68 with mild hypertrophic cardiomyopathy confirmed by echocardiography at age 45. Note the significant pathologic Q waves in V4–V6, and leads II, III, aVF. Incorrectly interpreted by computer as old inferolateral MI.

**Fig. 12-15.** LBBB.

**Fig. 12-16.** Wide QRS tachycardia, irregular rhythm: Atrial fibrillation with a ventricular response, 160 beats/min in a patient with Wolff-Parkinson-White syndrome. Clues to WPW syndrome antidromic, preexcited tachycardia: irregular, wide-complex tachycardia, often with runs at rapid rates exceeding 250 beats/min.

**Fig. 12-17.** Mimics inferior MI. The ST-T abnormality in lead 1 cannot be explained by an inferior MI. Fishhook-type pattern in II and concave ST segment elevation in III, aVF, are unlike acute MI. Treadmill cardiac nuclear perfusion imaging is normal; echocardiogram
shows mild LVH in this 31-year-old male known to have a VSD patch. Fig. 12-18. LVH, left atrial hypertrophy. Note the shape of the ST segment and deep T wave inversion V₄–V₆ and the extension of these changes to V₃ are in keeping myocardial ischemia and not simply hypertrophy. Also, consider apical HCM.

Fig. 12-19. Mimics dextrocardia. Error by technician: The reversed placement of arm leads is a common error; the reversal of V leads is a rare error and can cause incorrect interpretation of the ECG. Computer interpretation, dextrocardia. In this tracing, V₁ = V₆.

Fig. 12-20. Accelerated AV conduction, early transition: normal ECG.

Fig. 12-21. Tracing interpreted by computer as LBBB. Note absence of pacing spikes, because the muscle filter is activated. Deactivation of the computer muscle filter should expose the pacing spikes. (see Fig. 12-22.) The atypical IVCD, the negative concordance V₃–V₆ and in II, III, and aVF are pacing clues.

Fig. 12-22. Electronic pacing; the same patient as in Fig. 12-21 but with muscle filter turned off. Capture rate 61/min.

Fig. 12-23. WPW syndrome (type A). Note the tall R waves V₁–V₃.

Causes of tall R wave in V₁ include:

- Normal variants, thin chest wall, detroposition
- Misplacement of chest leads: V₆ placed in V₁ position; fortunately a rare technician’s error.
- WPW syndrome: type A pattern caused by lateral or posterior accessory pathways.
- RBBB
- Posterior MI; inferoposterior MI
- Right ventricular hypertrophy
- Hypertrophic cardiomyopathy
- Duchenne muscular dystrophy

Fig. 12-24. Normal ECG; tall peaked precordial T waves in a healthy 35-year-old with normal serum potassium.

Fig. 12-25. Severe myocardial ischemia; probable non–ST elevation MI?

Causes of ST depression include:

- Acute non–ST segment elevation MI (non–Q wave MI)
- Acute myocardial ischemia without infarction (subendocardial ischemia)
- Chronic myocardial ischemia
- Reciprocal ST depression associated with STEMI
- LVH with “strain” pattern
- Conduction defects: LBBB, RBBB, IVCD, WPW
- Digoxin and other drugs
- Hypokalemia
- Cardiomyopathy

**Fig. 12-26.** Normal ECG from a healthy 7 year old. ST-T changes V₁ to V₃ are normal findings.

**Fig. 12-27.** Hypertrophic cardiomyopathy. An asymptomatic 20 year old; played soccer, rugby, and hockey for the past 5 years. Routine physical revealed a grade II systolic murmur. Echocardiogram: asymmetric LVH; marked septal hypertrophy: septal thickness 36 mm (normal <11 mm).

**Fig. 12-28.** Right bundle branch block, left axis −60, left anterior fascicular block (hemiblock).

**Fig. 12-29.** Normal variant ST elevation V₂–V₅ in a 24-year-old male. Note the J-point fishhook in V₃. Normal variant ST elevation is common in males and rare in females. Many cardiologists and interpreters refer to the ST change as “early repolarization.”

