Rapid Interpretation of EKG's

Dr. Dubin’s classic, simplified methodology for understanding EKG’s

6th Ed.

Dale Dubin, MD
RAPID
INTERPRETATION
OF
EKG's

... an interactive course

by
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In 1790, an audience of usually sedate scientists gasped in disbelief as Luigi Galvani, with a flare of showmanship, made a dead frog’s legs dance by electrical stimulation.

Galvani knew that completing a circuit connecting dissimilar metals to the legs of a recently deceased frog would create a stimulating ________ current.

electrical

The resulting electrical current would stimulate the frog’s legs to jump, and with repeated stimuli he could make them ________.

dance

**Note:** But in those times, bringing a dead frog “back to life” was a shocking and ghastly “supernatural” feat. (And Galvani loved it!)*

* Get yourself a warm cup of coffee, relax and enjoy... the rest is just as easy and entertaining.
While conducting basic research around 1855, Kollicker and Mueller found that when a motor nerve to a frog’s leg was laid over its isolated beating heart, the leg kicked with each heartbeat.

"Eureka!" they thought, "the same electrical stimulus that causes a frog’s leg to kick must cause the heart to ______.”

So it was logical for them to assume that the beating of the heart must be due to a rhythmic discharge of _________ stimuli.

**Note:** And thus an association between the rhythmic pumping of the heart and electrical phenomena was scientifically established. Very basic and very important.
Mid 1880's, Ludwig and Waller

In the mid 1880's, while using a "capillary electrometer," Ludwig and Waller discovered that the heart's rhythmic electrical stimuli could be monitored from a person's skin.

This apparatus consisted of sensor electrodes that were placed on a man's ________ and connected to a Lippman capillary electrometer, which used a capillary tube in an electric field to detect faint electrical activity.

The fluid level in the capillary tube moved with the rhythm of the subject's ________-beat... very interesting.

This apparatus was a little too unsophisticated for clinical application, or even for economic exploitation, but it was ________ interesting.

Note: This momentous achievement opened the door for recording the heart's electrical activity from skin surfaces.
Enter Dr. Willem Einthoven, a brilliant scientist who suspended a silvered wire between the poles of a magnet.

Two skin sensors (electrodes) placed on a man were then connected to the ends of the silvered wire, which ran between the two poles of the ___________.

The silvered ________ (in the magnetic field) twitched to the rhythm of the subject's heartbeat.

This was also very interesting, but ________________ wanted a timed record.
Very clever, that Einthoven! The ____________ rhythmic
movements of the wire (representing the heartbeat)
created a bouncing shadow...

... that was recorded as a ____________ series
of distinct waves in repeating cycles.

He named the waves of each cycle (alphabetically)
P, QRS, and ____, T.

Note: “Now,” thought the clever Einthoven, “we can record a heart’s
abnormal electrical activity... and compare it to the normal.” And thus
a great diagnostic tool, his “electrokardiogram” (ElectroKardioGram),
evolved around 1901. Let’s see how it works...
The **electrocardiogram** (EKG) records the electrical activity of the heart, providing a record of cardiac electrical activity, as well as valuable information about the heart’s function and structure.

The electrocardiogram is known by the three letters ____; it provides us with a record of cardiac electrical activity and valuable information about the heart’s function and structure.

**Note:** Since the time of Einthoven’s “electrokardiogram,” the medical profession has used the letters EKG to represent the electrocardiogram. Some say that “ECG” is more correct, and you may see it used in some texts. However, Medicine honors tradition, and EKG has been used for years. Also, ECG sounds like EEG (the brain wave recording), and this can cause misunderstanding and confusion.

The EKG is inscribed on a ruled paper strip that gives us a permanent _________ of cardiac activity and the health status of the heart. Cardiac monitors and cardiac telemetry provide the same information in real time.
The EKG records the electrical activity of contraction of the heart muscle ("myocardium").

The information recorded on the EKG represents the heart’s electrical activity.

Most of the information on the EKG represents electrical activity of contraction of the myocardium.  

**Note:** The EKG also yields valuable information about the heart’s rate and rhythm.

When the myocardium (cardium = heart, myo = muscle) is electrically stimulated, it contracts.

**Note:** This illustration is intended to familiarize you with the simplified cross-section of the heart. The chambers are identified, and you should know them, for this diagram will be used often.
The interiors of heart muscle cells (myocytes*) are negative ("polarized") at rest, but when "depolarized" their interiors become positive and the myocytes contract.

While in the resting state, myocytes are polarized, the interior of every cell being __________-ly charged.

Note: In the strictest sense, a resting polarized cell has a negatively charged interior and a positively charged outside surface, but for simplicity we will consider only the negative interior.

The interiors of resting myocytes are negative, but when these cells are depolarized, their interiors become __________ and the cells contract.

"Depolarization" moves as a wave through the myocardium. As this wave of depolarization stimulates the heart’s myocytes, they become positive and __________.

* Just as the heart muscle is called the myocardium, its cells are called “myocytes”.
As a wave of depolarization progresses through the heart, it causes contraction of the myocardium.

Depolarization may be considered an advancing wave of _______ charges within the heart's myocytes. positive

Note: The depolarization wave initiates contraction of the resting myocytes as the charge within each cell changes to positive.

The advancing wave of depolarization causes progressive contraction of the myocardium as this wave of _______ charges positive passes through the interiors of the myocytes.

Note: The cell-to-cell conduction of depolarization through the myocardium is carried by fast-moving sodium (Na+) ions, the +'s in the illustration above.
The depolarization wave (cell interiors become positive), and a phase of *repolarization* (cell interiors return to negative) that follows, are recorded on the EKG as shown.

This stimulating wave of depolarization makes the interiors of the myocytes ___________ and stimulates them to contract. positive

Then the myocyte interiors regain their resting negative charge during the ________________ phase that follows. repolarization

**Note:** Repolarization is an electrical phenomenon that, in reality, begins immediately after depolarization. The broad hump that we see on EKG is the most active phase of repolarization.

Myocardial contraction is caused by ________________ of the myocytes, which records on the EKG as shown above. depolarization

The recovery phase that follows depolarization is known as ________________ (see illustration). repolarization
Sensors called “electrodes” are put on the skin to detect the heart’s electrical activity. The EKG machine records this activity on moving paper as an electrocardiogram.

Both depolarization and repolarization of the myocardium are ___________ phenomena caused by the movement of ions. electrical

The heart’s electrical activity may be detected and recorded from the _____ surface by sensitive monitoring equipment, skin including EKG machines, cardiac monitors, and telemetry devices.

The EKG records the electrical activity of the heart using skin sensors called _____________. electrodes
As the positive wave of depolarization within the myocytes flows toward a positive electrode, there is a positive (upward) deflection recorded on EKG.

**Note:** “Positive electrode,” of course, refers to a positive electrode actively recording a patient’s EKG.

A wave of depolarization advancing through the myocardium is a moving wave of ____________ charges. (Here come the Na⁺ ions!)

When this wave of positive charges (Na⁺ ions) moves toward a positive electrode, there is a simultaneous upward deflection recorded on the ________.

In general, when you see an upward wave on EKG, you know that it represents a depolarization wave moving toward a ______________ electrode.
The heart’s dominant pacemaker, the SA Node, initiates a wave of depolarization that spreads outward, stimulating the atria to contract as the circular wave advances.

**Note:** The SA Node (“Sinus Node”) is the heart’s dominant pacemaker, and its pacing activity is known as a “Sinus Rhythm.” The generation of pacemaking stimuli is automaticity. Other focal areas of the heart that have automaticity are called “automaticity foci.”

The SA Node, located in the upper-posterior wall of the right atrium, initiates a depolarization wave at regular intervals to accomplish its pacemaking responsibility.

Each depolarization wave of + charges (Na⁺ ions) proceeds outward from the SA Node and stimulates both atria to contract.

The ability of the SA Node to generate pacemaking stimuli is known as automaticity.  

**Note:** The depolarization wave flows away from the SA Node in all directions. Imagine the atria as a pool of water. A pebble dropped in at the SA Node produces an enlarging, circular wave (depolarization) that spreads outward. Atrial depolarization (and contraction) is a spreading wave of positive charges within the atrial myocardial cells. Let’s read this page again.
Each depolarization wave emitted by the SA Node spreads through both atria, producing a P wave on the EKG.

Note: The illustration depicts the positive wave of atrial depolarization advancing toward a positive skin electrode, producing an upward (positive) P wave on EKG.

The wave of depolarization sweeping through the atria can be detected by sensitive _______ electrodes.

Atrial depolarization is recorded as a ___ wave on EKG.

So when we see a P wave on an electrocardiogram, we know that, electrically speaking, it represents atrial ________________.

Note: The atria have a specialized conduction system, which we will examine later (page 101, if you’re curious).
Thus the P wave represents the electrical activity (depolarization) of both atria, and it also represents the simultaneous contraction of the atria.

As the wave of depolarization moves through both atria, there is a simultaneous wave of atrial _____________.

So the P wave represents the depolarization and contraction of both ___________.

**Note:** In reality, contraction of the atria lasts longer than the duration of the P wave. However, we’ll still consider that a P wave = atrial contraction. This simultaneous contraction of the atria forces the blood they contain to pass through the Atrio-Ventricular (AV) valves between the atria and the ventricles.
The Atrio-Ventricular (AV) valves prevent ventricle-to-atrium blood backflow, and they electrically insulate the ventricles from the atria… except for the AV Node, the only conducting path between the atria and the ventricles.

When the ventricles contract, the blood they contain cannot flow back into the atria due to the very efficient ___ valves. AV

The mitral and tricuspid (AV) valves lie between the atria and the ventricles, thereby acting to electrically _________ insulate the ventricles from the atria…

… leaving only the ___ Node as the sole pathway to conduct the depolarization stimulus through the fibrous AV valves to the ventricles.

Note: The AV Node is just above, but continuous with, a specialized conduction system that distributes depolarization to the ventricles very efficiently. Next we will review the movement of blood through the heart’s chambers.
Oxygen-depleted venous blood enters the right atrium. Atrial contraction forces blood through the *tricuspid valve* into the right ventricle, which pumps it into the lungs.

**Note:** Tricuspid is right side.

The right side of the heart (right atrium and right ventricle) receives under-oxygenated venous blood from all over the body, and pumps it into the **lungs**.

The right ventricle contracts, forcing the under-oxygenated venous blood through the **pulmonary valve** into the **pulmonary **artery, and thence to the lungs.

**Note:** Remember, both atria contract simultaneously, and also both ventricles contract together. However, the right and left sides of the heart have different responsibilities.
Oxygenated blood from the lungs enters the left atrium, which contracts to force blood through the mitral valve into the left ventricle. The powerful left ventricle, in turn, pumps blood through the aorta to all areas of the body.

**Note:** Mitral is left side.

The left atrium contracts, forcing oxygenated blood through the _______ valve into the left ventricle.

Then the muscular left ventricle contracts, forcing oxygenated blood through the aortic valve into the _______. (That's too easy!) aorta

Both atria contract simultaneously, then both __________ ventricles contract simultaneously.
When the wave of atrial depolarization enters the AV Node, depolarization slows, producing a brief pause, thus allowing time for the blood in the atria to enter the ventricles. Slow conduction through the AV Node is carried by calcium (Ca++) ions.

**Note:** Of course you remember that the AV Node is the only electrical conduction pathway between the atria and the ventricles.

Because depolarization slows within the AV Node, there is a brief delay or _________ before _________ pause _________ depolarization is conducted to the ventricles.

This brief pause allows the blood from the atria to pass through the AV valves and into the ________________. _________ ventricles

**Note:** At this point, we are correlating electrical activity with mechanical physiology. The atria contract, forcing blood through the AV valves, but it takes a little time for the blood to flow through the valves into the ventricles (hence the necessary pause that produces a short piece of flat baseline after each P wave on the EKG). Please review the illustration again.
Depolarization conducts slowly through the AV Node, but upon reaching the **ventricular conduction system**, depolarization rapidly shoots through the **His Bundle** and the **Left and Right Bundle Branches** and their subdivisions.

Depolarization conducts slowly through the AV Node, since it is carried by slow-moving Ca++ ions, but depolarization shoots rapidly through the ventricular conduction system beginning in the ____ Bundle. **His**

Depolarization conducts slowly through the AV Node, then rapidly through the His Bundle to the Right and Left _____________ Branches. **Bundle**

Depolarization shoots rapidly through the His Bundle and the Bundle Branches and their subdivisions, so depolarization is quickly distributed to the myocytes of the _____________. **ventricles**

**Note:** The ventricular conduction system originates at the His Bundle, which penetrates the AV valves, then immediately bifurcates (in the interventricular septum) into the Right and Left Bundle Branches. The His “Bundle” and both “Bundle” Branches are “bundles” of rapidly conducting **Purkinje fibers***. Like the myocardium, Purkinje fibers use fast-moving Na+ ions for the conduction of depolarization.

* Texts in the past (including this one) have *incorrectly* implied that only the terminal filaments were Purkinje fibers. Not so! Study “Note” and learn it correctly.
The terminal filaments of the Purkinje fibers rapidly distribute depolarization to the ventricular myocytes. Depolarization of the entire ventricular myocardium produces a QRS complex on EKG.

Note: The ventricular conduction system is composed of bundles of rapidly-conducting Purkinje fibers that carry depolarization away from the AV Node at high speed. The Purkinje fibers terminate in tiny filaments that directly depolarize the ventricular myocytes. The (rapid) passage of depolarization down the ventricular conducting system is too weak to record on EKG; however, depolarization of the ventricular myocardium records as a QRS complex.

Depolarization conducts slowly through the AV Node (using Ca^{++} ions), and then conducts rapidly (using Na^{+} ions) through the His Bundle to the Right and Left Bundle Branches into the terminal filaments of the Purkinje fibers, which depolarize the ventricular myocytes. ventricular

Note: The terminal filaments of the Purkinje fibers spread out just beneath the endocardium that lines both ventricular cavities, therefore ventricular depolarization begins at the lining and proceeds toward the outside surface (epicardium) of the ventricles. The Purkinje fibers branch and subdivide just beneath the endocardial lining, but they really do not penetrate into the myocardium. Since that's almost impossible to depict in a two-dimensional drawing, please recognize the limits of the illustration and remember it correctly.
The entire ventricular conduction system consists of rapidly conducting Purkinje fibers. The terminal filaments of the Purkinje fibers depolarize the ventricular myocardium, initiating ventricular contraction while inscribing a QRS complex on EKG.

The terminal filaments of the Purkinje fibers rapidly conduct depolarization to the myocytes that lie just beneath the endocardial lining of both ventricles.

**Note:** Remember, the entire ventricular conduction system, i.e., the His Bundle through the terminal filaments, is composed of Purkinje fibers that use fast-moving Na⁺ ions for conduction.

Depolarization of the ventricular myocytes produces a QRS complex on the electrocardiogram and initiates contraction of the ventricles.

**Note:** The QRS complex actually represents the beginning of ventricular contraction. The physical event of ventricular contraction actually lasts longer than the QRS complex, but we will still consider the QRS complex as generally representing the occurrence of ventricular contraction. So the QRS complex is an electrocardiographic recording of ventricular depolarization, which causes ventricular contraction. Still with me?
The Q wave is the first downward wave of the QRS complex, and it is followed by an upward R wave, however the Q wave is often absent on EKG.

The Q wave, when present, always occurs at the beginning of the QRS complex and is the first downward deflection of the complex.

The downward Q wave is followed by an upward ___ R wave.

Note: If there is any upward deflection in a QRS complex that appears before a “Q” wave, it is not a Q wave, for by convention, when present, the Q wave is always the first wave in the complex.

* It is now popular to use small (non-capital) letters to designate small waves in the QRS complex, for instance a “q” (small, lower case q) wave is a small wave.
The upward R wave is followed by a downward S wave. The entire QRS complex represents ventricular depolarization.

The first upward wave of the QRS complex is the _________.

Any downward wave PRECEDED by an upward wave is an _________.

The complete QRS complex can be said to represent ________ depolarization (and the initiation of ventricular contraction).

**Note:** An upward wave is always called an R wave. Distinguishing between the downward Q and downward S waves really depends on whether the downward wave occurs before or after the R wave. The Q occurs before the R wave, and the S wave follows the R. Just remember your alphabet.
Name the numbered waves in each QRS complex.

1. ____________

2. ____________

3. ____________

4. ____________

Q wave

R wave

S wave

QS wave

Note: Number 4 is a little unfair. Because there is no upward wave, we cannot determine whether number 4 is a Q wave or an S wave. Therefore it is called a **QS wave**, and it is considered to be a Q wave when we look for Q’s.
Following the QRS complex, there is a segment of horizontal baseline known as the ST segment, and then a broad T wave appears.

The horizontal segment of baseline that follows the QRS complex is known as the _____ segment. ST

After the QRS there is a segment of horizontal baseline, followed by a broad hump called the ____ wave. T

**Note:** The ST segment is horizontal, flat, and most importantly, it is normally level with other areas of the baseline. If the ST segment is elevated or depressed beyond the normal baseline level, this is usually an sign of serious pathology that may indicate imminent problems.

**Note:** The ST segment represents the “plateau” (initial) phase of ventricular repolarization. Ventricular repolarization is rather minimal during the ST segment.
The T wave represents the final, "rapid" phase of ventricular repolarization, during which ventricular repolarization occurs quickly and effectively.

Repolarization occurs so that the ventricular myocytes can recover their interior, resting negative charge, so they can be depolarized again.

Even though the T wave is usually a low, broad hump, it represents the rapid phase of ventricular repolarization.

Repolarization of the ventricular myocytes begins immediately after the QRS and persists until the end of the T wave.

Note: Repolarization (both phases) is accomplished by potassium (K⁺) ions leaving the myocytes.

Note: Ventricular systole* (contraction) begins with the QRS and persists until the end of the T wave. So ventricular contraction (systole) spans depolarization and repolarization of the ventricles. This is a convenient physiological marker.

* Pronounced "SISS-toe-lee"
The QT interval represents the duration of ventricular systole and is measured from the beginning of the QRS until the end of the T wave.

**Note:** The QT interval is a good indicator of repolarization, since repolarization comprises most of the QT interval. Patients with hereditary Long QT interval ("LQT") syndromes are vulnerable to dangerous (or even deadly) rapid ventricular rhythms. If you routinely examine the QT interval in all EKG's, eventually you will detect this anomaly, and probably save a patient's life during your career.

**Note:** With rapid heart rates both depolarization and repolarization occur faster for greater efficiency, so the QT interval varies with heart rate. Precise QT interval measurements are corrected for rate, so they are called QTc values. As a simple rule of thumb, the QT interval is considered normal when it is less than half of the R-to-R interval at normal rates.
Intermittent Mobitz (2° AV Block)

Occasional dropped QRS due to permanent BBB (one side) with intermittent BBB of the other side.

Simultaneous RBBB and LBBB prevents depolarization from reaching the ventricles; this is a complete (3°) AV block. So, block of one Bundle Branch with intermittent block of the other produces intermittent complete AV block, intermittent Mobitz.

Right BBB plus intermittent Left BBB will record on EKG as continuous Right BBB pattern QRS's with intermittent episodes of complete AV block (P waves without _____ response).

Left BBB plus intermittent Right BBB will record on EKG as continuous Left BBB pattern QRS's with intermittent episodes of complete AV block (P waves without _____ response).

Note: An EKG tracing or cardiac monitor display with a continuous BBB pattern QRS with an occasional missing QRS indicates intermittent complete AV block. The intermittent block may worsen, eventuating in a constant complete AV block. This intermittent Mobitz (exactly what it is) flashes an important warning sign. Intermittent Mobitz is the heart’s warning that eventually it will need an artificial pacemaker to drive the ventricles at a normal rate. Don’t let it slip by you unnoticed… for the patient’s sake!
Calcium (Ca\(^{++}\)) ions: cause myocyte contraction.

Potassium (K\(^{+}\)) ions: outflow causes repolarization of myocytes.

Sodium (Na\(^{+}\)) ion movement produces cell-to-cell conduction (of depolarization) in the heart...

except the AV Node, which depends on the (slow) movement of Ca\(^{++}\) ions.

The movement of three types of ions determines all aspects of cardiac conduction, contraction, and repolarization.

The release of free Ca\(^{++}\) ions into the interiors of the myocytes produces myocardial contraction.

Following depolarization, repolarization is due to the controlled outflow of K\(^{+}\) ions from the myocytes.

Cell-to-cell conduction (of depolarization) through the myocardium is carried by Na\(^{+}\) ions, however, AV Node conduction is due to the slow movement of Ca\(^{++}\) ions.

Note: This information may seem incongruous in an EKG text, however, during this millennium this page will prove to be the most important of all. Movement of these three ions is the very basis of cardiac physiology; this knowledge will serve you well in the future.

Note: Very soon, all health care professionals will understand (so easy!) cardiac function on the ionic-molecular level by reading Ion Adventure in the Heartland, which demonstrates how and why the electrical messages of the heart are displayed on EKG. See pages 331 and 332.
Chapter 2: Recording the EKG

The EKG is recorded on ruled (graph) paper. The smallest divisions are one millimeter (mm) squares.

The EKG is recorded on a long strip of ________ ruled (graph) paper, although some EKG machines record many different leads simultaneously on a large sheet.

The smallest divisions are one ________ long and one ________ high.

Between the heavy black lines there are ___ small squares. Each large square is formed by heavy black lines on each side, and each side is five mm long.

Note: As with all graphs, the time axis is horizontal and moves left to right, like we read. So timed events on EKG are measured left to right and similarly, cardiac monitors display a time axis that reads from left to right.
Innocuous Imitators of Intermittent Mobitz

dropped QRS at the end of a long Wenckebach series

non-conducted Premature Atrial Beat (P')

transient Sinus Block (P-QRS-T missing)

Since intermittent Mobitz may herald a complete AV block requiring a pacemaker, it is important that we recognize its characteristic span of clear baseline after a normal P wave. But, innocuous conditions can also produce a span of empty baseline.

A Wenckebach series (innocuous) produces a barren span of baseline after the terminal, punctual P wave, which is not conducted (review page 180).

A non-conducted Premature Atrial Beat (innocuous) strikes the AV Node while it is still refractory, so no stimulus is conducted to the ventricles (page 128); notice the peculiar, premature P' before the barren baseline.

A transient Sinus Block (usually innocuous, but the patient should be followed) can produce a pause before pacing resumes, or an automaticity focus may respond to the pause with an escape beat; in either case there is never an isolated P wave preceding the pause (review page 174).

**Note:** Simply stated:

- punctual P wave (no QRS response)... 2º AV block; Mobitz vs. Wenckebach
- premature P' wave (no QRS response)... non-conducted PAB
- missed P-QRS-T cycle... SA Node transiently blocked (Sinus Block)
Upward deflections are called “positive” deflections. Downward deflections are called “negative” deflections.

Positive deflections are __________ on the EKG.  

Negative deflections are __________ on the EKG.

**Note:** When a wave of stimulation (depolarization) advances toward a positive skin electrode, this produces a positive (upward) deflection on EKG. You will recall that depolarization is an advancing wave of positive charges within the cardiac myocytes. So with depolarization, the advancing wave of positive intracellular charges produces a positive deflection on EKG as this wave moves toward a positive electrode. Be positive!
The horizontal axis represents time.

Between the **heavy** black lines there are __ small squares.

The amount of time represented by the distance between two **heavy** black lines is ____________.

.2 of a second

Each small division (measured horizontally between two fine lines) represents ________________.

.04 of a second

(that's four hundredths!)
By measuring along the horizontal axis, we can determine the duration of any part of a cardiac cycle.

The duration of any wave may be determined by measuring along the horizontal _______.

Four of the small squares represents _____ of a second. .16
(sixteen hundredths)

The amount of EKG graph paper that passes out of the EKG machine in .12 second is _______ small squares. three (3)
(You don’t have to be a mathematician to read EKG’s.)
The standard EKG is composed of 12 separate leads.*

A standard EKG is composed of six limb leads, recorded by using arm and leg electrodes and...

... there are also six chest leads obtained by placing a suction cup electrode at six different positions on the chest.

**Note:** Leads not considered “standard” may be monitored from various locations on the body as required for special diagnostic purposes.

* Rhymes with seeds.
To obtain the **limb leads**, electrodes are placed on the right arm, the left arm, and the left leg. A pair of electrodes is used to record a lead.

By placing electrodes on the right and left arms and the left leg, we can obtain and record the _____ leads.

**Note:** Einthoven used these three locations for limb electrodes. They remain the conventional standard for recording the EKG.

The placement of these _____ is the same electrodes as originally used by Willem Einthoven.

**Note:** Two electrodes are used to record a lead. A different pair is used for each lead.
Each bipolar limb lead is recorded using two electrodes. So by selecting a different pair of electrodes for each lead, we create three separate bipolar limb leads (lead I, lead II, and lead III) for recording.

Each limb lead consists of a pair of electrodes, one is positive and one is __________, so these leads are called “bipolar” limb leads.

Lead I is horizontal, and its left arm electrode is __________, positive while its right arm electrode is negative.

When we consider lead III, the left arm electrode is now __________, and the left leg electrode is negative.

**Note:** The engineering wonders of the EKG machine permit us to make any skin electrode positive or negative depending on which pair of electrodes (that is, which lead) the machine is recording.

**Note:** The bipolar limb lead configuration is sometimes called “Einthoven’s triangle.”
Bipolar Limb Leads

moved to intersect at a center point

By pushing the three (bipolar) limb leads to the center of the triangle, we produce three intersecting lines of reference.

The triangle has a center, and each ______ may be lead
moved to that center point.

By pushing leads I, II, and III to the center of the triangle, three intersecting lines of ______ are formed. reference

Although the three bipolar limb leads may be moved to the ______ of the triangle, they remain at the same angles center relative to each other. (They’re still the same leads, yielding the same information.)
Another standard lead is the AVF lead. The AVF lead uses the left foot electrode as positive and both arm electrodes as a common ground (negative).

The AVF lead uses the left foot electrode as ___________.

In AVF both the right and left arm electrodes are channeled into a common ground that has a ___________ charge.

**Note:** Dr. Emanuel Goldberger, who designed and introduced the “Augmented” limb leads, discovered that in order to record a lead in this manner, he had to amplify (Augment) the Voltage in the EKG machine to match the wave magnitude of leads I, II, and III. He named this lead: A (Augmented), V (Voltage), F (left Foot), and he went on to produce two more leads using this same technique.

**Aside:** Your deductive mind tells you that lead AVF is a combination of leads II and III... just what Dr. Goldberger was trying to accomplish! Therefore lead AVF is a cross between (and oriented between) those two bipolar limb leads. Now, let’s create two more augmented leads.
The remaining two augmented limb leads, AVR and AVL, are obtained in a similar manner.

For the AVR lead the Right arm electrode is positive, and the remaining two electrodes are ___________.

To obtain the AVL lead, the Left arm electrode is made ___________; the other two electrodes are negative.

**Note:**
- AVR — Right arm positive
- AVL — Left arm positive
- AVF — Foot (left foot) positive

(These augmented limb leads are sometimes called the "unipolar" limb leads, stressing the importance of the positive electrode.)
The augmented limb leads, AVR, AVL, and AVF, intersect at different angles than those produced by the bipolar limb leads, and they produce three other intersecting lines of reference.

AVR, AVL, and AVF are the augmented (or “unipolar”) ___ limb leads.

These augmented limb leads intersect at 60 degree angles, but the angles differ from those formed by bipolar limb leads, I, II, and III.

Leads AVR, AVL, and AVF intersect at angles different from leads I, II, and III. In fact, leads AVR, AVL, and AVF split the angles formed by leads I, II, and III.
Six Limb Leads of the Standard EKG

All six limb leads (I, II, III, and AVR, AVL, and AVF) meet to form six intersecting leads that lie in a flat "frontal" plane on the patient's chest.

The six limb leads consist of the three bipolar leads, I, II, III, and three augmented leads, AVR, AVL, and AVF.

If the bipolar limb leads I, II, and III are superimposed on augmented limb leads AVR, AVL, and AVF, we have six intersecting leads in a flat plane on the patient's frontal plane.

Note: Don't get bedazzled by the kaleidoscope of limb leads. Bear with me for a few pages, and soon you will understand their utility, and a simplified way to visualize this concept.
Each camera position represents the positive electrode of a standard limb lead. Each limb lead (I, II, III, AVR, AVL, and AVF) records from a different angle (viewpoint), to provide a different view of the same cardiac activity.

Note: The heart’s electrical activity remains constant, but the positive electrode position changes from lead to lead. Therefore the tracing looks slightly different in each lead, as the angle from which we record the electrical activity changes with each lead. Remember, a wave of depolarization is a progressive wave of POSITIVE charges passing through the myocardial cells. So, when a depolarization wave moves toward a POSITIVE electrode, a POSITIVE (upward) deflection is produced on the EKG (or monitor) for that particular lead. (A little repetitious, but it is so important!)

The EKG records the same cardiac activity in each lead.

The waves look different in various leads because the heart’s electrical activity is recorded from a different angle (viewpoint) for each lead.

* If this were a video camera, it could record the information for a cardiac monitor.
It is conceptually necessary for you to visualize the six intersecting limb leads. Why? Can you identify this car?

**Note:** This page sure seems empty, doesn’t it?

**Note:** Automobile experts are encouraged not to recognize the car for the sake of understanding the concept.
By observing the same object from six different angles, you will obtain a great deal of information, and in this case, perhaps even identify the car.

**Note:** You can’t see the car’s rear bumper in the photo at top left. But with progressively different views, you can determine more about the bumper (or even the driver). Similarly, it may be difficult to see a specific wave in a given lead, but with six different lead positions, it is certain to show up better in other leads.

**Note:** Observation from six different angles is better than one. Thus recording cardiac electrical activity from six different angles gives us a much greater and more accurate perspective. At this point you can take a sip of coffee and relax. By the way, the car is a 1965 Ford Thunderbird, but it is far more important that the concept (not the car) always remain in your mind.
The importance of the positive electrode’s position is emphasized by the conventional grouping of limb leads. A positive left arm electrode is used to record “lateral leads” I and AVL, and a positive left foot electrode is used to record “inferior leads” II, III and AVF. The location of the positive electrode is the key.

**Leads I and AVL** are called the *lateral leads* (left lateral understood) because each has a __________ electrode positioned laterally on the left arm.

**Leads II, III, and AVF** are called the *inferior leads* because each of these leads has a positive electrode positioned inferiorly on the left ______.

**Note:** So now you can determine if depolarization is moving toward (or even away from) the patient’s left side, and the same for depolarization directed inferiorly toward (or even away from) the left foot. The “inferior leads” and the “lateral leads” include 5 of the 6 limb leads. These are not arbitrary designations. These terms are common cardiology parlance and have important clinical/diagnostic significance. Know and understand them.
To obtain the six standard chest leads, a positive electrode (suction cup) is placed at six different positions (one for each lead) on the chest.

The six chest leads are recorded from six progressively different positions around the _________. (See illustration.)

For each of the chest leads, the suction cup electrode that is placed on the chest is considered _________.

The chest leads are numbered from V₁ to V₆ and are positioned in successive steps from the patient’s right to the left side of the chest. Notice how the chest leads cover the ________ in its normal anatomical position within the chest.

Note: Traditionally a suction cup electrode records the chest leads, however adhesive electrodes are now commonly used. Because the electrode for the chest leads is always positive, a depolarization wave moving toward a given chest electrode produces a positive (upward) deflection in that chest lead of the EKG tracing.
In general, each of the chest leads\(^*\) is oriented through the AV node and projects through the patient’s back, which is negative.

**Note:** The plane of the chest leads (called the “horizontal” plane) cuts the body into top and bottom halves.

The electrode for each of the chest leads is always considered ________ (positive or negative).  
positive

If leads \(V_1\) through \(V_6\) are imagined to be the spokes of a wheel, the center of the wheel is the __________.  
AV Node

**Lead** \(V_2\) describes a straight line directly from the front to the back of the patient. In lead \(V_2\) the patient’s back is considered ________ (positive or negative).  
negative

\(^*\) The chest leads, also called the “precardial” (in front of the heart) leads, were introduced by Dr. Frank Wilson.
By examining an EKG, you will notice that the waves in the six chest leads show progressive changes from \( V_1 \) to \( V_6 \).

**Note:** When observing the chest leads from \( V_1 \) to \( V_6 \) you will see gradual changes in all the waves (as the position of the positive electrode changes for each successive lead).

In the illustration the \( V_1 \) chest lead, the QRS complex is mainly _______ (positive or negative) normally.

In chest lead \( V_6 \) the QRS complex is usually mainly _______ (positive or negative). Understand why.

Observing the \( V_6 \) chest lead, we know that the mainly positive QRS complex is produced by ventricular depolarization moving _______ the POSITIVE chest electrode of \( V_6 \) (if you’re a little unsure about this concept, take another look at page 12).
Leads $V_1$ and $V_2$ are oriented over the right side of the heart, while $V_5$ and $V_6$ are oriented over the left side of the heart.

Leads $V_1$ and $V_2$ are called the "________" chest leads.  

The two chest leads oriented over the left side of the heart are ___ and ___, (and are called the "left" chest leads).  

A depolarization wave moving toward the (positive) chest electrode in lead $V_6$ causes an ________ deflection on the EKG tracing of this lead. (Now you understand!)
Leads $V_3$ and $V_4$ are generally oriented over the *interventricular septum*. $V_3$ is nearer the right ventricle, and $V_4$ is nearer the left ventricle.

Leads $V_3$ and $V_4$ are oriented over the area of the interventricular septum.

**Note:** The interventricular septum is a common wall shared by the right and left ventricles, so this septum separates the cavity of the right ventricle from the cavity of the left ventricle. The Right and the Left Bundle Branches course through the interventricular septum.
On the standard EKG tracing there are six chest leads and six limb leads. This is the 12 lead electrocardiogram.

The six limb leads all lie in the ________ plane, which can be visualized on the patient’s chest.

The six chest leads lie in the horizontal plane and are arranged in progressive order from V₁ to _____.

The six chest leads are recorded using a positive electrode, which is placed at six specific anatomical positions on the chest, encircling the heart in the ________ plane.
The six limb leads also can be recorded by using carefully positioned electrodes on the trunk of the patient. The special electrode placement (above) used for exercise (“stress”) testing, can be used to record each of the twelve EKG leads.

**Note:** An EKG recorded from a carefully positioned trunk* electrode can record the same information (same accuracy and same amplitude) as an ankle or wrist electrode for a given limb lead. In this way, a standard twelve lead EKG can be recorded using trunk electrodes.

Cardiac monitoring in hospital rooms, as well as in the emergency department, surgery, recovery room, coronary care, and intensive care, is carried out using modified electrode positions on the patient’s trunk* to monitor classical limb (and other) leads.

Paramedics and many Emergency Medical Technicians (EMT’s) use trunk* electrodes for diagnostic purposes and also for telemetry transmission.

*These are “trunk” but not truly “chest” electrodes, for they often use the shoulders and abdomen as electrode locations. A variety of modifications are commonly used to monitor patients in various settings and circumstances (see page 346).
The **Autonomic Nervous System** (ANS) regulates vital functions of all organs by both reflex and central nervous system control, but *not* by conscious control.

Although the ANS controls all organs and organ systems, our main concern is autonomic control of the ________, heart and also of the systemic arteries as they relate to blood pressure.

**Note:** The ANS has two divisions that sometimes seem difficult to comprehend, because one division may stimulate an organ, yet inhibit another organ. However, these two divisions have well defined roles in the heart and the systemic arteries; *one division stimulates, and one division inhibits.* It's that simple! One division stimulates cell function, while the other division opposes that stimulation. Each division operates like an electrical system that controls its own terminal switches called “receptors” that modulate the function of cells.

**Note:** A stimulus originating in the ANS is transmitted to a **ganglion** of secondary nerve cells for processing. The nerves of the ganglion relay the stimulus to their nerve ends, each of which terminates as a disc called a **bouton** (bouton is French for button) that covers the receptors of a cardiac cell (or an arterial muscle cell). See next page.

*“Ganglion” is singular, “ganglia” is plural.*
The ANS consists of a sympathetic system and an opposing parasympathetic system. Each of these two systems secretes its own neurotransmitter from its terminal boutons in order to activate specific cell receptors in the cell membrane.

The terminal ends (boutons) of sympathetic nerves secrete Nor-epinephrine* (N-epi), an adrenaline-like neurotransmitter that activates specific ______ receptors called adrenergic receptors.

**Note:** In the heart, the sympathetic and parasympathetic nervous systems have opposite functions. Interestingly, the parasympathetic exercises some direct control of the sympathetic.

The terminal parasympathetic nerve ends (boutons) secrete the neurotransmitter Acetylcholine (ACh), which exclusively activates cell ___________ called cholinergic receptors.

* Nor-epinephrine, hyphenated for recognition purposes, will be “Norepinephrine” from now on. Although the abbreviation “N-epi” is used here, some texts use “NE.”
**Sympathetic System**

activates cardiac $\beta_1$ adrenergic receptors

**Cardiac Excitatory Effects**

- rate of SA Node pacing
- rate of conduction
- force of contraction
- irritability of foci

The heart is stimulated by the sympathetic system through its terminal boutons. The boutons deliver N-epi to the $\beta_1$ (adrenergic) receptors; this activates the $\beta_1$ receptors, producing an excitatory response at the cellular level.

**Norepinephrine** (N-epi), the neurotransmitter of the sympathetic system, activates the heart’s $\beta_1$ (adrenergic) receptors, stimulating the SA Node to pace faster.

N-epi also:

- improves AV Node conduction and accelerates conduction through the atrial and ventricular myocardium.
- increases the force of myocardial contraction.
- and increases the irritability of atrial and Junctional (page 123) automaticity and minimally affects ventricular foci.

**Note:** N-epi’s brother, epinephrine (“adrenaline”) is secreted into the blood by the adrenal glands. Epinephrine is an even more potent stimulator of the heart’s $\beta_1$ receptors.

$^*$ $\beta_1$ adrenergic receptors” is often shortened to “$\beta_1$ receptors,” but *adrenergic* is understood.
Parasympathetic System activates cholinergic receptors

Note: There are two vagus nerves, left and right. Each vagus nerve supplies the heart and g.i. tract.

Cardiac Inhibitory Effects
- rate of SA Node pacing
- rate of conduction
- force of contraction
- irritability of atrial and Junctional foci

Parasympathetic nerves release the neurotransmitter acetylcholine (ACh), which activates cardiac cholinergic receptors (most are within the atria) to produce a cardiac inhibitory effect. Conversely, the gastrointestinal tract is stimulated by its parasympathetic innervation.

Parasympathetic activation of cholinergic receptors by ACh:

...inhibits the SA Node, decreasing the heart rate,

...decreases the speed of myocardial conduction, and depresses the AV Node,

...diminishes the force of myocardial contraction,

...and depresses irritability of automaticity mainly those in the atria and AV Junction.

Note: The vagus nerves are the body’s main parasympathetic pathway, so “vagal” stimulation means parasympathetic stimulation, with the understanding that vagal “stimulation” of the heart is inhibitory.

Note: Despite the parasympathetic system’s inhibiting effect on the heart, parasympathetic activation of cholinergic receptors stimulates the gastro-intestinal tract. Recalling the agony of severe vomiting or an episode of painful, crampy diarrhea will help you remember the effect of excessive parasympathetic stimulation of the stomach and the bowel.
Besides controlling the SA Node’s pacing rate, the Autonomic Nervous System controls blood flow and blood pressure by regulating constriction and dilation of arteries throughout the body.

Sympathetic stimulation of arterial $\alpha_1$ (adrenergic) receptors constricts arteries throughout the body, increasing blood pressure and blood flow. The $\alpha_1$ receptors are more responsive to the neurotransmitter N-epi than to circulating epinephrine.

**Note:** By pulling both ends of the Greek alpha, the center loop of the “$\alpha$” constricts the artery (see arrows on the $\alpha$ in the illustration). Now you will always remember alpha adrenergic sympathetic effects on systemic arteries.

Parasympathetic activation of arterial (cholinergic) receptors dilates the same arteries as above, reducing blood pressure and blood flow. Besides the direct cholinergic inhibition of the arteries, there is also an indirect inhibitory parasympathetic effect on the sympathetic ganglia that send nerve fibers to the vessels.

**Note:** Blood flow is also very dependent on the heart rate: sympathetic stimulation increases the SA Node pacing rate, while the parasympathetic decreases it. Autonomic control of the heart rate and systemic blood pressure involves delicate regulation of the parasympathetic-sympathetic balance to maintain circulatory homeostasis (the ideal status quo).
Severe pain and/or seeing one’s own blood may induce a reflex parasympathetic response that causes syncope* (loss of consciousness).

Severe pain and/or seeing one’s own blood often initiates reflex parasympathetic activity that slows SA Node pacing, known as _______cardia.

The same reflex parasympathetic response dilates systemic arteries causing hypotension, as the blood _______ falls.

**Note:** A devastating injury, which causes excruciating pain/awareness of bleeding, can induce a parasympathetic response that dramatically lowers the blood pressure and slows the heart. This merciful reflex effectively reduces the brain’s blood supply to the point of syncope.

**Aside:** Perhaps you have encountered (oversensitive) patients who lose consciousness upon seeing their own blood drawn for lab tests, or if they experience minimal pain. Be compassionate; their body is only responding to a normal parasympathetic reflex.

*This and other types of vagally mediated syncope are sometimes called “vaso-vagal syncope.” Syncope is pronounced “SINK-oh-pee.”
Cardiovascular sensors provide (“afferent”) input for parasympathetic reflexes that counterbalance sympathetic effects. **Vagal maneuvers** may be employed to produce a reflex parasympathetic response for both diagnostic and therapeutic purposes.

**Gastrointestinal stimulation** (e.g., gag reflex) may be employed to produce a parasympathetic _________.

**Carotid sinus massage** may be used in carefully selected patients* to produce a _______________ response.  

**Note:** An induced parasympathetic response may be used therapeutically to depress an irritable focus in the atria or AV Junction. An induced parasympathetic response may be diagnostically employed to transiently slow AV Node conduction, or to make the AV Node more refractory to depolarization (see pages 160 and 183).

*Injudicious use of carotid sinus massage in some patients can dislodge a piece of atheromatous plaque, sending plaque emboli to the brain (careful!). Use discretion when employing diagnostic procedures that can cause iatrogenic stroke, which can disable a patient and incite a feeding frenzy of lawyers.
It seems as though standing would allow blood to gravitate into the lower extremities. However, standing produces a compensatory sympathetic response that constricts peripheral arteries to prevent distal blood pooling, and stimulates sinus pacing.

**Note:** The body has “pressure” receptors\(^a\) that detect low blood pressure, particularly with standing. These pressure receptors initiate a sympathetic reflex that constricts peripheral arteries and increases the heart rate slightly, thereby conserving blood flow to the brain.

Impaired function of this normal sympathetic response to standing can diminish blood flow to the brain, causing \underline{**syncope**}

**Orthostatic hypotension** is an abrupt fall in blood pressure caused by failure of these compensatory sympathetic mechanisms upon \underline{**standing**}

\(^a\) Careful! These “receptors” are cardiovascular sensors, called baroreceptors, that the body uses to detect changes in blood pressure. Please don’t confuse them with the cell membrane receptors that are activated by N-epi or ACh.
Standing produces a sympathetic vasoconstriction response to maintain adequate circulation. This compensatory mechanism may fail with prolonged standing in certain elderly patients, triggering a paradoxical parasympathetic response causing syncope.

Note: Pooling of blood in the lower extremities from prolonged standing is normally compensated by a reflex sympathetic increase in both blood pressure and heart rate. However, in some elderly patients, sinus pacing accelerates, but peripheral vasoconstriction is inadequate. So, the partially filled ventricles contract rigorously, stimulating parasympathetic mechanoreceptors in the left ventricle. This initiates an undesirable parasympathetic reflex that slows SA Node pacing and reduces blood pressure; so blood flow to the brain is reduced, causing syncope. This is neuro-cardiogenic syncope.

Neuro-cardiogenic syncope, a (paradoxical) parasympathetic response to prolonged standing, causes vasodilation and slowing of the pulse, resulting in a loss of ____________________ consciousness.

Under controlled circumstances, a Head Up Tilt ("HUT") test confirms the diagnosis of neuro-cardiogenic ____________ syncope.
1. Rate
2. Rhythm
3. Axis
4. Hypertrophy
5. Infarction

Knowing basic cardiac principles and understanding the autonomic nervous system ensures mastery these five general areas of routine EKG interpretation.

Proper interpretation of an _____ requires consideration of Rate, Rhythm, Axis, Hypertrophy, and Infarction. They are all equally important.

**Note:** Take a moment and examine page 334 to observe the simple methodology that will become your routine.

Before you begin each chapter, preview its summary (pages 335 to 346). Then, as you progress through the chapter, little “aha’s” of recognition will flash in your brain, and you will appreciate how each concept is carefully woven into this simplified methodology. Your understanding evolves rapidly; this is the foundation of your permanent knowledge.

Ready? Let’s go!
When reading an EKG, you should first consider the rate.

**Note:** The sign in this picture is not informing the driver* about the rate of his race car. The man holding the sign is a physician who has been monitoring the driver's transmitted EKG. The sign is telling the driver about his current heart rate (he's a little excited).

When examining an EKG, you should determine the ________ first.

The rate is read as cycles per ____________.

**Now,** let's examine where and how the normal heart rate originates...

*With a sincere dedication to Billy Occam, deceased long ago, who made simplicity a virtue of science.*
The SA Node (Sinus Node), the heart’s pacemaker and the dominant center of automaticity, generates a **Sinus Rhythm**. The SA Node paces the heart in the normal rate range of 60 to 100 per minute.

The heart’s normal pacemaker, the _______, generates a continuous series of regular, pacemaking stimuli (this is its "**automaticity**").

The SA Node is located within the upper-posterior wall of the right _______. The SA Node emits a regular series of pacemaking (depolarization) stimuli.

**Note:** The Sinus Node (SA Node) is the heart’s dominant center of automaticity, and the normal, regular rhythm that it generates is called the Sinus Rhythm.

At rest, the Sinus Rhythm maintains a rate of 60 to _____ beats per minute, which is the normal range of the pacing rate.
If the Sinus Node (SA Node) paces the heart at a rate slower than 60 per minute, this is Sinus Bradycardia.

**Note:** “Brady” = slow; “cardia” = heart.

A rhythm originating in the heart’s normal pacemaker, the SA Node, with a rate slower than 60 per minute is called Sinus _________________.

**Note:** Sinus Bradycardia most often results from parasympathetic excess, as we see in conditioned athletes at rest. Sometimes an extremely slow heart rate may reduce blood flow to the brain causing loss of consciousness (syncope). See pages 60 and 63.

Sinus Bradycardia is present if the SA Node produces a heart rate of less than one beat per ____________. (careful!)
If the Sinus Node (SA Node) paces the heart at a rate greater than 100 per minute, this is Sinus Tachycardia.

**Note:** “Tachy” = fast; “cardia” = heart.

A rhythm originating in the SA Node (Sinus Node) is called Sinus Tachycardia if the rate is greater than ______ per minute.

Exercise produces sympathetic stimulation of the SA Node; this is the most common cause of Sinus ______________. Tachycardia

**Note:** There are focal areas of automaticity in the heart known as automaticity foci.* They are potential pacemakers that are capable of pacing in emergency situations. Under normal circumstances, these foci are electrically silent (that’s why they are referred to as “potential” pacemakers).

* “Automaticity foci” refers to more than one “automaticity focus”; in fact, when the word “foci” is used alone, “automaticity foci” is understood. Foci is pronounced “FOE-sigh.”
If normal SA Node pacemaking fails, other potential pacemakers known as automaticity foci (also called "ectopic" foci) have the ability to pace (at their inherent rate). They are in the atria, the ventricles, and the AV Junction.

If the SA Node ceases to function, one of the potential pacemakers, known as an automaticity focus, will assume pacemaking activity at its inherent rate (only one focus assumes pacing responsibility). Rate

The atria have automaticity foci of potential pacemakers that are within the atrial conduction system (see page 101), and they are called atrial automaticity foci.

Note: The proximal end of the AV Node has no automaticity foci, however the middle and distal regions of the AV Node, an area known as the AV Junction does have automaticity foci that are called Junctional automaticity foci.