**Fig. 12-30.** Old inferior MI: Deep wide Q wave changes in II, III, aVF have persisted for more than 15 years. Also, features of LVH and anterolateral ischemia are present.

**Fig. 12-31.** Atrial flutter. Note: a ventricular rate of 150/min is a clue to atrial flutter with 2:1 AV conduction; the prominent sawtooth pattern in II, III, aVF is typically absent in I, V₅, V₆.

**Fig. 12-32.** Brugada syndrome. Note the atypical incomplete RBBB pattern with a curious coved ST segment elevation in V₁, V₂, and saddle-back type elevation in V₃. A 40-year-old Algerian male who collapsed; following an episode of syncope; an ICD was placed. Brugada and WPW syndrome should be considered along with assessments of blocks and is thus put as Step 3 of the 11-step strategies. Readers may wonder why interpreters should be on the watch for these rare conditions. They can cause death in young individuals, and these deaths and/or hospitalizations can be prevented.

**Fig. 12-33.** Nonsustained ventricular tachycardia in a 38-year-old male presenting with chest pain at 3:14:41 AM.

**Fig. 12-34.** Ventricular premature beats, triplets. ECG at 3:13:29 AM on presentation to the ER. Same patient as in Fig. 12-33; ECG taken a minute later.
Fig. 12-35. Acute inferior myocardial infarction. Note the reciprocal depression in leads I and aVL. Right bundle branch block.

Fig. 12-36. Extensive anterior infarct probably in recent past; age indeterminate; left anterior fascicular block (hemiblock).

Fig. 12-37. WPW mimics inferior MI.

Fig. 12-38. Atrial fibrillation, normal ventricular rate.

Fig. 12-39. Severe myocardial ischemia. A 52-year-old female with angiographic proven severe obstructed coronary artery disease. Current ECG similar to 4 years prior and unchanged over 6 years. Received PTCA and stent at age 48.

Fig. 12-40. Electronic pacing; capture rate 66/min. Note the premature beat in V₁ is followed by a correctly timed paced QRS and indicates that the pacemaker is sensing correctly. The paced beat after the premature beat occurs at the correct pacing interval equal to the distance between the pacing spikes.

Fig. 12-41. RBBB and Q waves II, III, aVF: probable old inferior MI.

Fig. 12-42. A 41-year-old African male with long-standing restrictive cardiomyopathy. T wave changes caused by myocardial disease mimic LVH and ischemia. Borderline IVCD.

Fig. 12-43. Old anterior MI. Left atrial abnormality; APB, left axis, left anterior fascicular block (hemiblock).

Fig. 12-44. VPBs, bigeminy.

Fig. 12-45. A 2:1 AV block. Note the P-P intervals are constant. Computer interpreted as nonconducted APBs. ECG from a 44-year-old female with some shortness of breath, no presyncope. ECG tracing November 30, 2005. Note the heart rate, 43/min, is identical in a tracing done 1 year later, shown in Fig. 12-9. If the heart rate is <45/min, screen for bradycardias.

The differential diagnosis for marked bradycardia, slow rate of <45/min include:

- Sinus bradycardia
- Nonconducted APBs (bigeminy)
- Sinoatrial block (SA block)
- A variety of AV block (2:1 AV block, 3:1 block, complete AV block, and atrial fibrillation or flutter with complete AV block during which the ventricular rate becomes regular because of an idioventricular rhythm)

Fig. 12-46. Acute MI. Marked diffuse ST segment depression; note the ST elevation in aVR and little less so in V₁, a clue to the diagnosis of left main coronary artery occlusion.
Fig. 12-47. Right atrial hypertrophy.

Fig. 12-48. Atrial flutter. Note: Usually there is little visible evidence of flutter waves in lead I; V₅ and V₆ also tend to be silent or may reveal negative P-like waves.

Fig. 12-49. LBBB in a man with mitral valve bioprosthesis >20 years duration, marked precordial rocky motion caused by left ventricular aneurysm.

Fig. 12-50. LVH, and ischemia, axis 50, left anterior hemiblock, IRBBB, in a 65-year-old man with severe aortic regurgitation.