Purkinje fibers have automaticity foci, so there are foci of these potential pacemakers in the His Bundle and in the Bundle Branches and their subdivisions; these foci are called ventricular automaticity foci.
The automaticity foci of each "level" (the atria, the AV Junction, and the ventricles are each a "level") have a general range of pacemaking rate. Although all foci of a given level pace within a general rate range, each individual automaticity focus has its own precise inherent rate at which it paces.

Each automaticity focus of the atria has a specific inherent rate at which it paces, but its inherent rate falls within the general range of ___ to 80 per minute.

The automaticity foci of the AV Junction all pace in the range of ___ to 60 per minute, but any single Junctional focus paces at its individual inherent rate.

Ventricular automaticity foci all pace in the ___ to 40 per minute range, but any specific ventricular focus has a distinct inherent rate of pacing.
A cardiac cycle is represented by the P wave, the QRS complex, the T wave, and the baseline that follows until another P wave appears. This cycle is repeated continuously. Please study the illustration to make certain that you understand every event in sequence.

**Note:** Physiologically, a cardiac cycle represents atrial systole (atrial contraction), followed by ventricular systole (ventricular contraction), and the resting stage that follows until another cycle begins.

Atrial depolarization (and contraction) is represented by the ___ wave.

Ventricular depolarization (and contraction) is represented by the ______ complex.

**Note:** In reality, atrial contraction lasts longer than the P wave, and ventricular contraction lasts longer than the QRS complex, but you already knew that.
**SA Node overdrive-suppresses all foci**
(since all foci have a slower inherent pacing rate)

- **Atrial foci** *(inherent rate 60 – 80 per min.)*
- **Junctional foci** *(inherent rate 40 – 60 per min.)*
- **Ventricular foci** *(inherent rate 20 – 40 per min.)*

*Rapid automaticity (pacemaking activity) suppresses slower automaticity* *(pacemaking activity)* – this is **overdrive suppression**, a very important fundamental characteristic of all automaticity centers.

**Note:** Overdrive suppression is characteristic of all centers of automaticity (including the SA Node and all automaticity foci). Simply stated: any automaticity center will overdrive-suppress* all others that have a slower inherent pacemaking rate.

The SA Node overdrive-suppresses the (slower) inherent pacemaking activity of all the automaticity foci below it; this provides the SA Node with the luxury of not having to compete with slower pacemaking activity of lower automaticity foci.

In fact, once an automaticity focus actively begins pacing, it will **overdrive-suppress** all lower (slower) foci, including slower foci at the same level…

…eliminating any competition. Well Designed!

*When used as a verb, “overdrive-suppress” is hyphenated… so says the publisher.
Overdrive suppression is the heart’s failsafe pacing mechanism, providing three separate levels of backup pacing, by utilizing automaticity foci in the atria, the ventricles, and the AV Junction.

**Note:** An automaticity focus actively pacing at its inherent rate, overdrive-suppresses all slower foci including slower foci at its own level.

Should normal SA Node pacing fail (pacemaker failure), a backup pacemaker (i.e., an automaticity focus from a lower level) – no longer overdrive-suppressed – will emerge to pace at its inherent rate; and it conveniently overdrive-suppresses potential pacemaking activity at all levels that are ________ it.

Therefore, an automaticity focus only emerges to function as a pacemaker when it is no longer __________-suppressed.

For instance, in SA Node failure...

...a focus from a lower level – no longer overdrive-suppressed by regular pacing stimuli from above – can emerge to pace.

*Very* well Designed!

Let’s do that once again, slowly.
The atria have automaticity foci of potential pacemakers, any one of which can assume active pacemaking responsibility in its inherent rate range of about 60 to 80 per minute, if normal pacemaking fails.

If SA Node pacing fails, an atrial automaticity focus can assume the active pacemaking responsibility in its inherent rate range of about 60 to ___ per minute (close to the SA Node’s normal rate). 80

If the SA Node fails, an atrial automaticity ______ focus (within the atrial conduction system) may then assume active pacing responsibility to become the dominant pacemaker. (The last sentence on this page explains “dominant” pacemaker.)

So without SA Node pacing, an atrial automaticity focus can emerge as an active backup pacemaker, and it becomes the dominant pacemaker by overdrive-suppressing all lower levels of foci, since they have slower inherent rates. inherent
The AV Junction has automaticity foci (potential pacemakers), one of which will emerge to actively pace in its inherent rate range of 40 to 60 per minute if there is an absence of regular pacing stimuli progressing down from the atria.

**Note:** The AV Junction is that portion of the AV Node that has foci of automaticity. The proximal end of the AV node has no foci. The AV Junction has foci of automaticity called “Junctional foci.”

An automaticity focus in the AV Junction begins active backup pacing only in the absence of pacing stimuli coming down from the atria. Then, no longer overdrive-suppressed, it emerges to actively pace in its inherent range of 40 to 60 per minute, and it overdrive-suppresses all lower (slower) automaticity foci, becoming the dominant pacemaker.

A Junctional focus actively pacing at its inherent rate (___ to 60 per minute range), produces an *idio-junctional* rhythm.

**Note:** A Junctional focus (that is, an automaticity focus in the AV Junction) emerges as the active pacemaker if it is no longer overdrive-suppressed by regular pacing stimuli from above. This can occur if the SA Node and all atrial foci fail. But wait — something else can prevent a Junctional focus from being depolarized by regular pacing stimuli from above. Next page!

* The prefix “idio” is of Greek origin, and it means “one’s own.”
* Idiojunctioonal is usually *not* hyphenated.
(AV) Junctional Automaticity Focus
(potential pacemaker)

If there is a complete conduction block in the AV Node above the AV Junction, then no regular paced depolarization stimuli from above reach the automaticity foci in the AV Junction.

**Note:** You will recall that the AV Node is the only conduction link between the atria and the ventricular conduction system below.

With a complete conduction block in the AV Node above the AV Junction, an automaticity _______ in the AV Junction focus
just below, receives no pacing stimuli from above...

...so, no longer overdrive-suppressed, the Junctional focus escapes to become the active pacemaker for the ventricles. And that Junctional focus paces the ventricles at its inherent rate of ____ - ____ per minute while overdrive-suppressing lower, 40 - 60 (slower) ventricular foci.

**Note:** It is possible for the AV Junction (together with all its automaticity foci) to suffer a complete block. In that instance, only an automaticity focus in the Purkinje fibers of the ventricles can come to the rescue to pace the ventricles. Let’s see how...
The ventricles have automaticity foci (potential pacemakers), any one of which will assume pacing in its inherent rate range of 20 to 40 per minute, if the usual overdrive suppression (due to regular pacing stimuli from above) is absent.

**Note:** *Ventricular automaticity foci* are composed of specialized Purkinje fibers. These pacemaking foci are in the His Bundle, the Bundle Branches, and all their subdivisions, since they are all composed of Purkinje fibers.

Without overdrive suppression from above, a ventricular automaticity focus emerges to actively pace in its inherent rate range of ____ to 40 per minute; this is an *idio-ventricular* rhythm.

**Note:** A ventricular focus emerges as the active ventricular pacemaker only if it is no longer overdrive-suppressed by regular, paced stimuli from above. This occurs:
- if all pacemaking centers above it have failed.
- or-
- if there is a complete block of conduction below the AV Node (including the AV Junction) that prevents any pacing stimulus above it (i.e., from the SA Node, an atrial focus, or a Junctional focus) from reaching the ventricles.

*Hyphenated here for ease of recognition, *idioventricular* should not be hyphenated.*
OVERDRIVE SUPPRESSION
provides emergency backup pacing at 3 separate levels

Emergency Failsafe Pacing Mechanism

If Failure...
  ➔ Atrial focus assumes pacing responsibility.

If Failure...
  ➔ Junctional focus assumes pacing responsibility.

If Failure...
  ➔ Ventricular focus assumes pacing responsibility.

Range of Inherent Pacing Rates of Automaticity Foci

<table>
<thead>
<tr>
<th>50</th>
<th>60</th>
<th>80</th>
<th>100</th>
</tr>
</thead>
<tbody>
<tr>
<td>Atrial</td>
<td>Junctional</td>
<td>Ventricular</td>
<td></td>
</tr>
</tbody>
</table>

If normal SA Node pacing fails, an automaticity focus in the atria, or the AV junction, or even the ventricles (in that order) is available to assume the pacemaking responsibility at its own inherent rate. This provides three levels of backup pacing.

If the SA Node should cease pacing, an atrial automaticity focus can pace in its inherent rate range of 60 to 80 per minute; failing that, backup pacing by a Junctional focus will assume the active responsibility in its (slightly slower) inherent rate range of 40 - 60 per minute.

The ventricles can be paced by a ventricular automaticity focus in a rate range of 20 to 40 per minute, if the focus is regularly depolarized by paced stimuli. The lack of properly paced stimuli to the ventricular focus can be due to failure of all automaticity centers above, or due to an intervening complete block that prevents pacing stimuli (from above) from conducting to the ventricles. What a miracle of Nature!

Note: In a physiological or pathological emergency, an irritable automaticity focus may suddenly discharge at a rapid rate. This emergency rate (150 to 250 per minute) is approximately the rate for foci of all levels.

Now let's try something real easy…
The height and depth of a wave are measured vertically from the baseline in millimeters, and this vertical amplitude represents a measure of voltage. *

The height or depth of waves is measured from the baseline in millimeters and is a measure of voltage.

**Note:** The deflection of a wave is the direction in which it records on EKG; for instance, the "upward deflection" or "downward deflection" of a wave. However, the amplitude of a wave is the magnitude (in millimeters) of upward deflection or downward deflection. The height or depth of a wave (i.e., its amplitude) is a measure of voltage.

The first wave in the illustration has an upward deflection of 3 mm in amplitude.

**Note:** The elevation or depression of segments of baseline is also measured vertically in millimeters, just as we measure waves.

* Ten millimeters vertically represents one millivolt (mV), however, in practice, one usually speaks of "millimeters" of height or depth (waves) and the same for elevation or depression of baseline segments.
Our main objective is to rapidly determine the heart rate.

After finishing this chapter you will be able to determine the _______ rapidly.

No special devices, calculators, rulers, or awkward mathematical computations are needed in order to determine the rate.

**Note:** In emergency situations, you probably will not be able to find, much less use a calculator; and you may not have the presence of mind (or the time) to do mathematical calculations.

Observation alone can give us the _______.

First: Find an R wave that peaks on a heavy black line (our “start” line).

To calculate rate, you should first look at the ___ waves.

Now find one that peaks on a heavy black line, and we will call it the “_______” line.

start
Next: Count off "300, 150, 100" for the three thick lines that follows the start line, naming each line as shown. Memorize these numbers.

An R wave peaks on a heavy black start line...

...the next heavy black line is named "______"... followed by "______" and "______" for the next two heavy black lines.

Note: The line that the R wave peaks upon is the start line; we only name the heavy lines that follow the start line.

The three lines following the start line (where the R wave falls) are named "______, ______, ______" in succession. (Say them out loud!)

Again!
Then: Count off the next three lines after "300, 150, 100" as "75, 60, 50."

The next three lines after "300, 150, 100" are named "______, 60, 50."

Remember the next three lines together as:
- "______, ______, ______." 75, 60, 50

Once more out loud, please.

Very good!
Now: Memorize these triplets until they are second nature. Make certain that you can say the triplets without using the picture.

These triplets, “300, 150, 100” and “75, 60, 50” must be memorized.

Be able to name the lines that follow the start line on which an R wave _______; it is easy to remember them as triplets, and so easy to use immediately. (Can hardly wait!)

Do not count those lines that follow the start line - name them with the ____________ as you go.
Where the next R wave falls, determines the rate. It's that simple.

Find an R wave peaked upon a heavy black (start) line, then look for the _____ R wave. next

Where the next R wave falls gives the _______. rate
There is no need for mathematical computations.

If the next R wave falls on "75"... the rate is 75 per _________. minute

**Note:** You may have noticed that the illustration shows the normal rate range of 75 to 100.
By knowing the triplets “300, 150, 100” then “75, 60, 50” you can merely look at an EKG and tell the approximate rate immediately.

The triplets are: first “_____, _____, _____." 300, 150, 100

then “_____, _____, _____." 75, 60, 50

By simply naming the lines following the start line, you can identify the rate immediately using the ________. triplets
Practice Tracing

This is an EKG tracing from a resting patient, whose heart rate is slower than the usual rate one would see with a Sinus Rhythm. Let's examine the rate.

The rate in the above tracing is about ____ per minute.

If you were told that this rhythm probably originated in an automaticity focus, by the rate alone, you would suspect the origin (pacemaker) to be in the ____ ___________.

AV Junction

Note: This is indeed a rhythm originating in the AV Junction, and that's why you don't see P waves. This elderly woman has a very diseased heart. Her SA Node failed, then all the atrial automaticity foci failed. Fortunately, a Junctional focus came to the rescue. This natural pacing backup system is wonderfully effective.
You do not need to depend on mathematical computations in order to calculate the rate. Observation alone will do it!

You can rapidly determine the rate on an EKG tracing by _________ alone.

There is no need to depend on annoying math or calculators (where did I put that darn thing?) in order to determine the _________.

**Note:** You will always have your brain with you (at least until that time when brain transplants would provide you with someone else’s brain). Just remember to name the lines that follow the *start* line using the triplets, and say: “300, 150, 100” then “75, 60, 50.” Enough, enough... let’s try it!
Now, let's determine the approximate rates of these EKG tracings.

A. 

B. 100

C. 150 or so

D. 60

E. 75

Note: As you may have discovered for yourself, any prominent wave (like the S wave in example B.) can be used to determine the rate.
The distance between the heavy black lines represents 1/300 min.

So two 1/300 min. units = 2/300 min. = 1/150 min.
(or 150/min. rate)

and three 1/300 units = 3/300 = 1/100 min.
(or 100/min. rate)

There is a logical explanation for the seemingly unusual rate determinations using the triplets.

**Note:** The unit of time (duration) between two heavy black lines is .2 sec., which is 1/300th of a minute.

The number of time units between five consecutive heavy black lines is ____.

So this represents 4/300 minute or a rate of ____ per minute.

Therefore if a heart contracts 75 times per minute, there will be a span equal to the distance between five heavy black lines between the ____ complexes.

**Note:** Reasonable instructors should not require students to master this page. As author, I have not personally memorized the text material on this boring page. Let’s keep it simple and practical.
The fine line divisions can provide more precise rate determination. Memorizing them is impractical, so when determining fast rates, most of us use a reference like the one provided in the Personal Quick Reference Sheets (page 335).

**Note:** It is admittedly a great task to memorize the fine line subdivisions, so you can use page 335 as a personal reference when you need it. Determining the rate range using the triplets is more than adequate in most cases.

**Note:** For rates less than sixty per minute, see the next few pages for a simple way to determine rate when you see a bradycardia.
Bradycardia  
(slow rates)

For very slow rhythms, there is an easy method for quickly determining the rate.

The proper term for slow heart rate is _________________.

Note: The triplets give us a very large range of rates. By using the triplets “300, 150, 100” then “75, 60, 50” you can determine rates ranging from 300 to 50. Bradycardia means a rate slower than 60 per minute.

For bradycardia you can use another easy method to determine the _____. I’ll show you on the next page...
On the top margin of every EKG strip, there are small marks that identify the "three second" intervals.

There are small marks above the graph portion of the EKG tracing. Find a strip of EKG tracing and examine it.

Two of these marks enclose a three second interval.

Note: Some EKG paper has "3 second intervals" that are marked with a dot, circle, triangle, or a vertical line.

When an EKG machine is running, the span of paper between two of these "3 second interval" marks passes under the stylus needle in 3 seconds.
Taking two of the three second intervals, we have a 6 second strip.

**Note:** A three second interval is obviously the distance between two consecutive 3 second interval marks.

Taking two of the three second intervals gives us a 6 second _______.

This 6 second strip represents the amount of paper used by the machine in six seconds (one-_______ of a minute).
Count the number of complete (R wave to R wave) cycles in this 6 second strip. With marked bradycardia, there will be few cycles per 6 second strip.

The length of a cardiac __________ can be measured from a specific wave until the wave is repeated again.

So R wave to ____ wave gives us the duration (length) of one cardiac cycle.

Count the number of cycles in the 6 second ______. strip

Next page…
Find the rate by multiplying the number of cycles in the six second strip by ten (10). Ten of the 6 second strips equals one ___________ (time) of EKG tracing.

The number of cycles per minute is the ________.

So cycles per 6 second strip multiplied by ___ equals the rate. Simple!
So, if there are 4 cycles per six sec. strip...

the rate is 40

You can just place a zero on the right of the number of cycles per six second strip, and you have the rate.

For very slow heart rates (bradycardia), you should first find a six second strip,

...count the number of cycles in this strip,

...and multiply by \( \frac{10}{4} \) to get the rate.

Note: Multiplying by ten may be done by placing a zero on the right side of the number of cycles per six second strip. For instance, five cycles (per six second strip) gives a rate of 50.
Let's determine the approximate rates of these EKG's.

Rates: No. 1. ____ per minute

No. 2. ____ per minute

No. 3. ____ per minute

Note: The general, average rates of irregular rhythms are usually determined using this method.

Why don't you obtain some EKG tracings and amaze yourself (and your friends) at how easily you can determine the rate.

Note: Take a minute to review the illustrations in this chapter, then turn to the Personal Quick Reference Sheets at the end of this book for a simplified summary of determining rate (page 335).
Chapter 5: Rhythm, Part I

The EKG provides the most accurate means of identifying a cardiac arrhythmia (abnormal rhythm), which can be diagnosed easily, once we understand the electrophysiology of the heart.

"Arrhythmia" literally means without ________, however ________, however rhythm

It is used to denote any abnormal rhythm. The term "arrhythmia" (bad rhythm) has the same meaning, and also is commonly used in medical literature.

The EKG records the heart’s electrical phenomena that may not be seen, felt, or heard on physical examination, so the EKG is a very accurate means of recording ________ disturbances. rhythm

Note: To understand the arrhythmias, you must first become familiar with the normal electrophysiology of the heart, including the normal conduction pathways.
The SA Node generates a regular* Sinus Rhythm that paces the heart. Each pacemaker impulse from the SA Node (Sinus Node) spreads through both atria as an advancing wave of depolarization.

It is the automaticity of the Sinus Node (SA Node) that generates the regular* cadence of depolarization stimuli for pace-________ activity.

Normally, the SA Node discharges regular pacing impulses (60 to 100 per minute) that depolarize the ______.

**Note:** We know that the SA ("Sino-Atrial") Node is the same as the Sinus Node, so we understand that the terms "Sinus" and "Sino" imply SA Node origin.

* The term "regular" indicates a rhythm of constant rate. See next page...
Normal (Regular) Rhythm

On EKG there is a consistent distance (duration) between similar waves during a normal, regular cardiac rhythm, because the SA Node’s automaticity precisely maintains a constant cycle duration between the pacing impulses that it generates.

**Note:** All automaticity foci pace with a regular rhythm. This is a characteristic of all automaticity centers.

The SA Node generates pacing impulses at a constant, unvarying rate, producing cycles of equal length, so the rhythm of the heart is said to be ________. This characteristic pattern of regularity is typical of SA Node pacing.*

And, because the sequence of depolarization is the same on each repeating cycle, there is a predictable regularity of all similar (named) waves. Therefore, irregularities at ________ are easy for you to spot on EKG. rhythm

**Note:** We can visually scan an EKG and appreciate the repetitive continuity of a regular rhythm. But breaks in that continuity, such as a pause, the presence of too-early (premature) beats, or sudden, dramatic change, immediately catch our attention, warning us of a rhythm disturbance.

*In reality, a Normal Sinus Rhythm varies imperceptibly with respiration.
A normal physiological mechanism, **Sinus Arrhythmia**, sounds pathological ("arrhythmia" = abnormal rhythm), but it functions in all humans at all times. The autonomic nervous system causes barely detectable rate changes in Sinus pacing that relate to the phases of respiration. This is not a true arrhythmia.

**Note:** Sinus Arrhythmia is a normal, but extremely minimal, increase in heart rate during inspiration, and an extremely minimal decrease in heart rate during expiration.

Sinus Arrhythmia represents normal, minimal variations in the SA Node's pacing rate in association with the phases of respiration.

**Note:** The slight increase in the heart rate is due to inspiration-activated sympathetic stimulation of the SA Node. The slight decrease in pacing rate is due to expiration-activated parasympathetic inhibition of the SA Node. Perhaps you knew that already, since Sinus pacing is regulated both divisions of the Autonomic Nervous System.

**Note:** This variability of Sinus Rhythm is normal. In fact, if the heart rate variability is reduced, this is pathological and is a valuable indicator of increased mortality, particularly after infarction. Parameters of "Heart Rate Variability" are being established for determining patient prognosis in many types of heart disease.
The atrial conduction system consists of three specialized internodal tracts in the right atrium (the Anterior, the Middle, and the Posterior), and one conduction tract known as Bachmann’s Bundle that innervates the left atrium.

Three conduction pathways in the right atrium course from the SA Node to the AV Node (thus the term “Intermodal”). They are the Anterior, the Middle, and the ____________ Posterior Internodal Tracts.

Bachmann’s Bundle originates in the SA Node and distributes depolarization to the left ______ atrium.

Depolarization passing rapidly through the atrial conduction system does not record on EKG; however, depolarization of the atrial myocardium produces a ___ wave on EKG. P

Note: Just as ventricular automaticity foci are within the ventricular Purkinje fibers, similarly, atrial automaticity foci are within the specialized atrial conduction system. Because there is a concentration of merging atrial conduction tracts in the immediate region of the AV Node near the coronary sinus, *considerable automaticity activity originates in that area.

* The heart’s own venous drainage (i.e., from the myocardium) empties into the right atrium via the coronary sinus.
When the depolarization stimulus (passing down from the atria) reaches the AV Node, the stimulus slows in the AV Node, producing a pause on EKG.

Atrial depolarization eventually reaches the AV Node, but conduction of depolarization slows within the AV Node, recording a ________ on EKG.

This pause (during which blood from the atria passes into the ventricles) is represented by the horizontal piece of baseline between the P wave and the ______ complex.

**Note:** The AV Node is named for its position between the Atria and the Ventricles (thus “AV”). The proximal end of the AV Node has no automaticity foci. However, the remainder of the AV Node, an area known as the AV Junction, does have automaticity foci. These foci are essential for backup pacing should there be a total failure of all pacemaking activity from above (SA Node as well as atrial foci), or (this is important) if a complete conduction block of the proximal end of the AV Node occurs, preventing all (SA Node or atrial foci) pacing stimuli from being conducted to the ventricles.
After passing slowly through the AV Node, depolarization proceeds rapidly through the His Bundle, Bundle Branches and their subdivisions, and through the terminal Purkinje filaments to distribute depolarization to the ventricles. Ventricular depolarization produces a QRS complex on EKG.

**Note:** The His Bundle and the Bundle Branches are "bundles" of rapidly conducting Purkinje fibers. Depolarization passing through the Purkinje fibers of the ventricular conduction system is too weak to record on EKG; this is a form of "concealed" conduction.

After creeping through the AV Node, depolarization shifts gears and races through the His ________, and... Bundle

...through the Right and Left Bundle Branches and their subdivisions to rapidly transmit depolarization via the terminal Purkinje filaments to the endocardial surface of the ____________ myocardium. ventricular

When the ventricular myocardium depolarizes, it produces a ________ complex on EKG. QRS

**Note:** The Purkinje fibers of the ventricular conduction system contain automaticity foci (you knew that already).
The Purkinje fibers of the ventricular conduction system rapidly conduct depolarization away from the AV Node to the endocardial surface of the ventricles; when the ventricles depolarize, it produces a QRS complex on EKG.

**Note:** Ventricular depolarization begins midway down the interventricular septum, where the Left Bundle Branch produces fine terminal filaments. The Right Bundle Branch does not produce terminal filaments in the septum. So left-to-right depolarization of the septum occurs immediately before the rest of the ventricular myocardium depolarizes. (Examine the illustration.)

Ventricular depolarization initiates ventricular contraction, which persists (through both phases of repolarization) to the end of the ___ wave.

Ventricular contraction begins and ends during the ___ interval.

**Note:** Repolarization of the Purkinje fibers takes longer than ventricular repolarization. That is, the end of the T wave marks the end of ventricular repolarization; however, repolarization of the Purkinje fibers terminates a little later — beyond the end of the T wave. The final phase of Purkinje repolarization may record a small hump, the U wave (following the T wave), on EKG.
SA Node overdrive-suppresses all foci (since all foci have a slower inherent pacing rate)

There are three levels of automaticity foci (atrial, junctional, and ventricular) that can assume backup pacemaker responsibility if pacing activity fails. The foci of each level have a characteristic inherent rate range, giving the SA Node a failsafe capacity of three levels of backup pacing.

Each level of automaticity foci has a consistent range inherent rate.

Note: The SA Node and all automaticity foci are centers of automaticity ("automaticity centers"), which means that they can generate regular pacing stimuli.

Overdrive suppression allows the automaticity center with the fastest rate to be the dominant pacemaker (no competition).

Should the highest pacemaking center fail, an automaticity focus from the next highest level (no longer overdrive-suppressed) emerges ("escapes") to actively pace at its inherent rate, and it then becomes the dominant pacemaker by overdrive-suppressing all automaticity foci below it.

Note: A very "irritable" automaticity focus may suddenly pace rapidly.
Arrhythmias

• Irregular Rhythms (page 107)
• Escape (page 112)
• Premature Beats (page 122)
• Tachy–arrhythmias (page 146)

The arrhythmias can be divided into a few general categories, according to the arrhythmia’s mechanism of origin. The best students, I’ve noticed, apply index tabs to the pages that begin each arrhythmia category (see above); try it – you will find it very helpful!

Note: Although arrhythmia literally means “without rhythm,” generally it describes any rhythm disturbance, that is, any variance from a Normal Sinus Rhythm. Some authors prefer the term “dysrhythmia” rather than arrhythmia.

Note: The illustration is a simplified arrhythmia classification that is categorized according to the mechanism of origin, so the arrhythmias will be easy for you to understand.

Note: The underlying mechanisms that are basic to the heart’s function are very satisfying to learn. But more importantly, conceptual understanding of the basic mechanisms facilitates and perpetuates your knowledge. Don’t memorize patterns; your knowledge will be vital to others! Lasting knowledge results from understanding.
Irregular Rhythms

- Wandering Pacemaker
- Multifocal Atrial Tachycardia
- Atrial Fibrillation

The irregular rhythms presented in this section are usually caused by multiple, active automaticity sites.

Rhythms that lack a constant duration between paced cycles are said to be ___________.

irregular

Note: The term “irregularly irregular” is an old designation that describes an irregular and chaotic rhythm that has no predictable recurring pattern.

Note: In some hearts with structural pathology or hypoxia, malfunctioning automaticity foci may suffer from entrance block, whereby any incoming depolarization is blocked, “protecting” them from passive depolarization by any other source. Such “protection” is not healthy. By being insensitive to passive depolarization, they cannot be overdrive-suppressed, while their own automaticity is still conducted to surrounding tissue. When an automaticity focus has entrance block, it is said to be parasystolic (the focus paces, but can’t be overdrive-suppressed).
**Wandering Pacemaker** is an irregular rhythm produced by the pacemaker activity wandering from the SA Node to nearby atrial automaticity foci. This produces cycle length variation as well as variation in the shape of the P' waves. The overall rate, however, is within the normal range.

**Note:** The P' (pronounced “P prime”) wave represents atrial depolarization by an automaticity focus, as opposed to normal Sinus-paced P waves.

**Note:** Each automaticity focus has a specific inherent rate at which it paces. In a given lead, each atrial automaticity focus produces its own morphological signature, that is, it produces a P' wave of a distinctive shape related to the anatomical location of that focus within the atria.

Wandering Pacemaker is an irregular rhythm (normal rate range); the pacemaking activity wanders from the SA Node to _____ foci...

... so the cycle lengths vary, and ___ wave morphology (shape) varies as the pacemaking center moves.

**Note:** Should the rate accelerate into a tachycardia (greater than 100 per minute), it becomes _Multifocal Atrial Tachycardia_. Next page...
Multifocal Atrial Tachycardia (MAT) is a rhythm of patients with Chronic Obstructive Pulmonary Disease (COPD). The heart rate is over 100 per minute with P' waves of various shapes, since three or more atrial foci are involved.

In MAT, we can recognize each P' wave from a particular focal atrial focus by its morphological signature, i.e., P' waves from the same focus look the same in a given lead.

Note: MAT is an arrhythmia of patients who are very ill with COPD. The atrial automaticity foci are also ill, showing early signs of parasystole (entrance block) by developing a resistance to overdrive suppression. That is why no single focus achieves pacemaking dominance, so they all pace together.

Because of the multifocal origin of MAT, each individual atrial focus paces at its own inherent rate, but the total, combined pacing of multiple unsuppressed foci produces a rapid, irregular rhythm...

... and in a given lead, each focus produces P' waves with a specific morphological signature, i.e., P' waves of a distinct shape (note that some P' waves are identical, since they're from the same focus). And remember, this is a tachycardia.

*MAT is sometimes associated with digitalis toxicity in patients with heart disease.
**Atrial Fibrillation**

**irregular rhythm**
- continuous chaotic atrial spikes
- irregular ventricular rhythm

*Atrial Fibrillation* is caused by the continuous rapid-firing of multiple atrial automaticity foci. No single impulse depolarizes the atria completely, and only an occasional, random atrial depolarization reaches the AV Node to be conducted to the ventricles; this produces an irregular ventricular (QRS) rhythm.

**Note:** Atrial Fibrillation is NOT an arrhythmia of healthy, young individuals. It is the result of multiple "irritable" atrial foci, suffering from entrance block, pacing rapidly. These multiple atrial foci are *parasystolic*, so they’re all insensitive to overdrive suppression; therefore, they all pace at once. What chaos!

During Atrial Fibrillation, no single impulse completely depolarizes both _____, so there are no P waves, just a rapid series of tiny, erratic spikes on EKG.

Only the occasional atrial impulse gets through the AV Node to initiate a _____ complex. The *irregular ventricular response* may result in either a slow or rapid ventricular rate, but it is always irregular.

**Note:** You must determine and document the general ventricular rate in Atrial Fibrillation (QRS’s per six second strip times 10).
This tracing was monitored from a patient with an irregular pulse.

This practice tracing has an irregular rhythm in which we see discernible ___ waves, so we know that it is not P' Atrial Fibrillation.

The "P" waves are not identical, and the rate does not gradually increase and gradually decrease, so we immediately know that this is not ______ Arrhythmia. Sinus

The rate is less than 100 (which rules out MAT), the rhythm is irregular, and the P' waves are of different shapes. This is most likely ______________ Pacemaker. Wandering

Easy, isn’t it!

Note: Just to solidify your knowledge of these irregular rhythms, study the simplified review and tracings of Irregular Rhythms on page 336.
Escape Rhythm - an automaticity focus escapes overdrive suppression to pace at its inherent rate:

- Atrial Escape Rhythm
- Junctional Escape Rhythm
- Ventricular Escape Rhythm

Escape Beat - an automaticity focus transiently escapes overdrive suppression to emit one beat:

- Atrial Escape Beat
- Junctional Escape Beat
- Ventricular Escape Beat

"Escape" describes the response of an automaticity focus to a pause in the pacemaking activity.

The SA Node’s regular pacing overdrive-suppresses all automaticity foci, but a brief pause in SA Node pacing permits an automaticity ______ to escape overdrive suppression.

If SA Node pacing ceases entirely, an automaticity focus will escape to pace at its inherent _____, thereby producing an Escape Rhythm. We will, however, need to identify the focus (atrial, Junctional, or ventricular) that escapes to actively pace.

If the pause in pacing is brief (only one cycle missed), an automaticity focus may ________ to emit a single Escape Beat before the normal Sinus rhythm returns. So, we will need to identify that focus (atrial, Junctional, or ventricular).
**Sinus (SA) Node pacing interrupted by Sinus Arrest**

Sinus Arrest occurs when a very sick SA Node ceases pacemaking completely. But the heart’s efficient, failsafe mechanism provides three separate levels of automaticity foci for backup pacemaking. Divine Design.

**Note:** With Sinus Arrest, the SA Node ceases pacing; then, absent overdrive suppression by the SA Node, an automaticity focus (with the fastest inherent pacing rate) escapes to become an active pacemaker. And since it has the fastest inherent rate, it overdrive-suppresses all foci below, to become the dominant pacemaker.

**Note:** An automaticity focus is overdrive-suppressed if it is regularly depolarized by a pacing rate faster than its own inherent pacing rate. But if an automaticity focus is not overdrive-suppressed – regardless of the cause – it escapes to initiate its own pacemaking activity.

**Note:** Each specific focus has its own individual, inherent rate of pacing. However, the inherent pacing rates of all foci of a given level (for example, the inherent rates of all Junctional foci) are within a rate range.

**Note:** With a Sinus Arrest, the SA Node ceases pacing, so absent overdrive suppression from above, an automaticity focus escapes to produce an Escape Rhythm. However with Sinus Block, the SA Node misses one pacing cycle, producing only a transient pause. So an automaticity focus escapes to emit an Escape Beat, which actually represents the first beat of the attempt by the focus to pace, but the return of SA Node pacing overdrive-suppresses it again.
With Sinus Arrest an atrial focus quickly escapes overdrive suppression to become the dominant pacemaker at its inherent rate. This is an **Atrial Escape Rhythm**.

With a Sinus Arrest, an automaticity focus in the highest level of foci, the ______, escapes overdrive suppression to become an active pacemaker in its inherent rate range of 60 to 80 per minute.

An Atrial Escape Rhythm originates in an atrial automaticity focus, so the P' waves are not identical to the previous P waves that were produced by the ______. (See illustration.)

**Note:** The active atrial automaticity focus overdrive-suppresses all lower, slower foci to become the dominant pacemaker. It also paces at its inherent rate, which differs from (i.e., is slower than) the previous Sinus rate. (See illustration.)

When an atrial focus assumes pacing responsibility in the absence of a Sinus Rhythm, this is an Atrial ______ Rhythm.
Sinus Rhythm  Junctional Escape Rhythm
"idiojunctional rhythm"

Absent regular pacing stimuli from above, an automaticity focus in the AV Junction may escape overdrive suppression to become an active pacemaker producing a Junctional Escape Rhythm in its inherent rate range: 40 to 60 per minute.

Note: A Junctional focus escapes the influence of overdrive suppression if there is a Sinus Arrest, and the atrial foci also fail to function properly...

... or if there is a complete conduction block in the proximal end of the AV Node. In either case, the Junctional focus is not regularly stimulated by pacing depolarizations from above.

When a Junctional focus is not overdrive-suppressed, it actively paces, producing a Junctional Escape Rhythm, and it becomes the dominant pacemaker of the ventricles at a rate ranging from 40 to ___ per minute (it’s also called an “idiojunctional rhythm”).* 60

A Junctional Escape Rhythm usually conducts mainly to the ventricles, producing a series of lone ___ complexes. QRS

But see the next page for an interesting exception.

*Sometimes the inherent Junctional pacing rate may accelerate beyond its usual range to produce an Accelerated Idiojunctional Rhythm.
A Junctional Automaticity Focus May Cause Retrograde Atrial Depolarization

Each P' is inverted in leads with an upright QRS

Because each Junctional automaticity focus is located within the AV Node, each pacing stimulus originating there will conduct to the ventricles as expected, but the paced stimuli may also (unexpectedly) depolarize the atria from below ("retrograde"), producing inverted P' waves in EKG leads with an upright QRS.

**Note:** The illustration shows that atrial depolarization and ventricular depolarization proceed in opposite directions from a pacing Junctional focus. Also, most EKG leads produce an upright QRS.

With a Junctional Escape Rhythm, every paced stimulus will depolarize the ventricles, but the pacing may also depolarize the atria from below in a retrograde fashion, producing inverted P' waves in EKG leads with an upright QRS.

**Note:** The AV Node conducts very slowly, so depolarization from a Junctional focus may delay either ventricular depolarization or retrograde atrial depolarization (if present)...

... as a result, if there is retrograde atrial depolarization from a Junctional focus, it may record on EKG with one of these three patterns:

- retrograde (inverted) P' wave immediately before each QRS
- retrograde (inverted) P' wave after each QRS
- retrograde (inverted) P' wave buried within each QRS (not shown)
Ventricular Escape Rhythm occurs when a ventricular automaticity focus is not regularly stimulated by paced depolarizations from above, so it escapes overdrive suppression to emerge as the ventricular pacemaker with an inherent rate in the range of 20 to 40 per minute* (so it is also called an “idiioventricular rhythm”). Notice the continuous ventricular complexes.

Ventricular Escape Rhythm usually results from one of two mechanisms:

- With complete conduction block high in the ventricular conduction system (but below the AV Node), the ventricular foci are not stimulated by atrial depolarizations from above (see P waves in illustration), so a ventricular focus escapes to pace the ventricles at its inherent rate.

- Total failure of the SA Node and all automaticity foci above the ventricles is a rare and grave condition called “downward displacement of the pacemaker”. In extremis, a ventricular focus escapes to become the active ventricular pacemaker in a final, futile attempt to sustain life.

Note: Pacing from a ventricular focus is often so slow that blood flow to the brain is significantly reduced to the point of unconsciousness (syncope). This is Stokes-Adams Syndrome. This unconscious patient’s airway must be monitored and maintained... constantly.

*Should this accelerate above the inherent rate range, it becomes an Accelerated Idioventricular Rhythm.
During a Sinus Rhythm, a transient Sinus Block makes the SA Node miss a pacing stimulus (one missed cycle), producing a pause in pacing. So an atrial automaticity focus escapes overdrive suppression to emit an Escape Beat.

With a transient Sinus Block, an unhealthy SA Node misses one pacing stimulus. This missed cycle produces a ______ during which the heart is electrically silent.

If this pause is long enough (see NOTE below), then an automaticity focus will "escape" ______ suppression.

**Note:** If there is a “sufficient” pause — longer than the inherent (pacing) cycle length of an automaticity focus — that focus will "escape" the SA Node’s overdrive suppression to emit a stimulus.*

If the SA Node misses only one cycle, it will then resume pacing, and the SA Node’s overdrive suppression of all automaticity _____ resumes as well.

* If you don’t understand the Note, don’t worry. Just be aware of the escape mechanism.
A transient Sinus Block of one pacemaking stimulus (SA Node misses one cycle) is a sufficient pause for an atrial automaticity focus to escape overdrive suppression and emit an Atrial Escape Beat. Notice that the P' wave differs from the Sinus-generated P waves.

A transient Sinus Block can prevent the ______ from discharging one pacemaking stimulus, thus producing a pause of electrical silence for one pacing cycle.

This pause, resulting from one missed SA Node pacing cycle, is sufficient enough to remove the overdrive suppression of an atrial automaticity ______ and...

...the atrial focus escapes to emit a single stimulus; this is an Atrial Escape Beat (on EKG, a pause followed by a P' that differs from the P waves). Then the SA Node quickly resumes pacing, so the atrial focus is ___________-suppressed again.
An unhealthy SA Node that suffers a transient Sinus Block misses one pacing cycle. This pause can induce a Junctional automaticity focus to escape overdrive suppression and emit a **Junctional Escape Beat**.

If the SA Node suffers a transient Sinus Block, it misses one pacing cycle, so a sufficient ______ results and...

... absent any atrial focal response, a Junctional automaticity focus will escape overdrive _________ to emit a ____________ to emit a Junctional Escape Beat.

The depolarization stimulus emitted by the Junctional focus passes down the ventricular conduction _______ to depolarize the ventricles in a normal fashion, so a normal QRS complex results. Then the SA Node resumes pacemaking, overdrive-suppressing the Junctional focus.

**Note:** A single Junctional Escape Beat may produce retrograde atrial depolarization that records an inverted P' immediately before the QRS or an inverted P' after the QRS.
A **Ventricular Escape Beat** originates in a ventricular automaticity focus that is no longer overdrive-suppressed by regular pacing stimuli from above. A ventricular focus typically produces this enormous ventricular (QRS) complex.

A ventricular automaticity _____ can escape overdrive suppression when it is not stimulated by pacemaking activity from above for at least one – maybe two cycles.

It seems a little unusual that the SA Node as well as all of the atrial foci and all the Junctional _____ would fail simultaneously. How then, is it that Ventricular Escape Beats are not so rare? Here’s how…

**Note:** Cardiac parasympathetic innervation inhibits the SA Node and also inhibits the atrial and Junctional foci (see illustration, page 58), but not the ventricular foci. Therefore, a burst of excessive parasympathetic activity depresses the SA Node (producing a pause) and also depresses the atrial and Junctional foci, which leaves only the ventricular foci to respond to the pause. So a ventricular automaticity focus escapes overdrive suppression and discharges, depolarizing the ventricles, producing an enormous ventricular complex. Such a burst of excessive parasympathetic activity is usually transient, so the SA Node resumes its pacemaking activity.

**Note:** Please study the organized review of *Escape* on page 337, with a focus on understanding.
Premature Beats

Premature Beat - an irritable focus spontaneously fires a single stimulus:

- Premature Atrial Beat
- Premature Junctional Beat
- Premature Ventricular Beat

A premature beat (premature stimulus) originates in an irritable automaticity focus that fires spontaneously, producing a beat (on EKG we see evidence of a depolarization) earlier than expected in the rhythm.

**Note:** Those things that make you irritable can do the same to an atrial or Junctional automaticity focus. Quickly peek at the next page and you'll see.

A premature beat, like a premature baby, appears earlier than ________.

When we see a premature beat, we recognize that it was fired by an irritable automaticity _______, so we need to identify the focus (atrial, Junctional, or ventricular).

**Note:** Ventricular automaticity foci are the world's most sensitive O₂ sensors. When they sense low O₂, they become irritable... and they react!

**Note:** Premature beats can cause peculiarities in the rhythm that may mimic more serious problems such as pathological conduction blocks. While some premature beats are not serious, others are a dire warning – we’ll explore them all. You should be cautious and know the difference – lives will depend on it! Understanding the basics provides answers, and understanding facilitates rapid judgement.
Atrial and Junctional foci become irritable because of:

- adrenaline (epinephrine) released by adrenal glands
- increased sympathetic stimulation*
- presence of caffeine, amphetamines, cocaine, or other $\beta_1$ receptor stimulants
- excess digitalis, some toxins, occasionally ethanol
- hyperthyroidism
  (direct stimulation plus heart oversensitive to adrenergic stimulants)
- stretch
  ... and to some extent, low $O_2$

* decreasing or blocking parasympathetic effects may accomplish this.

An automaticity focus in the atria or in the AV Junction may become irritable and spontaneously fire an impulse or even suddenly pace very fast. The cause of irritability in atrial and Junctional foci is usually adrenergic substances (page 57).

Should an atrial or Junctional automaticity focus become irritable,* it may fire a spontaneous impulse that depolarizes the surrounding tissue, so we can recognize it on ____ as a premature beat. EKG

But a very __________ atrial or Junctional focus may fire irritable series of rapid pacing impulses to become the dominant pacemaker, overdrive-suppressing all automaticity centers.

Note: Conditions/substances that can make an atrial focus (usually) or Junctional focus (occasionally) irritable:

- an excess of epinephrine or norepinephrine, the natural substances that activate the adrenergic receptors (of foci).
- adrenergic chemicals that mimic this effect.
- substances or conditions that increase the release of epinephrine or norepinephrine.

* Recalling an irritable person who suddenly yelled at you (too much adrenaline, or maybe too much coffee), you will remember that upper level foci can also become "irritable" (same causes) and spontaneously fire a stimulus.
A **Premature Atrial Beat** (PAB) originates suddenly in an *irritable* (see previous page) atrial automaticity focus, and it produces a P' wave earlier than expected. On EKG, P' is atrial depolarization by an automaticity focus.

A Premature Atrial Beat (PAB) originates in an irritable atrial automaticity focus that spontaneously fires a depolarization stimulus earlier than the normal ___ wave on EKG.

But because an atrial focus is the origin of this premature atrial depolarization (not the SA Node), the stimulus produces a premature and unusually shaped P' wave* that does not look like a normal (Sinus-generated) P _____.

**Note:** On EKG, a PAB records as a P'. The P' may be difficult to detect when it's hiding on the peak of a T wave; the giveaway is a too-tall T... taller than the other T waves in the same lead.

**Note:** Each PAB depolarizes the SA Node; the effect of this... (next page)

* Atrial depolarization from a focus near the SA Node produces a generally upright P' wave whereas a focus in the lower atrium depolarizes the atria in a "bottom-upwards" (retrograde) fashion to record an inverted P' wave in most leads.
An active automaticity center "resets" to continue pacing one cycle length from a Premature Stimulus

Pacing resets to begin in step with* the premature stimulus
*one cycle length after

All centers of automaticity reset, a characteristic of automaticity. A center of automaticity resets its rhythm when it is depolarized by a premature stimulus, so its pacemaking activity resets in step with the premature beat. Observing the illustration from left to right, helps clarify the concept.

Resetting occurs when the dominant automaticity center (usually the SA Node) is depolarized by a __________ beat, then... premature

... its pacemaking activity resets in step with the premature stimulus, so that the next pacing stimulus that it generates is one cycle length from the premature ______.

If the SA Node is depolarized by a premature beat, the SA Node pacemaking is reset, so that regular pacing resumes one cycle length from the ________ stimulus.

Note: In order to reset, the dominant (active) center of automaticity must be depolarized by the premature beat. When there is a premature stimulus that does not reach the dominant pacing center, its pacing is not reset.
A Premature Atrial Beat from an irritable atrial automaticity focus produces a too-early depolarization of the atria that depolarizes the SA Node as well. So the SA Node resets its rhythm in step with the Premature Atrial Beat (P').

**Note:** The P' on EKG is the funny-looking atrial depolarization wave produced by an automaticity focus. It appears different from all SA Node-generated P waves in the same EKG lead, but a normal QRS follows.

If a Regular Sinus Rhythm produced by the SA _____ is interrupted by a spontaneous Premature Atrial Beat (from an atrial automaticity focus), the SA Node, which lies within the atria, is depolarized as well so...

... the SA Node resets, making the P' the first beat of its (reset) ______. The “?” in the illustration marks where the P wave would have occurred, if the SA Node weren’t reset.

**Note:** The reset rhythm of the SA Node resumes the same rate (same cycle length) as before the premature stimulus, but it continues one cycle length from (i.e., in step with) the P'. The pacing rate of the SA Node before and after the PAB remains the same.

**Note:** In reality, the first cycle after a PAB is a little lengthened due to a transient (baroreceptor) parasympathetic effect on the SA Node, which resumes pacing during systole. (Understanding the mechanism is not important.)
The ventricular conduction system is usually receptive to being depolarized by a Premature Atrial Beat, but one Bundle Branch may not have completely repolarized (that is, it's still a little refractory) when the other is receptive. This "aberrant ventricular conduction" produces a slightly widened QRS for that premature cycle only.

**Note:** When a Premature Atrial Beat (P') is conducted to the ventricles, the ventricles are also depolarized earlier than usual.

Sometimes a Premature Atrial Beat can produce aberrant ventricular conduction, because one of the Bundle Branches is not completely repolarized, and therefore is temporarily refractory to depolarization.

So, depolarization of one ventricle is immediate, while depolarization of the other ventricle is slightly delayed.

The non-simultaneous depolarization of the ventricles records as a slightly widened QRS complex after the P' on the EKG. Normal ventricular conduction resumes with normal cycles.
At times, the AV Node is completely unreceptive to a premature atrial depolarization stimulus because it reaches the AV Node prematurely, that is, while the AV Node is still in the refractory period of its repolarization. This results in a "non-conducted" (to the ventricles) Premature Atrial Beat.

A Premature Atrial Beat may be unable to depolarize the AV Node if it (the AV Node) is not fully repolarized and still __________ to an extra-stimulus.

On EKG, this records as a too-early, unusual ___ wave that has no ventricular (QRS-T) response.

**Note:** Warning! Although a "non-conducted" PAB (on EKG, a premature P wave without a QRS response) does not depolarize the ventricles, it does depolarize the SA Node, which resets its pacemaking one cycle length after the premature stimulus. The combination of reset pacing plus the missing QRS-T creates a harmless, but dangerous-looking, span of empty baseline... which has the sinister appearance of a "some-kind-of-block." And one day you will have the satisfaction of correcting someone who guessed the wrong diagnosis.
Occasionally, an irritable automaticity focus fires a Premature Atrial Beat (P') that couples to the end of a normal cycle, and repeats this process by coupling a PAB to the end of each successive normal cycle. This is **Atrial Bigeminy**.