Fig. 12-51. Old anterior and lateral MI; left atrial hypertrophy; left axis −60, small q in I, small r in III: left anterior hemiblock.

Fig. 12-52. Complete heart block. Rate 38/min.

Fig. 12-53. Junctional tachycardia, rate 148/min. Note: P wave inverted in II, III, and aVF, positive in aVR, aVL.

Fig. 12-54. Atrial premature beats, with runs, also, junctional escape beats in V₄–V₆.

Fig. 12-55. RBBB with abnormal Q waves V₁, V₂, V₃: old anterior MI; APB, left axis −75, left anterior fascicular block (hemiblock).

Fig. 12-56. Left atrial hypertrophy: bifid P lead II, left atrial abnormality shown in V₁. Right axis, small r wave in lead I, small q in lead III: indicates probable left posterior hemiblock.

Fig. 12-57. Sinus bradycardia 44/min. Left ventricular hypertrophy and left atrial hypertrophy. A 50-year-old Vietnamese female with well controlled very mild hypertension for 10 years; semigiant T wave inversion V₅–V₆ is likely caused by apical hypertrophic cardiomyopathy as unlikely to be caused by very mild controlled hypertension. Echocardiogram shows some apical hypertrophy.

Fig. 12-58. RBBB; small Q in I, small r in III, and left axis −70 = left anterior fascicular block (hemiblock), atrial premature beat.


Fig. 12-60. Acute anterior MI. ST elevation V₁ through V₄ (STEMI).

Fig. 12-61. WPW syndrome mimicking RBBB; ECG from a 26-year-old male. A good reason to assess for WPW early in the interpre-
tive sequence (Step 3) done soon after the assessment for RBBB and LBBB.

**Fig. 12-62.** Acute inferior MI (STEMI); abnormally shaped high ST segment in inferior leads. Note the reciprocal depression in leads I, aVL, V₁, V₂.

**Fig. 12-63.** Ventricular tachycardia.

**Fig. 12-64.** Anteroseptal MI; age indeterminate. ECG from a 60-year-old man; ECG done during annual assessment, silent MI; the patient had a normal ECG 1 year earlier.

**Fig. 12-65.** Extensive anterior MI in a 50-year-old female. Note ST elevation in 8 leads.

**Fig. 12-66.** A 2:1 AV block, IRBBB; erroneously read by computer as APBs nonconducted. Note: With second degree AV (type I or type II block), the PP interval remains constant and the P wave morphology is unchanged. Note the P waves stuck to the T waves are not premature in time, and with nonconducted APBs, the PP interval will vary. Non-conducted APBs should not be mistaken for second degree AV block and vice versa. A 2:1 AV block can be either type I or type II.

**Fig. 12-67.** Accelerated junctional rhythm; IRBBB.

**Fig. 12-68.** Sinus tachycardia 125/min; APB.

**Fig. 12-69.** Acute anterior MI (SEMI). Sinus bradycardia 49; ECG from a 39-year-old male.

**Fig. 12-70.** Old inferior MI. Note the Q waves in II, III, aVF may be interpreted as “non diagnostic inferior Q waves noted.” The tracing is similar to 5 years prior. ECG from a 55-year-old female with severe hyperlipidemia (total cholesterol >8 mmol/L, 300 mg/dL from age 20 to 30). She had a proven inferior MI at age 32 with typical inferior Qs. Subsequent angina and CABG. Stable for the past 15 years. LDL maintained <2.5 mmol/L past 20 years. Wide inferior Q waves have become narrower and less deep over a 5-year period postinfarction.

**Fig. 12-71.** Atrial flutter.

**Fig. 12-72.** Acute anterior MI (STEMI).

**Fig. 12-73.** Old inferior MI. Note the Q waves in II, III, and aVF are distinct and diagnostic (see Fig. 12-70).

**Fig. 12-74.** Apical hypertrophic cardiomyopathy. Note the giant T wave inversion in keeping with apical HCM seen mainly in Japanese people. Despite the sinister looking ECG with giant T waves and high precordial QRS voltage, an outflow tract gradient does not develop and the prognosis is good compared with obstructive HCM. ECG from an 80-year-old Vietnamese woman with minimal cardiac symptoms over 10 years, during which time the ECG remained similar.
Fig. 12-75. Atrial fibrillation, ventricular rate 140/min.