**Note:** The cycle containing the premature beat together with the cycle or cycles to which it couples, is called a “couplet.”

When an irritable atrial focus repeatedly couples a PAB to the end of each (otherwise normal) cycle, this is a run of ______ Bigeminy.* Atrial

Sometimes, an irritable atrial focus may prematurely fire after two normal cycles; when this couplet ________ continuously, repeats it is a run of **Atrial Trigeminy**.

**Note:** With both Atrial Bigeminy and Atrial Trigeminy, each premature stimulus (from the irritable atrial focus) depolarizes the SA Node and resets it, producing a span of clear baseline between the couplets. So a series (“run”) of couplet groups called “group beating” is often seen with Atrial Bigeminy, Atrial Trigeminy, etc. Just look for the premature (P') beat in each couplet. It’s that simple! This is mentioned because group beating may occur with a type of AV conduction block to be discussed later (page 180).

* As you may have noticed, there is a slightly widened (aberrant) QRS after each P' in the upper tracing. Aberrant ventricular conduction can occur after any premature atrial (or Junctional) beat.
Can you determine what is occurring on each of these practice EKG tracings?

Tracing A:
This tracing is from a medical student who had a few cups of coffee in order to study late. She went to the Emergency Room because her pulse seemed irregular.
The intern on duty thought that the tracing showed "intermittent complete AV Block" and was about to call the attending physician (at 4:00 am) to schedule an emergency artificial pacemaker implantation. Explain the EKG strip to the intern using only what you have learned so far (before he wakes the attending physician and discovers the real meaning of "irritable").

Tracing B:
This transmitted telemetry tracing is from a known drug abuser who took a large quantity of amphetamines before his emergency ride to the hospital.
Someone in the ambulance suggested what sounded like "Winky bok block," when the telemetry was transmitted. Utilizing only what you have read and understand so far in this book, you will recognize things that you have just learned. Notice that in each grouping only two of the P waves are identical. Carefully analyze what you see, so you can explain it to others.

**Note:** Carefully examine each tracing and contemplate its subtle information. The answers will appear as you continue reading... somewhere.
A Premature Junctional Beat (PJB) occurs when an irritable automaticity focus in the AV Junction suddenly fires a premature stimulus that is conducted to, and depolarizes, the ventricles (and sometimes the atria, in retrograde fashion).

When an irritable focus (see page 123) in the AV Junction spontaneously fires a stimulus, this produces a Premature Junctional Beat on EKG.

Note: After heart tissue depolarizes, it immediately repolarizes, and during repolarization that tissue is refractory to another stimulus (premature stimulus). As the ventricles repolarize, one Bundle Branch may repolarize slower than the other. So the too-early depolarization from a PJB may conduct through one Bundle Branch, but the impulse is temporarily delayed in the other, still refractory, Bundle Branch (usually the Right). So, instead of depolarizing simultaneously, one ventricle depolarizes punctually and the other is delayed, producing a slightly widened QRS complex typical of a Premature Junctional Beat with aberrant ventricular conduction.

If you see a premature QRS complex that is slightly widened, you should consider that it may be due to a Premature Junctional (or Premature Atrial) Beat with aberrant ventricular conduction.

Previous page answers: A. Non-conducted PAB. B. Atrial Trigeminy with non-conducted PAB.
A Premature Junctional Beat originates in an irritable Junctional focus within the AV Node. We expect such a premature stimulus to conduct to the ventricles, but it may also depolarize the atria in a bottom-up “retrograde” fashion that records as an inverted P' wave in EKG leads with an upright QRS.

Since atrial and ventricular depolarization move in opposite directions from the Junctional focus, the premature P' wave is __________ i.e., opposite the upright QRS.

If a PJB produces retrograde atrial depolarization, it may record an inverted P' wave immediately before the premature _____ complex.

Sometimes an inverted ___ wave associated with a PJB follows the QRS. Occasionally the P' disappears within the QRS when atrial and ventricular depolarization occur simultaneously (not shown in illustration).

**Note:** Retrograde atrial depolarization from a PJB usually depolarizes the SA Node as well. So the SA Node pacing is reset in step with the retrograde atrial depolarization.
(AV) Junctional Bigeminy

An irritable focus in the AV Junction may initiate a Premature Junctional Beat after each normal (SA Node-generated) cycle. This is Junctional Bigeminy. When a PJB is coupled with two consecutive, normal cycles in a repeating series of these couplets, this is Junctional Trigeminy.

An irritable Junctional automaticity focus may fire a premature stimulus coupled to the end of each normal (SA Node paced) cycle to produce Junctional Bigeminy.

An irritable Junctional focus may fire a stimulus after two consecutive, normal Sinus-generated cycles. A repeating series of these couplets is Junctional Trigeminy.

Note: Don’t forget that on EKG you may see an inverted (retrograde) P wave (arrows in upper tracing) with every PJB in either Junctional Bigeminy or Junctional Trigeminy. Also, the SA Node will reset its pacing with each retrograde atrial depolarization; this can produce alarming (but innocent) gaps of empty baseline between couplets.
A ventricular focus can be made irritable by:

| Low O₂ | Airway obstruction  
|        | Absence of air  
|        | (near-drowning or suffocation)  
|        | Air with poor O₂ content  
|        | Minimal blood oxygenation in lungs  
|        | (pulmonary embolus or pneumothorax)  
|        | Reduced cardiac output  
|        | (hypovolemic or cardiogenic shock)  
|        | Poor to absent coronary blood supply  
|        | (coronary insufficiency or infarction)  

| Low K⁺ | Reduced serum potassium ("hypokalemia")  

| Pathology | Mitral Valve Prolapse, stretch, myocarditis, etc.  

... and to a lesser degree, epinephrine-like substances  
(β, adrenergic stimulants).

A ventricular focus may become irritable from under-oxygenation ("hypoxia") due to various circumstances and conditions. Hypokalemia, QT-prolonging medications, mitral valve prolapse, cardiac pathology, and stretch, can do the same.

Poor oxygenation (hypoxia) can make a ventricular automaticity become irritable and fire a spontaneous impulse, producing a premature ventricular beat on EKG.

A very irritable ventricular focus may be so excessively provoked by hypoxia or "ischemia" (diminished blood supply) that it suddenly fires a series of rapid impulses, overdrive-suppressing the normal Sinus rhythm...

... so it becomes the heart's dominant pacemaker.

Note: If you study the illustration for a moment, you will quickly realize that there are numerous mechanisms that can reduce the oxygen supply to these sensitive ventricular automaticity foci. In a clinical setting, most (but not all) "deadly" ventricular tachycardias are due to coronary insufficiency or infarction. Know the other causes of ventricular focal irritability (see illustration).

Note: Cocaine is known to make atrial and Junctional foci irritable, but it has more dangerous effects. Cocaine causes coronary spasm, making ventricular foci hypoxic **and** very irritable; dangerous ventricular arrhythmias may ensue.
A premature ventricular beat called a **Premature Ventricular Contraction** (PVC) originates suddenly in an *irritable* ventricular automaticity focus and produces a giant ventricular complex on EKG.

An irritable (quickly review the previous page, please) ventricular focus may suddenly fire a stimulus and produce a ___________ Ventricular Complex* (PVC) on EKG. **Premature**

**Note:** PVC’s occur early in the cycle. Easily recognized by their great width and enormous amplitude (height and depth), they are usually opposite the polarity of the normal QRS’s (e.g., if QRS’s are upward, PVC’s are mainly downward).

The most likely reason for a ventricular automaticity focus to become irritable is under___________ (hypoxia). **oxygenation**

**Note:** PVC denotes a ventricular “contraction.” When you see a PVC, remember that this represents a (premature) ventricular contraction, and an associated premature pulse beat, albeit weaker than normal (the prematurely stimulated ventricles are not completely filled).

* PVC may stand for Premature Ventricular “Contraction” or “Complex.”

This issue remains unresolved.
The PVC originates in an automaticity focus within the ventricular conduction system, usually in a ventricular wall. Thus, one area of the ventricular wall begins to depolarize before the rest of the ventricle, long before the other ventricle depolarizes.

**Note:** After a normal, Sinus-generated depolarization stimulus has passed through the AV Node, the stimulus is quickly transmitted to the entire endocardial lining of both ventricles at once. This simultaneous depolarization of both ventricles produces a slender QRS complex.

**Note:** When an irritable ventricular automaticity focus suddenly fires an impulse, that ventricular region depolarizes before the rest of the ventricle, and then the depolarization wave creeps to the other ventricle, which then depolarizes... producing an enormously wide ventricular complex.

**Note:** Normally, ventricular depolarization passes through the entire thickness of both ventricles at once. Left ventricular depolarization in the leftward direction tends to be counterbalanced by the simultaneous right ventricular depolarization in the opposite direction. This minimizes the QRS amplitude. But depolarization originating in a remote ventricular focus (as with a PVC), gradually spreads without simultaneous opposition from the other side, and in its slow course, produces (unopposed) deflections of immense amplitude.
The PVC is an enormous ventricular complex that is much wider, taller, and deeper than a normal QRS. There is a pause after the PVC, but it is not caused by resetting of the SA Node; in fact, sometimes you can see the punctual, but ineffective, P wave within the PVC (see P in the illustration).

The PVC is a gigantic ventricular complex that jumps out at you from the EKG, warning you that there is a ventricular focus that is irritable, usually because of __________. hypoxia

PVC’s depolarize only the __________, not the SA Node, ventricles
so the SA Node discharges on schedule. In fact, by measuring P-P cycles, you can often locate the punctual P wave within a PVC.

But that timely P wave occurs while the ventricles are still refractory (from the PVC) and not fully __________. repolarized
When this normal stimulus arrives, they can’t depolarize...

...so there is a ______ as the ventricles finish repolarizing, pause*
making them receptive to the next Sinus-generated cycle.

Note: Interpolated PVC’s are rare, but are somehow sandwiched between the beats of a normal rhythm, producing no pause and no rhythm disturbance.

* The pause, sometimes called a “compensatory” pause, doesn’t “compensate” for anything.
Multiple PVC's from an irritable focus

Numerous PVC’s may emanate from the same ventricular focus, warning that the focus is very irritable, usually because of its poor state of oxygenation. Six or more PVC’s per minute is considered pathological.

While monitoring a given lead, you notice identical PVC's appearing quite often; since each PVC is identical, we know that they all originated in the same irritable focus. (These are “unifocal” PVC’s.) Frequent unifocal PVC’s usually indicate poor oxygenation of a single ventricular focus – often because the blood supply to that focus is diminished. Remember, six PVC’s per minute is pathological. Don’t ignore this patient!

Note: There are situations when the coronary blood flow is adequate, but the blood is poorly oxygenated (e.g., drowning, pneumothorax, pulmonary embolus, tracheal obstruction, etc.). When a highly irritable ventricular focus is warning you with multiple PVC’s… you must respond! Low serum potassium, as well as certain medications, can also irritate a ventricular focus. In addition, adrenergic stimulants like epinephrine can aggravate the situation.
A very irritable ventricular automaticity focus may fire a stimulus that couples to one or more normal cycles to produce **Ventricular Bigeminy**, or **Ventricular Trigeminy**, etc.

**Note:** By convention, 6 PVC’s per minute is considered pathological. A continuous run of any of these couplet patterns quickly exceeds that criterion and usually indicates that a very irritable focus is hypoxic and calling for help.

When a PVC becomes coupled to a normal cycle, and the pattern continues with every cycle in succession, we identify this as Ventricular Bigeminy.

If you see a repetitive pattern of a PVC coupling with every two normal cycles, this is a run of Ventricular Trigeminy.

**Note:** Ventricular automaticity foci are the heart’s hypoxia early warning system. Respond!
**Ventricular Parasystole** is produced by a ventricular automaticity focus that suffers from entrance block (but is not irritable). The parasystolic focus is not vulnerable to overdrive suppression, so it paces at its inherent rate, and the ventricular complexes that it generates poke through the dominant Sinus Rhythm.

**Note:** A solitary ventricular focus suffering from entrance block is "parasystolic", that is, it can't be overdrive-suppressed yet it can deliver pacing stimuli at its inherent rate.

**Note:** A parasystolic ventricular focus suffers from entrance block, insulating it from depolarization by outside sources. Absent overdrive suppression, it paces at its inherent rate. The result is a dual rhythm with pacing from two sources, the SA Node and the ventricular focus.

When you see PVC's that appear to be coupled to a long series of normal cycles, you should suspect Ventricular ________ Parasystole.

**Note:** Because this represents two unrelated, independent rhythms (from two different pacing locations), the interval between the normal cycle and the large ventricular complex is not always consistent. Occasionally a large ventricular complex may fail to appear because the ventricular focus happens to discharge during the refractory period of the (Sinus-paced) ventricles.
A very irritable ventricular focus can emit consecutive stimuli

One PVC

Run of 3 PVC's (VT)

Run of 6 PVC's (VT)

An irritable ventricular automaticity focus may fire once or, if extremely irritable (under-oxygenated), it may fire a rapid series of impulses to produce a run of PVC's.

A single irritable ventricular focus may fire one spontaneous impulse, but if its oxygen supply decreases further, the focus may be provoked into firing a series of discharges in rapid succession.

Of course, runs of PVC's that emanate from an irritable ventricular focus are more serious than occasional PVC's from the same focus, particularly in patients suffering an acute myocardial infarction.

Note: A run of three or more PVC's in rapid succession is really a run of Ventricular Tachycardia (VT). Two of the examples in the above illustration are VT. If a run of VT lasts longer than 30 seconds, it is called "sustained" VT.
Each irritable focus produces its own distinctive PVC

Severe cardiac hypoxia can cause **Multifocal PVC’s** – a desperation measure produced by multiple, exceptionally irritable (hypoxic), ventricular foci. Each focus produces its own unique, identifiable PVC every time it fires.

In a given lead, PVC’s originating in a specific ventricular focus all appear the ______.

**Note:** Severe cardiac hypoxia can cause numerous **multifocal PVC’s** to appear. This is indeed dangerous and requires immediate intervention. Because a single irritable ventricular focus can suddenly fire a series of rapid discharges to produce a dangerous tachy-arrhythmia (e.g., Ventricular Tachycardia), the presence of numerous multifocal PVC’s means that a number of extremely irritable foci are discharging, and trouble is imminent. The chance of developing a dangerous or even deadly arrhythmia (e.g., Ventricular Fibrillation) under these dire circumstances is obviously enhanced. With infarction patients, this is an alarm of crisis proportions!
Mitral Valve Prolapse (MVP) causes PVC's, including runs of VT and multifocal PVC's, yet it is considered a benign condition. With MVP the mitral valve is "floppy" and billows into the left atrium during ventricular systole.

**Note:** First described by Dr. J. B. Barlow in 1968 (MVP is also called "Barlow Syndrome"), this is quite common; 6% to 17% of females, and about 1.5% of males. Females with MVP typically have a slender body with a slight chest deformity, experience "dizzy" spells, and are anxiety prone. They first experience symptoms after age 20.

During ventricular systole, the billowing valves pull on the chordae that tether them to the papillary muscles in the left ventricle. In the author’s opinion, this traction on the papillary muscles causes localized stretch and ischemia, irritating adjacent ventricular automaticity.

**Note:** MVP patients usually have a mid-systolic click with a decrescendo murmur on auscultation.
If a PVC falls on a T wave... watch this patient closely.

If a PVC falls on a T wave ("R on T phenomenon"), particularly in situations of hypoxia or low serum potassium, it occurs during a "vulnerable period" and dangerous arrhythmias may result. Notice how a PVC hits the second T wave directly in its vulnerable period (ouch!)... and see what happens!

PVC’s are, of course, premature and usually occur just after the ____ wave of a normal cycle.

When a PVC falls on the peak of a T wave or on the initial part of its downslope, it catches the ventricles during a vulnerable period, particularly in the presence of _______ (often caused by cardiac ischemia from a narrowed coronary artery) or in the presence of low potassium.

Note: Repolarization of Purkinje fibers (along with their vulnerable period) extends beyond the T wave, so a PVC falling just after the T wave may, in fact, occur during the vulnerable period of ventricular foci. Ischemia can extend Purkinje repolarization even further.

Note: This well known warning sign, "R on T" is often noted after the fact, during the review of an EKG strip from a patient who suffered a dangerous or deadly arrhythmia. By being cautious and vigilant, you can respond quickly.
The discerning eye of a coronary care nurse detected a beat that appeared a little too early on the EKG strip taken from a patient’s monitor. Let’s determine the location of the irritable focus that produced the premature beat.

The last QRS complex in the strip catches your eye because it occurs prematurely, and it is not preceded by a ___ wave.  

The last QRS complex looks the same as the other QRS’s, so we know that, although premature, the last QRS resulted from depolarization that passed (in a normal fashion) down the ventricular conduction system. Therefore, it is not from a ___________ focus.  

Carefully examining the EKG strip, we don’t see a P’ (with a little baseline) before the premature QRS, so we know that the QRS did not emanate from an atrial focus. Therefore the irritable automaticity focus that produced the premature QRS must be in the ___________.  

**Note:** Sure you understand this, but it probably would be a good idea to take a minute to review this section, and study the simplified review of *Premature Beats* on page 337. Then, let’s take a break, and I’ll be right here when you return.
A "tachy-arrhythmia" originates in a very irritable focus that paces rapidly. Sometimes more than one active focus is generating pacing stimuli at once.

**Note:** Tachy-arrhythmia ("rapid arrhythmia"), hyphenated for recognition purposes here, is not usually hyphenated, so henceforth the hyphen will be omitted.

The rate ranges of the tachyarrhythmias are:

- **Paroxysmal Tachycardia** ........... ___ to ___ /minute. 150 to 250
- **Flutter** ......................... ___ to ___ /minute. 250 to 350
- **Fibrillation** ....................... ___ to ___ /minute. 350 to 450

**Note:** A tachyarrhythmia is easily recognized by rate alone, but the specific diagnosis requires that we identify the origin, that is, we must determine the location of the irritable automaticity focus (atrial, Junctional, or ventricular). You already have a solid understanding* of normal conduction in the heart, so we merely need to get up to speed (pun intended) in learning the behavior of very irritable automaticity foci, and how they record on EKG.

* "Understanding is a kind of ecstasy." Carl Sagan (from *Broca's Brain.*)
Paroxysmal (sudden) Tachycardia

a very irritable automaticity focus
suddenly paces rapidly:

- Paroxysmal Atrial Tachycardia
- Paroxysmal Junctional Tachycardia
- Paroxysmal Ventricular Tachycardia

Paroxysmal ("sudden") tachycardia ("rapid heart rate") indicates rapid pacing (250 to 250 per minute) by a very irritable automaticity focus. Once we recognize a paroxysmal tachycardia, we need only identify the focus (atrial, Junctional, or ventricular) of its origin.

The medical term for rapid heart rate is ___________.

Paroxysmal means ________.

sudden

Note: Paroxysmal tachycardia arises suddenly from a very irritable automaticity focus. Generally speaking, stimulants like epinephrine make higher level foci irritable, whereas more threatening physiological conditions like hypoxia (or low potassium) make ventricular foci irritable. There is some overlap, however. In addition, a single premature stimulus from another focus can provoke an irritable focus into a run of paroxysmal tachycardia.

In contrast, Sinus tachycardia is the SA Node's gradual response to exercise, excitement, etc. Although the SA Node's rate of pacing may eventually become quite rapid, Sinus tachycardia is neither sudden nor does it originate in an automaticity focus, so by definition, it is not a ___________ tachycardia.
Paroxysmal Tachycardia

The rate range of the paroxysmal tachycardias is 150* to 250 per minute, so they are easy to recognize. Locating the causative irritable focus (atrial, Junctional, or ventricular) gives us the diagnosis.

When calculating rate, we find an R wave that peaks on a heavy black "start" line. The next three heavy black lines are called "300, 150, ____.”

The fine line immediately to the right of the heavy black "300" line is the thin "250" line. Therefore, if an R wave falls on the "start" line (see illustration) the next R wave will fall within the shaded area during paroxysmal ______________
tachycardia.

You can instantly recognize a paroxysmal tachycardia by noting the rate range of ____ to 250 per minute. Now you have to determine at which of three levels there is a very irritable automaticity focus causing the tachycardia. Easy enough!

* Some authors now set the lower rate limit of paroxysmal tachycardia at 125 per minute.
**Paroxysmal Atrial Tachycardia** (PAT) is caused by the sudden, rapid firing of a very irritable atrial automaticity focus. You may see the beginning of this arrhythmia only occasionally, so become familiar with its general appearance.

Paroxysmal Atrial Tachycardia has a rapid heart rate that originates suddenly in a very irritable focus in one of the ________. The rate range is usually 150 to 250 per minute, so it overdrive-suppresses the SA Node and all other automaticity foci.

Because the origin of this tachyarrhythmia is a very irritable atrial focus, the atrial depolarizations of PAT are ___ waves that do not look like the Sinus-generated P waves.

Each depolarization impulse from the rapidly-pacing, irritable atrial focus depolarizes the atria and then is conducted down the ventricular conduction system to the ventricles, producing normal-appearing P’-QRS-T cycles.

**Note:** A premature stimulus from another focus may set off PAT.

*Quickly review page 123.*
PAT with (AV) Block

- rapid rate, spiked P' waves
- 2:1 ratio of P':QRS

Suspect digitalis excess or toxicity.

*Paroxysmal Atrial Tachycardia with AV block* has more than one P' wave spike for every QRS response. Suspect digitalis excess or toxicity; atrial foci are very sensitive to the irritating effects of digitalis preparations.

**Note:** Excess digitalis can provoke an atrial focus into such an irritable state that it suddenly paces rapidly. At the same time, digitalis markedly inhibits the AV Node, so that only every second stimulus conducts to the ventricles (every-other atrial stimulus is blocked in the digitalis-inhibited AV Node).

PAT with block* is a tachyarrhythmia that has two P' waves for each QRS response on EKG, because the _____ blocks the conduction of every-other atrial stimulus.

PAT with block is usually a sign of digitalis excess or toxicity, particularly if the patient has a low serum potassium, so careful administration of intravenous potassium can help. Also, digitalis antibodies can be employed to reduce toxicity.

* The "AV" is sometimes omitted, but is always understood.
**Paroxysmal Junctional Tachycardia** (PJT) is caused by the sudden rapid pacing of a very irritable automaticity focus in the AV Junction. The Junctional focus may suddenly initiate tachycardia pacing, because of marked irritability induced by stimulants and/or by a well-timed premature beat from another focus.

Paroxysmal Junctional Tachycardia is due to a very irritable* focus in the AV Junction that paces at the rate of 150 to 250 per minute.

**Note:** A rapidly pacing (irritable) Junctional focus may also depolarize the atria from below in retrograde fashion to record:

- an inverted P' immediately before each upright QRS, or (see illustration page 132)
- an inverted P' after each upright QRS, or
- an inverted P' buried within each QRS (difficult to detect).

**Note:** Each stimulus from a rapidly pacing (irritable) Junctional focus may occur at a time in the cycle when the Left Bundle Branch has fully repolarized (i.e., recovered from its refractory period), but the Right Bundle Branch is still refractory (in some patients, the reverse occurs). As a result, this *aberrant ventricular conduction* depolarizes the left ventricle before the right, to produce somewhat widened QRS's during the tachycardia.

*One more look at page 123, and I’ll never bother you again.*
Another type of Junctional Tachycardia is AV Nodal Reentry* Tachycardia (AVNRT). In theory, a continuous reentry circuit develops (which includes the AV Node and the lower atria) and rapidly paces the atria and ventricles.

**Note:** A theoretical “reentry circuit” may continuously circle (like perpetual motion) through the AV Junctional region, giving off a depolarization stimulus to the atria and to the ventricles with each pass in the circuit. This is “circus reentry,” an aptly named tachycardia that looks suspiciously like PJT.

**Note:** In AVNRT, each pacing stimulus first records from an origin near the coronary sinus — an area loaded with automaticity foci. Although the putative reentry circuit includes a broad area around the AV Node, only catheter ablation of the focus-laden region can successfully eliminate this tachycardia (very suggestive of focal automaticity origin). Dogmatic loyalty to this theoretical reentry model persists. The jury is still out.

* Pronounced “ree-EN-tree”.
Supraventricular Tachycardia

The very irritable* automaticity foci that produce both Paroxysmal Atrial Tachycardia and Paroxysmal Junctional Tachycardia originate above the ventricles, so these arrhythmias are known as Paroxysmal Supraventricular Tachycardia.

Supraventricular tachycardia (the word “paroxysmal” is often omitted) is a general term, which includes both PAT and _____.

PJT

The term “supraventricular” imparts the understanding that all atrial and all Junctional foci are above the ____________ ventricles.

Paraoxysmal Atrial Tachycardia can be so rapid that the P waves run into the preceding T waves to become indistinguishable. This can make differentiation between PAT and PJT very difficult. But since treatment for both is so similar, the umbrella term Supraventricular Tachycardia (SVT) suffices, and further distinction between the two is unnecessary. Certain conditions may widen the QRS’s in SVT, so it may then resemble Ventricular Tachycardia (next page).

*Usually an atrial or Junctional focus is made irritable by adrenergic stimulants, but a focus may be further provoked into tachycardia pacing by a premature stimulus from another irritable focus.
Paroxysmal Ventricular Tachycardia (PVT or VT)* is produced by a very irritable ventricular automaticity focus that suddenly paces in the 150 to 250 per minute range. It has a characteristic pattern of enormous, consecutive PVC-like complexes. Please conscientiously review page 134 now.

Paroxysmal Ventricular Tachycardia originates suddenly in a very ________ ventricular automaticity focus, producing a (ventricular) rate of 150 - 250.

Sudden runs of Ventricular Tachycardia* resemble a rapid series of _____’s (which in reality, they are).

Note: During Ventricular Tachycardia, the SA Node still paces the atria, but the large, dramatic ventricular complexes hide the individual P waves that can be seen only occasionally. So, there is independent pacing of the atria and the ventricles… a type of AV dissociation.

* The “Paroxysmal” is often left off, so “Ventricular Tachycardia” or “VT” are used commonly.
During Ventricular Tachycardia, the SA Node continues to pace the atria (AV dissociation), but an occasional atrial depolarization catches the AV Node in a receptive state, and then this depolarization stimulus conducts to the ventricles.

Occasionally (during VT), one of the regular (Sinus-paced) atrial depolarizations finds the AV _____ receptive to depolarization and... Node

that stimulus passes through the AV Node to depolarize the ventricles via the ventricular ___________ system. conduction

Note: On occasion during VT, a (Sinus-paced) depolarization stimulus from the atria finds the entire ventricular conduction system receptive to depolarization and produces a normal-appearing QRS (capture beat) in the midst of the ventricular tachycardia. More commonly during VT, an atrial depolarization finds a receptive AV Node, but ventricular depolarization only proceeds so far before it meets ventricular depolarization progressing from the ventricular focus. This produces a fusion beat, which is a blending on EKG of a normal QRS with a PVC-like complex (see illustration). The presence of “captures” or “fusions” confirms the diagnosis of Ventricular Tachycardia, because they could occur during SVT.
Runs of (Paroxysmal) Ventricular Tachycardia may signify coronary insufficiency (ischemia) or other causes of cardiac hypoxia that make a ventricular automaticity focus very irritable.

Ventricular Tachycardia appears like a run of ____'s.

Paroxysmal Ventricular Tachycardia often indicates coronary __________, causing poor oxygenation insufficiency of the heart (and ventricular foci). For other causes of cardiac hypoxia see page 134.

**Note:** This rapid ventricular rate suddenly erupts from an irritable (hypoxic) ventricular focus. The rapid rate is too fast for the heart to function effectively, particularly in the elderly with compromised coronaries. It should be treated quickly (but cautiously) in patients with a myocardial infarction.

**Caution:** A rapid (Junctional or atrial) Supraventricular Tachycardia with aberrant conduction can produce a tachycardia with widened QRS's that mimics VT. Also, pre-existing Bundle Branch Block with SVT will widen the QRS complexes to give the same impression. NEVER give medications for SVT to a patient with VT.
Distinguishing
Wide QRS complex SVT from Ventricular Tachycardia

<table>
<thead>
<tr>
<th>Helpful Clues</th>
<th>Wide QRS Complex SVT</th>
<th>Ventricular Tachycardia</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patient with coronary disease or infarction</td>
<td>uncommon</td>
<td>very common</td>
</tr>
<tr>
<td>QRS width (duration)</td>
<td>less than .14 sec.</td>
<td>greater than .14 sec.</td>
</tr>
<tr>
<td>AV dissociation showing captures or fusions</td>
<td>rare</td>
<td>yes</td>
</tr>
<tr>
<td>Axis: Extreme Right Axis Deviation (R.A.D., see page 231)</td>
<td>rare</td>
<td>yes</td>
</tr>
</tbody>
</table>

A few clues and good judgement can help you distinguish between VT and wide QRS complex SVT (with aberrant ventricular conduction). Begin with the history, and get a 12 lead EKG.

The patient with VT is most likely elderly and suffering from diminished coronary blood flow, reducing the oxygen supply to the ventricular foci.

Signs of AV dissociation (e.g., presence of fusions or captures, or Extreme R.A.D. (-90° to -180°) are characteristic of VT.

Note: If the QRS complex can be measured with accuracy, the QRS in SVT, even if widened by aberrant ventricular conduction, is usually .14 sec. or less in duration. However the ventricular complexes in VT are very wide, .14 sec. or greater. There are many criteria for distinguishing VT from SVT with aberrancy, but probably the most reliable to date are in:

Torsades de Pointes is a peculiar form of (very) rapid ventricular rhythm caused by low potassium, medications that block potassium channels, or congenital abnormalities (e.g., Long QT syndrome), all of which lengthen the QT segment. The rate is a variable 250 to 350 per minute, usually in brief episodes.

Note: Torsades* de Pointes means “twisting of points,” which refers to the series of ventricular complexes that are upward-pointing then downward-pointing in a repeating continuum. In 1966, Dr. F. Dessertenne presented the first scientific description of this arrhythmia. He theorized that it was caused by two competitive, irritable foci in different ventricular areas – an explanation that seems quite plausible.

The rate of this arrhythmia is 250 to ____ per minute, but fortunately it usually occurs only in brief self-terminating bursts, for at that rate there is no effective ventricular pumping.

Note: On EKG, the amplitude of each successive complex gradually increases and then gradually decreases, so when viewed as a whole, the general outline or silhouette of the tracing looks like a series of end-to-end spindle shapes. Some say the tracing outline resembles a twisted ribbon. If unresolved, it can lead to a deadly arrhythmia.

* This is the correct spelling; don’t forget the “s” at the end of “Torsades”, even though it is not pronounced.
Atrial Flutter originates in an atrial automaticity focus. The rapid succession of identical, back-to-back atrial depolarization waves, “flutter” waves, suggest a reentry pathway as some experts (see the last paragraph on the next page).

In atrial flutter an extremely irritable atrial automaticity focus fires at a rate of 250 to 350 per minute, producing a rapid series of ______ depolarizations.

**Note:** On EKG, atrial flutter is characterized by consecutive, identical “flutter” waves in rapid back-to-back succession. The baseline appears to vanish between the back-to-back flutter waves, and because the waves are identical, they are described as having the appearance of the teeth of a saw or a “saw tooth” baseline. Turn back to PAT with (AV) block to make sure that you recognize the difference.

**Note:** The AV Node has a long refractory period, so only one in a series of flutter waves conducts to the ventricles. Therefore, this very rapid series of atrial depolarizations cannot drive the ventricles at the same excessive rate; perhaps only one of two (or commonly, one of three, as above) atrial depolarizations reaches the ventricles in atrial flutter.
Identification of Atrial Flutter

may require inverting the tracing...

inverted tracing

...or employing a vagal maneuver.

vagal maneuver

flutter waves revealed

Inverting a tracing of suspected atrial flutter can help in its identification. Also, vagal maneuvers can be an effective diagnostic aid (see page 61.)

When in doubt about atrial flutter, inverting the _____ tracing may be helpful.

Note: With atrial flutter there may be a rapid QRS response rate, particularly in 2:1 ratios (flutter waves:QRS response), masking the flutter waves. Vagal maneuvers increase AV Node refractoriness, allowing fewer flutter waves to be conducted to the ventricles. This produces a longer series of flutter waves that are easier to identify.

Note: The “Maze” surgical procedure cuts (and resutures) the atria into a maze of channels that provides a continuous pathway from the SA Node to the AV Node. This procedure eliminates any possibility of reentry circuits. Yet a study of patients recovering from the maze procedure, revealed that 47% developed atrial flutter (or atrial fibrillation) postoperatively. This raises considerable doubt that the origin of atrial flutter could be reentry.
Ventricular Flutter is produced by a single ventricular automaticity focus firing at an exceptionally rapid rate of 250 to 350 per minute. It produces a rapid series of smooth sine-waves of similar amplitude.

Ventricular flutter is caused by a highly irritable ventricular focus that is desperately discharging at a rate of ______ to ______ per minute.

The ventricular rate in ventricular flutter is so rapid that the ________ ventricles hardly have enough time to fill – even partially, so this arrhythmia rapidly deteriorates into a deadly arrhythmia.

The smooth _____-wave pattern of ventricular flutter is its ______ distinguishing characteristic.

Note: Ventricular flutter produces a rapid series of smooth sine-waves of similar amplitude, whereas the waves of Torsades de Pointes get progressively larger, then smaller, producing a general outline of connected spindle shapes (page 158). Ventricular Flutter rarely self-resolves and is nearly always a prelude to a deadly arrhythmia... see next page.
True ventricular flutter almost invariably deteriorates into ventricular fibrillation, which requires immediate Cardio-Pulmonary Resuscitation and defibrillation.

**Note:** During ventricular flutter, the ventricles are contracting at an alarming rate. The above (separated but continuous) tracing shows ventricular flutter at a rate of about 300 per minute, which is five contractions per second. Blood is a viscous fluid, and the ventricles cannot properly fill (and empty) at a rate of 5 times per second; in fact, they hardly fill at all. For this reason, there is no effective cardiac output. Therefore, the coronary arteries are not receiving blood, and the heart itself has no blood supply. Ventricular Fibrillation ensues, as many profoundly hypoxic ventricular automaticity foci desperately try to compensate... in vain.
A monitored patient became very concerned about a sudden pounding in his chest.

By the history and the rate (which you determined by observation), you identify the rhythm as a paroxysmal _________.

Now, we'll determine the causative irritable automaticity focus.

Because this paroxysmal tachycardia has narrow, normal looking QRS's, it could **not** have originated in an irritable _____ focus; therefore it must be some type of supraventricular tachycardia.

There appear to be P' waves present, so we are probably dealing with an automaticity focus in the _____.. You have already ruled out a Junctional focus, because any retrograde depolarizations that it might have produced, would record as inverted P' waves (which are usually adjacent to the QRS when they precede it).

**Note:** This is Paroxysmal Atrial Tachycardia (PAT), and because each P' wave produces a QRS response, it could not be PAT with block. Quickly review the illustrations for the paroxysmal tachycardias and flutter before we go on. Take your time.
**Fibrillation**

multiple foci discharging rapidly

350 - 450 (discharges) /min.

"Fibrillation" is a totally erratic rhythm caused by continuous, rapid rate discharges from numerous automaticity foci in either the atria or in the ventricles.

**Note:** Fibrillation is caused by rapid discharges from numerous profoundly irritable automaticity foci in the atria (**Atrial Fibrillation**), or due to numerous profoundly irritable foci in the ventricles rapidly discharging (**Ventricular Fibrillation**). Both types represent a pathological condition: these *irritable* foci all suffer from entrance block, so they are *parasystolic*. Since they cannot be overdrive-suppressed, they all pace rapidly at once. The resulting rhythm is so erratic and uncoordinated that distinct, complete waves are not distinguishable, thus rates are impossible to determine. The involved chambers merely twitch rapidly.

**Note:** The "rate" 350 to 450 per minute is not a true rate, since many of the foci discharge simultaneously. The number and the tachy-rate of individual foci is conjectural. The range of "rate" is more relative and hypothetical than real, because fibrillating chambers do not effectively pump at all.
Atrial Fibrillation (AF) is caused by many irritable parasystolic atrial foci (with entrance block) firing at rapid rates, producing an exceedingly rapid, erratic atrial rhythm. The atrial "rate" is 350 to 450 per minute. Notice the irregular ventricular response.

Atrial fibrillation occurs when many irritable atrial foci* fire rapidly, but since they are parasystolic, none of them can be overdrive-suppressed; they all rapidly pace at once to produce an excessively rapid series of tiny, erratic spikes on EKG.

Note: Because so many atrial foci are rapidly firing, no single depolarization spreads very far. Only a small portion of the atria is depolarized by any one discharge from an atrial focus. Depolarizations from foci near the AV Node conduct to the ventricles, producing a very irregular ventricular rhythm. See rhythm strip, page 351.

Note: With a Normal Sinus Rhythm, each pacing impulse that the SA Node generates spreads through the atria like an enlarging, circular wave; much like a pebble dropped into a pool of water. However, the multiple erratic depolarizations of atrial fibrillation are analogous to a rain shower striking the same pool.

* Atrial fibrillation is usually initiated by parasystolic foci in the pulmonary vein ostia of the left atrium.
Atrial Fibrillation often appears as a wavy baseline without identifiable P or P' waves. The QRS response is not regular and may be fast or slow.

Atrial Fibrillation may cause such small, erratic spikes that it appears like a wavy baseline without visible __ waves (and without distinguishable P' waves either).

**Note:** Only discharging foci near the AV Node can (occasionally) stimulate it, but the AV Node sorts out a normal ventricular rate.

The AV Node is irregularly stimulated during atrial fibrillation, so the ____________ (QRS) response is irregular. On EKG, you may see only random QRS's (see illustration), so the pulse is irregular also.

**Note:** With Atrial Fibrillation, the ventricular rate depends on the AV Node’s duration of refractoriness after it is stimulated. During AF, the AV Node usually allows a relatively normal range of ventricular rate, albeit always irregular. Sometimes the AV Node permits an increased number of depolarization stimuli to pass through, producing a rapid ventricular rate that may require pharmacological control. Always determine the ventricular (pulse) rate (QRS’s per 6-second strip times 10) and document it. If the ventricular rate is out of a safe range for the patient, it should be treated appropriately.
**Ventricular Fibrillation** (VF) is caused by rapid-rate discharges from many variable, parasystolic ventricular automaticity foci, producing an erratic, rapid twitching of the ventricles (ventricular “rate” is 350 to 450 per minute).

Ventricular Fibrillation is due to numerous parasystolic ventricular foci pacing rapidly (each of them suffering from entrance block, so they can not be __________-suppressed); this produces an __________ overdrive erratic twitching of the ventricles.

Because there so many ventricular ______ firing rapidly, each one repeatedly depolarizes only a small area of ventricle. This results in a rapid, ineffective twitching of the ventricles.

This erratic twitching of VF has been called a “bag of worms,” for this is the way the ventricles actually appear. On EKG the tracing is totally erratic, without identifiable _____, and the ventricles do not provide mechanical pumping. Emergency!
Ventricular Fibrillation is easily recognized by its totally erratic appearance and lack of any identifiable waves on the electrocardiogram.

**Note:** These three strips are from a continuous tracing of the same patient’s dying heart. Notice how the amplitude of the deflections diminishes as the heart dies.

We recognize Ventricular Fibrillation by its completely erratic appearance on the EKG tracing. Even with large deflections, there are no identifiable ____.

There is no predictable pattern with __________ Fibrillation. As you can see, it looks different at every moment, but it is so erratic that it is difficult to miss, thank Goodness!

If you do recognize any repetition of pattern or regularity of deflections, you probably are not dealing with ____.
Ventricular Fibrillation is a type of cardiac arrest, for there is no pumping action by the heart; this is a dire emergency! VF requires immediate CPR and defibrillation, with some type of electrical defibrillator.

Ventricular Fibrillation is a type of cardiac arrest. There is no effective cardiac output, because the ventricles are only twitching erratically. There is no ventricular pumping, so there is no circulation.

**Note:** VF requires immediate defibrillation. Cardiac Arrest is an emergency that demands immediate intervention. Cardio-Pulmonary Resuscitation (cardiac massage and assisted respiration) is carried out in order to circulate oxygenated blood by external mechanical means. The technique of CPR was originally taught only to hospital and ambulance personnel, but it is imperative that every person master this technique. Only when CPR skills are universally known, can all victims of cardiac arrest get immediate lifesaving care at any location.

**Note:** There are two other types of cardiac arrest. Cardiac Standstill (“Asystole”) occurs when there is no detectable cardiac activity on EKG. This is a rare circumstance when the SA Node and the escape mechanisms of all the foci at all levels are unable to assume pacing responsibility. Pulseless Electrical Activity (PEA) is present when a dying heart produces weak signs of electrical activity on EKG, but the moribund heart cannot respond mechanically (no detectable pulse).
There are now computerized defibrillators that can detect Ventricular Fibrillation and immediately deliver a defibrillating shock. One type, the AED, is a portable unit that can be operated by the public. Another type, the ICD, is a small unit that is implanted under the chest skin to automatically defibrillate appropriate patients as needed.

**Note:** The *Automated External Defibrillator* (AED) is a small portable unit. When its electrodes are placed on the chest of an unconscious person, it is programmed to identify VF and deliver a defibrillating shock.

**Note:** The *Implantable Cardioverter Defibrillator* (ICD) is implanted under the chest skin of patients likely to develop Ventricular Fibrillation. Wire leads from the ICD are attached to the heart to detect VF and deliver a defibrillating shock. This mini-computer can also identify other arrhythmias and treat them with timed electrical stimuli, and it can pace if a bradycardia ensues. A technological wonder!

Please review all “fibrillation” illustrations.
An abnormal, accessory AV conduction pathway, the *bundle of Kent*, can "short circuit" the (usual) delay of ventricular conduction in the AV Node. This prematurely depolarizes ("pre-excites") a portion of the ventricles (producing a delta wave on EKG) just before normal ventricular depolarization begins.

The accessory Bundle of ______ causes ventricular pre-excitation in Wolff-Parkinson-White (WPW) syndrome.

The delta wave creates the illusion of a "shortened" PR interval and "lengthened" QRS. The delta wave actually records the depolarization of an area of ___________ pre-excitation.

**Note:** WPW syndrome is very important because persons with such an accessory pathway can have paroxysmal tachycardia by three possible mechanisms:

- rapid conduction — supraventricular tachycardia (including atrial flutter or atrial fibrillation) may be rapidly conducted 1:1 through this accessory pathway producing dangerously high ventricular rates.

- some Kent Bundles have been found to contain automaticity foci that can initiate a paroxysmal tachycardia.

- re-entry — ventricular depolarization may immediately restimulate the atria in a retrograde fashion via the accessory pathway causing a theoretical circus re-entry loop.
In patients with **LGL syndrome**, the AV Node is bypassed by an extension of the Anterior Internodal Tract. Absent the conduction delay in the AV Node, this "James bundle" conducts atrial depolarizations directly to the His Bundle without delay. This can pose a serious problem with rapid atrial arrhythmias like atrial flutter.

Ordinarily the AV Node filters rapid supraventricular rates, in order to transmit depolarization to the __________ at a physiologically reasonable rate.

Absent the filtering effect of the AV Node, patients with LGL syndrome can transmit rapid atrial rates directly (1:1) to the His Bundle, driving the ventricles at very _____ rates.

With this syndrome, the AV Node is bypassed by the James tract, so there is no significant PR interval delay; the P ______ are adjacent to their QRS’s on the EKG.

**Note:** You may now review *Rhythm, Part I* by turning to the Personal Quick Reference Sheets on page 336-338, and relate this to the simplified methodology that is summarized on page 334.
Chapter 6: Rhythm, Part II

Before you begin, look at this chapter's summary on pages 334 and 339.

Rhythm, Part II

Blocks

- Sinus Block
- AV Block
- Bundle Branch Block
- Hemiblock
  (begins on page 295, Chapter 9)

Blocks retard or prevent the conduction of depolarization; they can occur in the SA Node, the AV Node, or in the larger divisions of the ventricular conduction system. The general public and the media often refer to them as "heart blocks".

Blocks may develop in any of these areas: the SA Node, the AV Node, the His Bundle, the Bundle Branches, or in either of the two subdivisions of the Left Bundle Branch (Hemiblock).

These are blocks of electrical conduction that prevent (or retard) the passage of _____________ stimuli. depolarization

Note: When examining the rhythm on a tracing, always check for all the varieties of block, because the same patient can have more than one type of block.
An unhealthy SA Node (Sinus Node) may temporarily fail to pace for at least one cycle, this is **Sinus Block**, then the SA Node resumes pacing. Notice that the missed cycle has no P wave; a very important feature.

With **Sinus Block** (also called “**SA Node Block**” or simply “**SA Block**”), an unhealthy SA Node* stops its pacing activity for at least one complete ______, so the block is usually transient.

After the pause of Sinus Block, pacing resumes at the same rate (and timing) as prior to the block. The SA Node resumes its pacing responsibility in step with its previous rhythm. However, the pause may induce an escape beat from an automaticity ______.

**Note:** The P waves before and after the pause are identical, since they originate in the SA Node. The SA Node continues to generate atrial depolarizations with the same timing as before the block. However a long pause may elicit an escape beat from an automaticity focus before the SA Node can resume pacing (see pages 119-121).

* Some experts claim that the SA Node does generate a stimulus, but that it is blocked from leaving the SA Node. This is referred to as **Sinus “Exit” Block**.
Sick Sinus Syndrome (SSS) is a wastebasket of arrhythmias caused by SA Node dysfunction associated with unresponsive supraventricular (atrial and Junctional) automaticity foci, which are also dysfunctional and can’t employ their normal escape mechanism to assume pacing responsibility.

**Note:** Sick Sinus Syndrome most often occurs in elderly individuals who have heart disease. It is usually characterized by marked Sinus Bradycardia, but without the normal escape mechanisms of atrial and Junctional foci. SSS may also present as recurrent episodes of Sinus Block or Sinus Arrest associated with faulty (or absent) escape mechanisms of all supraventricular foci.

**Note:** Because of the exclusive parasympathetic innervation to the SA Node and all supraventricular (atrial and Junctional) foci, excessive parasympathetic activity depresses the pacing rate of the SA Node, and depresses the atrial and Junctional foci as well. Therefore, young, healthy individuals (e.g., conditioned athletes like marathon runners) who often have parasympathetic hyperactivity at rest, appear to exhibit convincing signs of SSS ("pseudo" Sick Sinus Syndrome).

**Note:** Patients with SSS may develop intermittent episodes of SVT (sometimes even Atrial Flutter or Atrial Fibrillation) mingled with the Sinus Bradycardia. This is Bradycardia-Tachycardia Syndrome.
AV Block

1° (first degree) AV Block

2° (second degree) AV Block

3° (third degree) AV Block

AV (Atrio-Ventricular) blocks either retard or eliminate (or both!) conduction from the atria to the ventricles.

Minor AV blocks lengthen the brief pause between atrial depolarization and ventricular depolarization.

Most AV blocks completely block some (or all) supraventricular impulses from reaching the ventricles.

Note: The AV blocks are:

• **first degree (1°) AV block** (lengthens the delay between atrial and ventricular depolarization)

• **second degree (2°) AV block** (Wenckebach and Mobitz types)

• **third degree (3°) AV block** (completely blocks the conduction of atrial stimuli to the ventricles)

Note: Whether “first degree” is written out or the shorthand notation, 1°, is used, both have the same meaning. This book will alternately use both methods for all degrees of AV block, since you will see both in the current literature. Although it is currently popular to omit the “AV” it is always understood.
First degree (1°) AV block retards AV Node conduction, prolonging the PR interval more than one large square (.2 sec.) on EKG.

**Note:** Technically, a “segment” is a portion of baseline, while an “interval” contains at least one wave. So the PR interval includes the P wave and the baseline that follows, up to the point where the QRS complex begins. Therefore, the PR interval is measured from the beginning of the P wave to the beginning of the QRS complex.

The delay caused by 1° AV block prolongs the ___ interval. \[\text{PR}\]

The PR interval normally should be less than one large square, which is less than ___ second. \[.2 \text{ (2/10)}\]

**Note:** You must observe (and record) the PR interval for every EKG. Some kind of AV block is present, if the PR interval is longer than one large square anywhere on an EKG.
1° AV Block

"Measure" PR by observation (one large square).

PR remains consistently lengthened cycle-to-cycle.