Fig. 12-76. Nonsustained ventricular tachycardia.

Fig. 12-77. A and B. Wide complex regular tachycardia, rapid rate 235 to 260/min. Computer incorrectly interpreted Holter record as ventricular runs. Note in B the tachycardia is triggered by an APB. The wide complex rapid rate suggests preexcited antidromic tachycardia. An accessory pathway was documented and ablation was successful in this 28-year-old with 3-year duration of recurrent palpitations. He had presented once to ER with atrial fibrillation, ventricular rate 160/min.

Fig. 12-78. Hypertrophic cardiomyopathy. Poor R wave progression V₂–V₃, nonspecific ST-T wave changes, borderline IVCD, left anterior hemiblock, left atrial hypertrophy.

A constellation of abnormal findings in a 51-year-old female with shortness of breath. Echocardiogram showed asymmetric septal hypertrophy, septal thickness 1.7 cm, posterior wall 1.4 cm, left atrium 5.0 cm, systolic anterior motion of the mitral valve (SAM) with leaflets touching the septum, resting outflow gradient 75 mm Hg, increasing to 146 mm Hg after amyl nitrate.

Fig. 12-79. APBs. Atrial bigeminy.

Fig. 12-80. Complete AV block. Ventricular rate 28/min.

Fig. 12-81. ECG from a 74-year-old man who presented with chest pain. The marked diffuse ST segment depression in 10 leads accompanied by ST elevation in aVR greater than in V₁ suggested probable acute left main coronary occlusion and proved true on coronary angiograms.

Fig. 12-82. RBBB: note the prolonged duration of the S wave in lead 1, V₅, V₆, >30 ms.

Left axis −60; left anterior fascicular block.

Fig. 12-83. Sinus tachycardia 140/min. Non diagnostic inferior Q waves noted in a 31-year-old male with chest infection.

Fig. 12-84. ST segment elevation V₂–V₅ (fish hook feature in V₃): normal variant in a 30 year old male.

Fig. 12-85. Tracing from a healthy 60–year-old female. Poor R wave progression V₂, V₃ is a not uncommon finding caused by lead placement of V₂, V₃ in females. Mimics a probable old anteroseptal MI.

Fig. 12-86. Sinus rhythm, RBBB; atrial premature beats nonconducted: These are a common cause of an unexpected pause. It is preferable to use the term non conducted APB rather than blocked APB.

Fig. 12-87. A. ECG from a 47-year-old man. Age corrected Sokolow index (SV₁ + RV₅ or V₆ = 57 mm, 5.7 mV). The abnormal ST–T
change in V3 and the abnormal coving of the ST segment in V4–V6 should prompt a diagnosis of ischemia. See Figure 12-87 B: definitely LVH.

12-87 **B.** LVH proven in a 50-year-old female with long duration hypertension. Note the so-called typical “strain pattern” in V4 to V6: asymmetric ST segment depression; the T wave has a gradual descending and a steep ascending limb, a hallmark of LVH.

**Fig. 12-88.** RBBB, pathologic Q waves in V1–V4 indicates definite old anteroseptal MI.

In the presence of RBBB a q wave in V1–V2 may occur in the absence of MI. Also, the tracing shows left anterior fascicular block.

**Fig. 12-89.** Figure 12-82. WPW syndrome: changes mimic incomplete LBBB. The atypical bundle branch block, or conduction delay should prompt search for short PR and delta waves (I, II, aVL).

**Fig. 12-90.** RSR′ in V1–V2 suggests incomplete RBBB, but there is no slurred or widened S wave (the S wave is not of prolonged duration) in leads 1, V5 or V6 to indicate true RBBB. This should alert the interpreter to assess for atypical RBBB, a feature of Brugada syndrome. Scrutiny of the ST segment in V1, V2 reveals a coved and saddle-back deformity, characteristic features of the syndrome.
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