A first degree (1°) AV block is characterized by a PR interval greater than .2 second (one large square). The PR prolongation is consistent in every cycle.

Once you recognize a prolonged PR _________, you should determine the type of AV block that is present.

Some type of AV block is present if any PR interval is longer than ___ second, anywhere in the tracing.

A 1° block* is present when the PR interval is consistently prolonged the same amount in every _____, and the P-QRS-T sequence is normal in every cycle also.

* Whenever you hear “1° block,” understand that it means “1° AV block.”
Second degree AV blocks allow some atrial depolarizations (P waves) to conduct to the ventricles (producing a QRS response), while some atrial depolarizations are blocked, leaving lone P waves without an associated QRS. There are two general types of 2° AV block; those that occur in the AV Node, and those that occur below the AV Node.

Note: There are two types of 2° AV blocks:

- 2° blocks of the AV Node are “Wenckebach” (formerly called “Type I”). They produce a series of cycles with progressive blocking of AV Node conduction until the final P wave is totally blocked in the AV Node, eliminating the QRS response. Each repeating Wenckebach series has a consistent P:QRS ratio like 3:2, 4:3, 5:4, etc. (one less QRS than P waves in the series).

- 2° blocks of Purkinje fiber bundles (His Bundle or Bundle Branches) are “Mobitz” (formerly called “Type II”). They usually produce a series of cycles consisting of one normal P-QRS-T cycle preceded by a series of paced P waves that fail to conduct through the AV Node (no QRS response). Each repeating Mobitz series has a consistent P:QRS ratio, like 3:1, 4:1, 5:1, etc.

Note: Don’t be intimidated by these descriptions, once you see the tracings on the next few pages, you will understand immediately.

* Pronounced “WINKY-bok”
Wenckebach $2^\circ$ AV block occurs in the AV Node. On EKG, the PR interval gradually lengthens in successive cycles, but the last P wave of the series fails to conduct to the ventricles (the final P lacks a QRS response). This series repeats.

On EKG, Wenckebach ($2^\circ$ AV block) gradually prolongs the PR interval in each successive cycle, until the final P wave of the series fails to produce a ____ response ("dropped QRS").

Each P wave and its associated QRS get progressively farther apart in successive cycles; the last P wave stimulus (totally blocked in the AV Node) stands alone at the end of the series. This typical Wenckebach pattern ("footprint") consists of anywhere from two to eight or more _____.

**Note:** Wenckebach is usually located in the AV Node. Wenckebach is sometimes caused by parasympathetic excess (inhibits the AV Node) or drugs that mimic or induce parasympathetic effects. Carefully examine EKG’s for this characteristic, progressive lengthening of the PR in consecutive cycles, ending in a final lone P wave (see page 329). Repeating short series of Wenckebach footprints can produce "group beating" that looks somewhat like couplets of premature beats. Don’t be fooled.
“Mobitz” 2º AV Blocks

2:1 Mobitz AV block

3:1 Mobitz AV block

Mobitz 2º (AV) Block totally blocks a number of paced atrial depolarizations (P waves) before conduction to the ventricles is successful. This produces 2:1 (two P waves to one QRS) or 3:1 (three P waves to one QRS) or even higher AV ratios. The series repeats. Mobitz is a serious problem; notice the extremely slow ventricular rates, which may produce loss of consciousness (syncope).

Mobitz (2º AV Block) may appear (at a normal Sinus rate) as two P waves to one _____ response, often referred to as “2:1 AV block” (or simply “2:1 block”).

Note: Mobitz sometimes blocks three atrial depolarizations (P waves) producing a single ventricular response (QRS); this is written “3:1 AV block,” or just “3:1 block”, which describes the mechanism of conduction. Poorer conduction ratios (e.g., 4:1, 5:1, etc.) relate to increased severity of the block and are sometimes called “advanced” Mobitz block.

Warning! With Mobitz, every cycle that is missing its QRS has a regular, punctual P wave — but never a premature P’ wave (see Note, page 128). This distinction is critical!

* Previously called “Type II” or “Mobitz II.”
2:1 AV Block
Wenckebach vs. Mobitz

Most likely Wenckebach...

if the PR interval is lengthened, but the QRS is normal.

Most likely Mobitz...

if the PR interval is normal, but the QRS is widened.

Both Wenckebach and Mobitz have missing ("dropped") QRS's, so how can we differentiate between 2:1 Wenckebach and 2:1 Mobitz? Wenckebach is considered innocuous and Mobitz is considered pathological, so we should differentiate.

Note: On EKG, a 2:1 AV block could be a short, two-cycle Wenckebach. For example, if the first cycle is fairly normal but in the second cycle the PR lengthens just enough to prevent conduction through the AV Node, this is 2:1 Wenckebach. But by its appearance alone, most of us would probably interpret (correctly?) a 2:1 block as Mobitz. Perhaps the following will help...

Because Wenckebach commonly originates in the AV _____, a 2:1 AV block of this origin often has an initial lengthened PR with no wide QRS pattern* (typical of Bundle Branch Block).

But since Mobitz originates below the AV Node, in the His Bundle or the Bundle Branches, we recognize that it often has a normal PR with a widened _____ (Bundle Branch Block) pattern.*

Note: Since differentiating between these two types of 2:1 AV Block is clinically very important, we may need to employ bedside diagnostic techniques to make the distinction (next page).

* The wide QRS pattern, typical of Bundle Branch Block, is soon explained (pages 191-202)
2:1 AV Block
Wenckebach vs. Mobitz

Differentiation of 2:1 Wenckebach (AV Node block) and Mobitz (AV conduction system block) is important clinically. In order to determine which type of 2:1 (2° AV) block a patient has, we can carefully employ vagal maneuvers (see page 61).

Occasionally an EKG of 2:1 block (like the one on the cover of this book) has criteria (i.e., for PR length and QRS width) that fit both Wenckebach and Mobitz. This may require the judicious use of a vagal maneuver to differentiate between the two.

The AV Node is richly supplied with parasympathetic innervation, so vagal __________ inhibits the AV Node, making it more refractory.

Therefore, vagal maneuvers increase parasympathetic inhibition of the AV __________, increasing the number of cycles/series to produce 3:2 or 4:3 Wenckebach.

But if the 2:1 block is Mobitz (i.e., the block is in the ventricular conduction __________), vagal maneuvers either eliminate the block, producing 1:1 AV conduction, or they have no effect.
On every EKG check:

1. **PR Interval**
   - increased consistently in 1° AV block
   - progressively increases in each series of cycles with Wenckebach
   - totally variable in 3° AV block
   - decreased in WPW and LGL syndromes

2. **P without QRS response**
   - Wenckebach and Mobitz 2° AV blocks
   - 3° AV block - independent atrial and ventricular rates

Let’s take a moment to see why routine EKG examination requires both checking the **PR interval** and looking for **P waves missing their QRS response**. Routinely checking these two parameters can reveal the entire spectrum AV conduction problems.

A prolonged **PR interval** can alert you to the existence of 1° AV block, 2° AV block, and 3° AV ______.

An EKG with **P waves lacking a QRS response** can expose 2° AV blocks and 3° ____ block.

**Note:** Let’s pause to contemplate how these two diagnostic parameters relate to each type of AV block. Really, take a moment and try this. It’s not just a meaningless exercise. You have consumed a great deal of practical knowledge about AV blocks. Now you can easily detect these blocks by checking both parameters on every EKG you see. In each instance, you should consider not only the anatomical origin of the problem, but its prognostic significance to the patient as a person. Congratulations on your progress; you should take pride in your knowledge.
An examining physician noticed that a patient had an irregular pulse. The doctor was surprised to feel a group of three pulse beats followed by a pause, and this group of beats seemed to repeat over and over. Let's share the patient's EKG.

First we scan the PR intervals and discover that the third cycle has a PR interval that exceeds .2 sec., so we suspect some kind of ___ Block.

While looking for P waves that lack a QRS response, we notice a lone P wave with no ____ response following the last complete cycle.

Upon close examination, we see that the PR interval normal at first, but becomes progressively longer with each successive cycle. We now recognize _________ block.
Complete (3°) AV Block

When the conduction of supraventricular depolarizations to the ventricles is totally blocked...

focus escapes to pace the ventricles

an automaticity focus escapes to pace the ventricles at its inherent rate.

Complete (3°) AV block is a total block of conduction to the ventricles, so atrial depolarizations are not conducted to the ventricles. Therefore, an automaticity focus below the complete block escapes to pace the ventricles at its inherent rate.

3° block is a complete block that prevents sinus-paced atrial depolarizations from reaching the ventricles.

A single block of the AV Node or the His Bundle can be “complete,” but more distally in the ventricular conduction system, there must be complete blocks of all subdivisions (branches) to eliminate conduction to the ventricles.

Absent paced depolarizations from above, an automaticity focus below the complete block escapes to pace the ventricles at its inherent rate.

Note: The location of the escaping focus depends on the location of the complete (3°) block. Next, let’s look at the possibilities.
Forms of Complete (3°) AV Block

Complete block in the upper AV Node leaves Junctional foci to escape and pace the ventricles.

Complete block of the entire AV Node or in the His Bundle leaves only a ventricular focus to pace.

Below the His Bundle, all paths are completely blocked, so a ventricular focus escapes.

Complete (3°) AV block occurs either in the upper AV Node, allowing a Junctional focus (below the block in the AV Node) to escape and pace the ventricles, or... the complete block may be below the AV Node, leaving only a ventricular focus to escape and pace the ventricles. To be a "complete block," all avenues of AV conduction must be blocked.

If a complete block is high in the AV Node, a Junctional focus (the fastest-pacing focus below the block), escapes to pace the ___________ at its inherent rate.

ventricles

If a complete block destroys the entire AV Node or is below the AV Node (for instance in the His Bundle), that leaves only ventricular foci to assume pacing responsibility...

...so a ventricular focus escapes to pace at its (slow) ________ rate. inherent

Note: Regardless of the location of the focus that escapes to pace the ventricles, the atria remain independently paced by the SA Node. From EKG, we see a Sinus-paced atrial (P wave) rate and a totally independent, focus-paced, slow ventricular (QRS) rate. Complete AV block produces this "AV dissociation" that records on EKG as an (usually normal) P wave rate superimposed over an independent, slower QRS rate. The AV dissociation (on EKG or cardiac monitor) acts as off that there is probably a complete AV block.
If a complete AV block occurs above the AV Junction (i.e., in the upper AV Node), then a Junctional focus, no longer overdrive-suppressed, escapes to pace the ventricles. On EKG, we see Sinus-paced P waves and a slower, independent QRS rate, usually with normal QRS complexes.

**Note:** If the complete AV block is in the AV Node, above the AV Junction, then a Junctional focus, no longer overdrive-suppressed, escapes to pace the ventricles. This is an “idiojunctional rhythm.”*

With a complete AV block, if the QRS’s appear normal (because each pacing stimulus passes down the ventricular conduction system), we know that a Junctional focus must be pacing the ventricles.

**Note:** Sometimes paced depolarizations from a Junctional focus may have to pass through diseased regions in the ventricular conduction system, delaying depolarization in some areas of the ventricles, producing wide QRS complexes.

If the ventricular rate ranges between 40 and 60, then a focus in the _____ is probably pacing the ventricles.

* Pacing by a Junctional focus may accelerate to become an accelerated idiojunctional rhythm.
When a complete AV block occurs below the AV Junction, a ventricular focus escapes overdrive suppression to pace the ventricles at its slow inherent rate of only 20 to 40 per min.; so slow, in fact, that cerebral blood flow is compromised and syncope may ensue.

**Noticing AV Dissociation** (separate atrial P wave) and ventricular (QRS) rates, you should check the morphology of the QRS's. **When we see large, wide, PVC-like complexes, we know that the ventricles are probably being paced by a ________ focus.** ventricular

We also see that the ventricular rate is within the inherent rate range (20 to 40 / min.) of a ventricular ______. focus

**Note:** We understand that a ventricular focus could only escape to pace if there were no Junctional focus available above it. So the complete AV block either obliterated the entire AV Node or it occurred below the AV Junction (i.e., below the AV Node).

**Note:** In 3rd (complete) AV block the ventricular rate may be so slow that blood flow to the brain is inadequate, and the patient may lose consciousness (syncope). This is *Stokes-Adams Syndrome.* Patients with complete AV block need continuous surveillance and maintenance of airway... many die needlessly without. Respond! Patients with 3rd AV block eventually need an artificial pacemaker.
Downward displacement of the pacemaker

No visible supraventricular activity

prognosis worse for:
  • wider complexes  • diminished amplitude  • slower ventricular rate

The above tracing is not caused by a 3º AV block. Don’t be trapped by assuming that wide complex bradycardia is always due to a 3º block. Can you see signs of independent atrial activity? In practice, you should check all leads.

Bradycardia with wide ventricular complexes is not always pathognomonic for complete AV block, so identify AV dissociation (independent atrial and ventricular activity) before calling any wide complex bradycardia a 3º AV _______

Note: The absence of atrial activity with wide complex bradycardia indicates that neither the SA Node nor supraventricular foci are viable enough to pace the atria. This failure of all automaticity centers above the ventricles, called “downward displacement of the pacemaker” usually carries an unfavorable prognosis. Before pronouncing this “downward displacement,” make certain that the flat baseline is not due to atrial fibrillation.

Note: Extremely high serum K⁺ concentrations “hyperkalemia” can severely depress the SA Node and supraventricular foci, producing the same EKG findings. Hyperkalemia can cause cardiac asystole, a form of cardiac arrest.

Why don’t we all take a break. Then, a little refreshed, we can look at Bundle Branch Block... next page.
Bundle Branch Block (BBB) is caused by a block (of conduction) in the Right or in the Left Bundle Branch. The blocked Bundle Branch delays depolarization to the ventricle that it supplies.

Normally, the Right Bundle Branch quickly conducts the stimulus of depolarization to the right ventricle, and the Left Bundle Branch does the same to the left ventricle. The depolarization stimulus is conducted to both ventricles at the same time (i.e., simultaneously).

A block of one of the Bundle Branches produces a delay of depolarization of the ventricle that it supplies.

Note: Ordinarily both ventricles are depolarized simultaneously. But with Bundle Branch Block, the unblocked Bundle Branch conducts normally, while depolarization in the blocked Bundle Branch has to creep slowly through the surrounding muscle (which conducts more slowly than the specialized Bundle Branch) to stimulate the Bundle Branch below the block. After the delay, depolarization proceeds rapidly again below the block. However, the delay in the blocked Bundle Branch allows the unblocked ventricle to begin depolarizing before the blocked ventricle (see next page).
Bundle Branch Block

Therefore in Bundle Branch Block, one ventricle depolarizes slightly later than the other, causing two “joined QRS’s” to appear on EKG.

When a Bundle Branch Block is present, either the left or the right ventricle may depolarize late, depending on which Bundle ________ is blocked.

**Note:** Individual depolarization of the right ventricle and depolarization of the left ventricle are still of normal duration. Because the ventricles do not depolarize simultaneously, this produces the “widened QRS” appearance that we see on the EKG. The two “out-of-sync” QRS’s are superimposed on one-another, and the machine records this combined electrical activity as a widened QRS with two peaks.

**Note:** Because the “widened QRS” represents the non-simultaneous depolarization of both ventricles (one punctually depolarized, the other slightly delayed), we usually see two R waves named in sequential order: R and R’. The R’ (pronounced “R-prime”) represents delayed depolarization of the blocked ventricle.
In Bundle Branch Block the “widened QRS” increases in duration to three small squares (.12 sec.) or greater, and two R waves (R and R') appear. The R' designates the delayed depolarization of the blocked ventricle.

**Note:** Simultaneous depolarization of the ventricles normally occurs in less than twelve hundredths second, producing a QRS that is less than three small squares in duration.

The diagnosis of Bundle Branch Block is mainly based on the widened ______ (.12 sec. or more duration).

In order to make the diagnosis of Bundle Branch Block, the QRS complex should be at least ______ small squares wide (.12 second). Check the QRS width of every EKG that you read!

**Note:** The needle that records the EKG tracing moves rapidly enough to record most of the heart’s electrical activity accurately. However, with great deflections the needle lags a bit mechanically, sometimes giving us an exaggerated duration on the tracing. Therefore, it is best to check the limb leads for QRS duration (where QRS amplitude is minimal), rather than the chest leads where the QRS deflections are often great.

**Note:** If a patient with BBB develops a supraventricular tachycardia, the rapid succession of widened QRS’s may imitate Ventricular Tachycardia. Careful!
In *Left Bundle Branch Block* (LBBB), left ventricular depolarization is delayed. In *Right Bundle Branch Block* (RBBB), right ventricular depolarization is delayed.

In Bundle Branch Block, you first notice the widened _____ complex. Then you should be able to find the R,R’ configuration in the chest leads.

In Right Bundle Branch Block, the left ventricle depolarizes punctually, so the R represents left ventricular depolarization, and the R' represents delayed _____ ventricular depolarization.

In Left Bundle Branch Block, left ventricular depolarization is delayed, so the right ventricle depolarizes punctually (R), and the R' represents delayed _____ ventricular depolarization.

Kind of easy to understand, isn’t it?
If there is a Bundle Branch Block, look at leads $V_1$ and $V_2$ (right chest leads) and leads $V_5$ and $V_6$ (left chest leads) for the R,R'.

When the QRS complex is wide enough (.12 sec. or more) to make the diagnosis of BBB, we immediately look at the right and left chest leads for the R,R'.

**Note:** During ventricular depolarization and just afterward (up to the peak of the T wave), any additional stimulus cannot depolarize the ventricles, that is, they are refractory to a premature stimulus. The Bundle Branches have a refractory period, but the Left and Right Bundle Branch refractory periods are not identical, so with a supraventricular tachycardia one Bundle Branch is receptive to stimulation before the other. At a certain critical rapid rate, one Bundle Branch conducts before the other, producing non-simultaneous depolarization of the ventricles. So this rate-dependent Bundle Branch Block produces a tachycardia with wide QRS's that imitates Ventricular Tachycardia.

The right chest leads are $V_1$ and ____.

$V_2$
Right Bundle Branch Block produces an R,R' in the right chest leads, V₁ or V₂.

With a wide ____ (and a diagnosis of BBB), check the right and left chest leads for R,R'.

Then, if there is an R,R' in the right chest leads V₁ or V₂ there is probably a ______ Bundle Branch Block.

In Right Bundle Branch Block, the right ventricle is depolarizing slightly later than the left ventricle, so the R' in the above illustration represents the delayed depolarization to the (blocked) ______ ventricle.
Left Bundle Branch Block

With a BBB, an R,R' in the left chest leads V₅ or V₆ means that Left Bundle Branch Block is present. The R' represents delayed depolarization of the left ventricle.

The chest electrode is located over the left ventricle in left chest leads ___ and V₆. 

Occasionally, the R,R' in V₅ or V₆ will appear only as a flattened peak with two tiny points in ____ Bundle Branch Block. (Examine the QRS in V₅ in the illustration).

In LBBB the right ventricle depolarizes before the left ventricle, so the first portion of the wide QRS represents ____ ventricular depolarization.

Notes: Make a mental note of the typical QRS pattern (i.e., shape) of Right and Left BBB. A diagnosis is often made by appearance alone. These patterns are important because sometimes a PVC or the ventricular complexes in VT are said to have a “RBBB” or “LBBB” pattern; you should understand what that means. The same is true of the ventricular complexes (on EKG) produced by artificial pacemaker electrodes.

Notes: The Left Bundle Branch has two subdivisions (“fascicles”); blocks of these fascicles are called Hemiblocks (pages 295 - 305).
Remember, a wide QRS (three small squares) indicates BBB, and you should identify which Bundle Branch is blocked by checking the left and right chest leads.

To diagnose BBB, the QRS complex must be at least 0.12 seconds in duration. Now, just for smiles, let’s identify the type of BBB in the illustration on page 193.

**Note:** In some individuals recovery from refactoriness (during the last stage of repolarization) differs slightly in duration between bundle branches. So only at a particular critical rate of tachycardia, one ventricle depolarizes after the other to produce a rate-dependent Bundle Branch Block (see Note, page 195).

The R,R' pattern may occur in only one chest lead. It is often difficult to see the R', but usually it can be found in the right chest leads V₁ or V₂ or in the left chest leads V₅ or V₆.

**Note:** Occasionally you will see an R,R' in a QRS of normal duration. This is called “incomplete” BBB.
Intermittent Mobitz (2° AV Block)

Intermittent dropped QRS

Occasional dropped QRS due to permanent BBB (one side) with intermittent BBB of the other side.

Simultaneous RBBB and LBBB prevents depolarization from reaching the ventricles; this is a complete (3°) AV block. So, block of one Bundle Branch with intermittent block of the other produces intermittent complete AV block, intermittent Mobitz.

Right BBB plus intermittent Left BBB will record on EKG as continuous Right BBB pattern QRS’s with intermittent episodes of complete AV block (P waves without _____ response).

Left BBB plus intermittent Right BBB will record on EKG as continuous Left BBB pattern QRS’s with intermittent episodes of complete AV block (P waves without _____ response).

**Note**: An EKG tracing or cardiac monitor display with a continuous BBB pattern QRS with an occasional missing QRS indicates intermittent complete AV block. The intermittent block may worsen, eventuating in a constant complete AV block. This intermittent Mobitz (exactly what it is) flashes an important warning sign. Intermittent Mobitz is the heart’s warning that eventually it will need an artificial pacemaker to drive the ventricles at a normal rate. Don’t let it slip by you unnoticed... for the patient’s sake!
Diagnosis: BBB

Check for

Right

\[ V_1 \rightarrow V_2 \]

Left

\[ V_5 \rightarrow V_6 \]

Remember, a wide QRS (three small squares) indicates BBB, and you should identify which Bundle Branch is blocked by checking the left and right chest leads.

To diagnose BBB, the QRS complex must be at least [ ] of a second in duration. Now, just for smiles, let's identify the type of BBB in the illustration on page 193.

**Note:** In some individuals recovery from refractoriness (during the last stage of repolarization) differs slightly in duration between bundle branches. So only at a particular critical rate of tachycardia, one ventricle depolarizes after the other to produce a *rate-dependent* Bundle Branch Block (see Note, page 195).

The R,R' pattern may occur in only one chest [ ]. It is often difficult to see the R', but usually it can be found in the right chest leads \( V_1 \) or \( V_2 \) or in the left chest leads \( V_5 \) or \( V_6 \).

**Note:** Occasionally you will see an R,R' in a QRS of normal duration. This is called “incomplete” BBB.
Rhythm: always observe

... for AV Block
(also P's missing a QRS response)

QRS... for Bundle Branch Block

Remember that you must always visually measure the duration of the PR intervals and the duration of the QRS complex when examining the rhythm on an EKG. Observation will suffice.

On all EKG's, you must always measure the PR intervals because if one is prolonged more than one large square, then there is some kind of ___ Block present (and, of course, look for missing QRS's, which indicate that a 2° or 3° AV Block is present).

On all EKG's, the QRS duration must be measured for if it is prolonged to .12 second or ____ there is a Bundle Branch Block.

Note: Always check the PR intervals and the QRS duration when scrutinizing the rhythm on any EKG. This must be part of any EKG interpretation. The spontaneous appearance of Mobitz AV block or Bundle Branch Block may be an early warning of impending infarction.

Note: Hemiblocks commonly occur with infarction, so they are described in the Infarction chapter. A hemiblock is a block of one of the two subdivisions ("fascicles") of the Left Bundle Branch.

*Once you visually check these criteria on EKG, you should record the precise PR and QRS duration.
Bundle Branch Block

Vector = ?

Ventricular Hypertrophy?

The Mean QRS Vector “Axis” will get to this in the next chapter and ventricular hypertrophy cannot be determined accurately in the presence of Bundle Branch Block.

Note: Because the Mean QRS Vector represents the general direction of the simultaneous depolarization of the ventricles, with BBB it is very difficult to represent such a vector. This is because with BBB the ventricles do not depolarize simultaneously, so there are really two separate (right and left) ventricular vectors.

Note: The criteria for ventricular hypertrophy (enlargement) are based on a normal QRS. Bundle Branch Block produces large QRS deflections because each ventricle lacks the (usual) simultaneous electrical opposition from depolarization of the other ventricle. Therefore the EKG diagnosis of ventricular hypertrophy should be very guarded with BBB. However, atrial hypertrophy can be diagnosed in the presence of BBB.

Note: Let’s review all the illustrations in this chapter. Then see “Blocks” in the Personal Quick Reference Sheets on page 339, and relate this to the simplified methodology that is summarized on page 334.
Axis refers to the direction of the movement of depolarization, which spreads throughout the heart to stimulate the myocardium to contract.

Note: The axis around which the earth rotates has nothing to do with electrocardiography, but we can borrow the large arrow ("Axis") in the illustration.

The progressive depolarization of the myocardium moves in a certain direction.

Axis refers to the direction of depolarization as it passes through the heart.

* Sometimes called "electrical axis."
Direction Of

Vector

Electrical Stimulus

To demonstrate the direction in which depolarization is moving, we use an arrow that is called a "vector."

We can demonstrate the general direction of the movement of depolarization by using a _________.

The vector shows the direction in which ________ is moving.

When interpreting EKG's, a vector shows the general ________ of depolarization in the heart.
The QRS complex represents the depolarization of the ventricular myocardium.

The QRS complex represents the simultaneous depolarization of both ____________ ventricles.

Ventricular depolarization and contraction can be said to occur at the same time, (but we know that ____________ contraction lasts a little longer).

Depolarization of the ventricles and their contraction is represented by the ____ complex. QRS
We can use small vectors to demonstrate ventricular depolarization, which begins at the endocardium that lines both ventricles and proceeds toward the outside surface (epicardium) in all areas at once.

**Note:** Once depolarization is beyond the AV Node, the ventricular conduction system conducts this stimulus to the ventricles with great speed. In this way, ventricular depolarization begins at the endocardial lining of the ventricles and proceeds through the thickness of the ventricular wall in all areas at about the same time. (We will not yet address depolarization of the ventricular septum).

The Purkinje fibers transmit depolarization to the myocardial cells just beneath the endocardium that lines both ventricles; this occurs so fast, that depolarization begins at the general level of the ____________ in all areas at about the same time. ____________

Depolarization of the ventricles generally proceeds from the endocardial lining to the outside (epicardial) surface through the full thickness of the ____________ wall in all areas at once. ____________

(See small vectors in the illustration).

**Note:** Notice that the thicker left ventricular wall has larger vectors.
If we add up all the small vectors of ventricular depolarization (considering both direction and magnitude), we have one large "Mean QRS Vector" that represents the general direction of ventricular depolarization.

The Mean QRS Vector is the sum of all the smaller vectors of ventricular depolarization.

By convention we consider the origin of the Mean QRS Vector to be the AV Node, so the "tail" of the Vector is always the AV Node.

Because the small depolarization vectors of the thicker left ventricle are larger (previous page), the Mean QRS Vector points more toward the left.

Note: Remember that a vector represents both direction and magnitude of depolarization... bigger vectors represent greater magnitude.
The Mean QRS Vector normally points downward and to the patient’s left, because this is the general direction of ventricular depolarization.

The ventricles are in the left side of the chest and angle downward and to the ______.

The ______ Vector points downward and toward the patient’s left side.

**Note:** From now on, we occasionally will use the word “Vector” (with a capital “V”) to represent the Mean QRS Vector, which depicts the general direction and magnitude of ventricular depolarization. Visualize the Vector over the patient’s chest, and remember that it begins in the AV Node.

**Note:** Depolarization is an advancing wave of Na⁺ ions.
The position of the Mean QRS Vector is described in degrees within a circle drawn over the patient’s chest. This circle is in the frontal plane. The limb leads are used to determine the position (“Axis”) of the Mean QRS Vector in the frontal plane.

We can locate the position of the Mean QRS Vector within a large _______ around the heart.  

The center of the circle is the _______.  

The Vector normally points downward and to the patient’s left, that is, between 0 and _____ degrees.  

(Note: The “axis” of the heart is simply the Mean QRS Vector when located by degrees in the frontal plane. For example, the axis of the heart in the above illustration is about +40 degrees. Review the illustration and note that 0° is on the patient’s left, and that the lower half of the circle is “positive” degrees. The top half of the circle is “negative” degrees. Axis is often denoted in medical literature by an “A” as in “A +30°” or “A = +30°”, and it may be called “electrical axis.”)
If the heart is displaced, the Mean QRS Vector is also displaced in the same direction. The AV Node is always the tail of the Vector.

If the heart is rotated toward the patient’s right side, then the Mean QRS Vector moves toward the _____ as well. This is common in tall, slender individuals (see illustration).

In very obese people the diaphragm is pushed up (and also the heart), so the Mean QRS Vector may point directly to the patient’s ______. (See illustration).

The tail of the Vector is the _______.

Note: In obese individuals the increased abdominal pressure often pushes the diaphragm upward so the position of the displaced heart may be called a “horizontal heart”. By the same token, a tall, slender individual may have a so called “vertical heart”.
With hypertrophy (enlargement) of one ventricle, the greater depolarization activity of the hypertrophied side displaces the Mean QRS Vector toward the hypertrophied side.

There is increased depolarization in a _______________ hypertrophied ventricle.

So, the Mean QRS Vector deviates toward the ___________ ventricle that is hypertrophied.

**Note:** A hypertrophied ventricle has more (and larger) vectors, which draw the Mean QRS Vector in that direction.
In myocardial infarction there is a necrotic (dead) area of the heart that has lost its blood supply and does not depolarize. The unopposed vectors from the other side draw the Mean QRS Vector away from the infarct.

**Note:** Myocardial infarction occurs when a branch of one of the coronary arteries (the heart's own source of blood supply) becomes occluded. The area of myocardium supplied by this blocked coronary artery has no blood supply and becomes electrically dead (can't depolarize).

In myocardial infarction (coronary occlusion) there is an area in the ventricles that has no ______ supply. This infarcted area cannot depolarize, and therefore it has no vectors.

Since there is no depolarization (and no vectors) in the infarcted area, the vectors from the opposite side are unopposed, so the Mean QRS Vector tends to point away from the ______.
Now you understand why the Mean QRS Vector is diagnostically so valuable. "Axis" is the Mean QRS Vector when given in degrees, and the normal axis range is 0° to +90° in the frontal plane.

The Mean QRS Vector should point downward and to the patient's ______, that is, in the 0° to +90° range. This is the range of normal axis.

The Mean QRS Vector gives us valuable information about the position of the ______, and...

... insight into ventricular ____________, and it also provides us with valuable information concerning myocardial ____________.

**Note:** The Mean QRS Vector tends to point toward ventricular hypertrophy, and away from myocardial infarction. These basic principles of axis are so logical and easy to understand that you should employ this diagnostic® tool whenever a twelve lead EKG is available.

* The diagnosis of Hemiblocks (pages 295-305) is based on changes in QRS Axis.
To determine the direction of the Vector, visualize a sphere surrounding the heart, with the AV Node at the center of the sphere.

Visualize a large _________ surrounding the heart.

The AV Node is the _________ of the sphere.

**Note:** The Mean QRS Vector has the AV Node as its tail, and the tip of the arrow touches somewhere on the surface of this hypothetical sphere.
With the sphere in mind, consider lead I (left arm with the positive electrode, right arm with the negative electrode).

**Lead I** uses the right and left _____ for recording.

If lead I is introduced into the sphere, the patient’s left side (left arm) is ________.

In lead I the right arm is ________.

**Note:** Lead I passes through the center of the sphere, which is the AV Node.
Still considering lead I, the patient’s left-hand side of the sphere is positive, and the right half is negative. The center of the sphere is the AV Node.

We are considering only lead ___ at this time.

We will now consider the lead I sphere in two ________.

The patient’s right half of the sphere is __________.
As the positive wave of depolarization within the myocardial cells moves toward a positive (skin) electrode, there is a simultaneous upward (positive) deflection recorded on EKG.

An advancing wave of depolarization may be considered a moving wave of __________ charges.

When this wave of positive charges is moving toward a __________ skin electrode, there is a simultaneous upward (positive) deflection recorded on the EKG.

If you see a _________ (upward) wave on EKG, it means at that instant a depolarization stimulus is moving toward a positive skin electrode that is being used to record the EKG.
The positive electrode used to record the \textit{inferior} limb leads, II, III, and AVF, is on the left “foot.” The positive electrode that is used to record the \textit{lateral} limb leads, I and AVL, is on the left arm.

Let’s focus our attention on the only horizontal lead, that is, lead I, which uses a positive electrode on the _____ arm.

Next, we will look at the only vertical lead, AVF, which uses a \underline{__________} electrode on the left leg (“Foot”).

That was fast... let’s move on.
If the QRS complex is positive (mainly upright) in lead I, the Mean QRS Vector is pointing somewhere into the patient’s left half (i.e., the positive half) of the sphere.

Obtain an EKG tracing and check the ______ complex in lead I.

**Note:** We check the QRS complex because it represents ventricular depolarization on the EKG tracing.

If the QRS in lead I is mainly upright, it is ________ (positive or negative)...

... and if the QRS is positive in lead I, then the Mean QRS Vector points positively, that is, into the ______ half of the sphere (toward the positive skin electrode on the patient’s left arm).

**Note:** This point becomes clearer if you go back and reread the previous page and continue directly with this page. It comes into focus better on the second go ‘round.
Still considering lead I on the tracing, if the QRS complex is mainly negative (downward), the Vector points to the patient’s right side.

In lead I, if the QRS complex is mainly below the baseline, it is _________ (positive or negative).

Now checking the lead I sphere surrounding the patient, a Vector pointing into the negative half of the sphere points to the patient’s _____ side.

So if the QRS in lead I is mainly negative, then the Mean _____ Vector points to the patient’s right side (away from the positive electrode on the patient’s left arm).
If the QRS is negative in lead I (Vector toward the right), this is Right Axis Deviation.

If the Mean QRS Vector points toward the right, we expect the QRS complex in lead I to be negative.

If the Mean QRS Vector points to the patient's right side (to the right of a vertical line drawn through the AV Node), this indicates Right ______ Deviation.

So if the QRS complex is negative in lead ___, this indicates that there is Right Axis Deviation (R.A.D.).
By simple observation, we can tell whether the Mean QRS Vector points to the patient's left or right side.

In Lead I

QRS Negative
- \[ - \]

Vector to patient's Right

R
- [+]

Right Axis Deviation

QRS Positive
+ [+]

Vector to patient's Left

L
- [+]

Lead I is the best lead for detecting Right ______ Deviation.

If the QRS complex is positive in lead I (which it usually is), this indicates that there is no R.A.D., because the Vector is pointing to the _____ side of the patient.

When we record lead I on an EKG, the patient's left arm has the _________ electrode.
The left foot has a positive electrode in lead AVF. Imagine a sphere around the patient for lead AVF.

Ignore the lead discussed on the previous page. We are considering only lead _____ at this time.

**Note:** We are now considering a completely different sphere – the one that surrounds the body when we record lead AVF on the EKG machine. We need to re-orient ourselves as to the positive and negative halves of the sphere in AVF.

When we switch the EKG machine to monitor lead AVF, the machine makes the electrode on the left _____ positive.

The lower half of this sphere is ________ positive.

The center of this sphere is the _____ AV Node.
For AVF the lower half of the sphere is positive, and the upper half is negative.

The lower half of the AVF sphere is the location of the positive left foot electrode, so we know that the lower half of this sphere is _________.

The upper portion of the sphere (above the AV Node) is _________ (positive or negative).

The sphere in AVF has two halves, the upper half is _________.

... and the lower half of the AVF sphere is _________.

positive

negative
Considering lead AVF of the EKG, if the QRS is mainly positive on the tracing, then the Mean QRS Vector points downward into the positive half of the sphere, toward the positive (lead AVF) electrode.

If the Mean QRS Vector points downward, then the QRS complex in lead AVF is ________ upright (or positive).

**Note:** Don’t get confused just because the positive QRS is upright, yet the Vector points downward. You must remember that the Vector points into the positive half of the sphere (toward the positive left foot electrode) when the QRS is positive. The lower half of the sphere just happens to be the positive half in lead AVF.
In AVF, if the QRS is negative, the Vector points into the negative half of the sphere.

The center of the sphere is the ___ _______.

The upper half of the (lead AVF) sphere is _____________.

A negative QRS complex in lead AVF tells us that the Mean QRS Vector points ___________ into the negative half of the sphere (i.e., it is pointing away from the positive electrode on the left foot).
Chapter 7: Axis

Follow the illustration closely. If the QRS is positive in lead I and also positive in AVF, the Vector points downward and to the patient’s left. This is the normal axis range; the area that is both yellow and blue (yellow plus blue equals green).

A mainly positive QRS in lead I indicates that the Mean QRS Vector points to the _______ side of the patient, and...

... a mainly positive QRS complex in lead AVF means that the Vector points ____________.

In the same patient, if the Vector points leftward and also points downward, the Vector must be in the only quadrant of the __________ that satisfies both criteria (and it happens to be the normal range).

**Note:** Since the ventricles angle downward to the left, and ventricular depolarization moves downward and leftward, it should not surprise you that this is the normal range of the Vector. Remember, Vector position is stated in terms of the patient’s left or right. If the QRS is upright in both lead I and AVF (the “two thumbs up” sign), the Vector (“Axis”) is within the normal range.
In the frontal plane, there are four possible axis quadrants where the Mean QRS Vector may point. Visualize this large circle on the patient’s chest in the frontal plane.

**Note:** In the *frontal* plane, we determine if there is any *Deviation* of Axis out of the normal range.

If the Vector points upward (from the AV Node) and to the patient’s left, this is Left ______ Deviation (L.A.D.).

If the Vector points to the patient’s right side, this is ______ Axis Deviation (R.A.D.).

If the Vector points downward to the patient’s left, it is in the _______ range (i.e., Normal Axis).

**Note:** Remember, Axis is merely the position (that is, the direction) of the Mean QRS Vector, which indicates the general direction of ventricular depolarization.
For any patient, when we find into which quadrant (of the frontal plane) the Vector points, we know in which direction the ventricular depolarization is going. The small type in the illustration relates to the patient’s right or left.

**Note:** This is how you should visualize the four axis quadrants in a large circle (AV Node is the center) drawn on the patient’s chest in the frontal plane. On some EKG charts the Mean QRS Vector is depicted in a similar circle (which represents the frontal plane).

The upper left quadrant represents _____ Axis Deviation (L.A.D.).
If the QRS is *positive* in lead I, and *negative* in AVF, that places the Vector in the upper left quadrant. This is *Left Axis Deviation*.

If the QRS in lead I is upright, the Vector points to the patient’s _____.

If the Vector is pointing upward, then the QRS in lead AVF is mainly ______ the baseline.

And when the Vector points upward and to the patient’s left, this is Left _____ Deviation (L.A.D.).
Now, by looking at the QRS complex in I and AVF, you can locate the Mean QRS Vector in an Axis quadrant (in the frontal plane as it relates to the patient).

Any time the QRS complex is negative in lead I, there is ______ Axis Deviation (R.A.D.); and when the Vector also points upward (and to the patient’s right), this is commonly called “Extreme” R.A.D.

But if the QRS is positive in lead I and negative in lead AVF, there is Left Axis ________.

So if the Mean QRS Vector points downward and to the patient’s left, we expect the QRS complexes in leads I and AVF to be mainly ________ (upright). And of course, they usually are, positive since this is normal.

Note: You also can calculate the vector for a portion of a QRS complex (for instance, the initial or terminal .04 sec.) in exactly the same manner as for the Mean QRS Vector.
When depolarization moves in a direction perpendicular to the orientation of a lead, the deflection is minimal and/or “isoelectric.” An isoelectric QRS records equal magnitudes of upward (positive) and negative (downward) deflection.

Depolarization that moves perpendicular to the orientation of a lead, is directed negligibly toward either electrode, so the recorded deflection is as much negative as positive and is called ____________.

The word “isoelectric” literally means “same voltage,” so it is used when the positive and the negative portions of the QRS complex are about ______.

Although the positive and negative deflections of an isoelectric QRS are equal in amplitude, they are generally small in the limb ______.

**Note:** First, locate the Mean QRS Vector in an axis quadrant (i.e., Normal, L.A.D., R.A.D., or Extreme R.A.D.). Then, find the limb lead in which the QRS is the most isoelectric, so you can more precisely locate the Vector in degrees (Axis). The Axis is about 90° from the orientation of the most isoelectric lead. It is really very easy... next page.
Left Axis Deviation

<table>
<thead>
<tr>
<th>Most Isoelectric</th>
<th>Axis</th>
</tr>
</thead>
<tbody>
<tr>
<td>I ...............</td>
<td>-90°</td>
</tr>
<tr>
<td>AVR .............</td>
<td>-60°</td>
</tr>
<tr>
<td>II ..............</td>
<td>-30°</td>
</tr>
<tr>
<td>AVF .............</td>
<td>0°</td>
</tr>
</tbody>
</table>

Normal Range

<table>
<thead>
<tr>
<th>Most Isoelectric</th>
<th>Axis</th>
</tr>
</thead>
<tbody>
<tr>
<td>AVF .............</td>
<td>0°</td>
</tr>
<tr>
<td>III .............</td>
<td>+30°</td>
</tr>
<tr>
<td>AVL .............</td>
<td>+60°</td>
</tr>
<tr>
<td>I ...............</td>
<td>+90°</td>
</tr>
</tbody>
</table>

To locate the position of the Vector (Axis) more precisely (i.e., in degrees) in the frontal plane: first locate the axis quadrant, and then note the limb lead in which the QRS is most isoelectric.

Note: Please refer to the illustration on this page (and the page that follows) to determine the exact position of the Mean QRS Vector (Axis) in degrees. For exams and in "real life" situations you need a reference. Accuracy is far more important than memory. You may copy page 340; it's yours for real life use.

Note: Let's review. First, locate the appropriate axis quadrant. Then, to determine the exact position of the Vector (Axis), find the lead where the QRS is most isoelectric. Refer to the illustration as you contemplate the hypothetical examples below.

A patient with Left Axis Deviation has a Mean QRS Vector of between 0 and ____ degrees (QRS positive in I and negative in AVF). Check the illustration. (don't forget the negative)

A young lady has a Mean QRS Vector in the normal range. If the QRS in lead III is isoelectric, then she has an electrical axis of ____. Please don't proceed to the next page until you feel comfortable with this exercise.
The exact position of the Vector (Axis) can be located in a similar way for Right Axis Deviation and Extreme Right Axis Deviation. Refer back to the illustration for each sentence below.

**Note:** After the axis quadrant is determined, the limb lead with the most isoelectric QRS is noted.*

Consider a patient with R.A.D. You find that the QRS is isoelectric in lead II, so the Axis is _____.

You have a patient with numerous widened, premature QRS’s, and you need to know whether it is a PVC or an aberrant Junctional beat. The wide QRS is negative in I and AVF, which places its Vector in the Extreme ____ quadrant (how could that be?)…

… the wide QRS is also isoelectric in AVL, so its Axis is ____. For ventricular depolarization to progress in that direction, it must have originated in a focus (or pacemaker electrode) at the apex of the left ventricle, rather than from a Junctional focus. Let’s think about that.

**Note:** An Axis of 180° is either + or - depending on whether the Vector is in the R.A.D. or Extreme R.A.D. quadrant respectively.

* This is summarized for you (page 340) of your Personal Quick Reference Sheet.
The Mean QRS Vector, which represents normal biventricular depolarization, points downward and to the patient’s left. The P wave vector, which represents normal atrial depolarization, points downward to the left side of the patient.

A vector is used to represent the general direction of depolarization. When depolarization (a wave of positive charges) moves toward a positive electrode, it records __________-ly (upward) on the EKG. positive

Note: The P wave vector points generally downward, toward the positive electrode on the patient’s left foot (for inferior leads II, III, and AVF), so the P waves are usually upright in those leads. The P wave vector also points leftward, toward the positive electrode on the patient’s left arm (for leads I, and AVL), producing generally upright P waves in those leads. So, if we see an inverted “P wave” in any of those leads, it is probably a P’ depolarizing upwards from a low atrial focus, or retrograde atrial depolarization moving upward from the AV Node.

Note: Most PVC’s emanate from a peripheral focus in a ventricular wall, depolarizing the ventricles in a general bottom upward direction, so they are usually mostly negative in the inferior and lateral limb leads where the QRS is usually upward. Exception: PVC’s that are mainly upward like the QRS’s in those leads, probably originate in a septal ventricular focus and follow a near-normal path.
The sphere has three dimensions, so it is important to note the general position of the Mean QRS Vector in the horizontal plane as well.

The horizontal _______ divides the body into top and bottom halves.

The chest leads form the ____________ plane.

**Note:** To determine changes ("rotation") of the Mean QRS Vector in the horizontal plane, we examine the chest leads.

**Note:** Although the Axis may "deviate" in the frontal plane, the Vector is said to "rotate" in the horizontal plane. This is conventional (universally accepted) terminology used in communication and in medical literature.
Chest lead $V_2$ is obtained by placing a positive electrode on the chest along the left side of the sternum (at the fourth interspace).

The chest electrode used for recording lead $V_2$ is always _________ (positive or negative).

**Note:** The electrode for the chest leads is a suction cup that is moved to a different position on the chest for each of the six chest leads (which form the horizontal plane). In each case, the suction cup electrode is positive.

The position of the (suction cup) electrode for recording lead $V_2$ places it in front of the heart, at the fourth interspace to the left of the sternum, so it is just _________ to the AV Node.

**Note:** We already know that metal electrodes affixed with conductive gel are often used for recording the chest leads, so let's refocus on the conceptual material.
Considering the sphere for lead V₂ we can see that the front half is positive and the back half is negative.

Considering a lead V₂ sphere, we view the patient from the side. The center of the ________ is still the AV Node.

The patient’s back is _________ (negative or positive), when considering lead V₂.

The front half of the sphere is _________ in lead V₂.
Normally the QRS in lead $V_2$ is negative. Therefore, the Mean QRS Vector points backward, because of the (generally) posterior position of the thick left ventricle.

Considering lead $V_2$ on the standard EKG, the QRS complex is usually \underline{negative} (below the baseline).

Therefore, the Mean QRS Vector usually points \underline{backward} (posteriorly) into the negative half of the sphere.

Normally, most of the ventricular depolarization moves away from the positive $V_2$ electrode, toward the thicker and more posteriorly positioned \underline{left} ventricle.
The orientation of chest lead $V_2$ makes it the most informative lead for the determination of both Anterior and Posterior Infarction.

The orientation of lead $V_2$ projects through the anterior wall and the posterior wall of the _____ ventricle.

So $V_2$ reflects the most reliable information concerning Anterior Infarction and __________ Infarction of the left ventricle.

**Note:** As you will soon see, both ventricular depolarization and repolarization should be scrutinized in the right chest leads, because they reveal subtle vector changes caused by both anterior and posterior infarctions (of the left ventricle).
In the chest leads, there is a gradual transition from the generally negative QRS in $V_1$ to the generally positive (upright) QRS in $V_6$.

The QRS is mainly negative in lead $V_1$ and mostly ________ in lead $V_6$. positive

Examining chest leads $V_1$ through $V_6$ to observe the gradual transition of QRS complexes, we notice that the QRS usually becomes as much positive as negative or “________” isoelectric in lead $V_3$ or $V_4$. This is the transitional zone.

Note: You will recall that an isoelectric QRS is $90^\circ$ away from the Mean QRS Vector. So a shift ("rotation") of the Vector in the horizontal plane is reflected as a similar change in position of the "transitional" (isoelectric) QRS in the chest leads. You will better understand and appreciate this, when you see the next page.

As the Vector changes its position (rotates) in the horizontal plane, the Vector’s tail remains anchored to the _______. AV Node
Rotation of the Vector in the horizontal plane is described from the patient’s point of view as “rightward” or “leftward.” Check the chest leads for the isoelectric QRS.

**Note:** The Vector can rotate in the horizontal plane with its tail anchored to the AV Node. When the isoelectric (“transitional”) QRS has rotated to the patient’s right (into leads V₁ or V₂) this is rightward rotation. But if the transitional QRS is found in the patient’s left chest leads, V₅ or V₆ this is leftward rotation. Anatomically speaking the heart is not capable of much rotation in the horizontal plane. But, we do know that the Vector shifts toward Ventricular Hypertrophy and away from Infarction.

**Note:** In older literature you may still see the terms “clockwise” (meaning leftward) rotation or “counterclockwise” (meaning rightward) rotation of the Vector in the horizontal plane. These terms have become obsolete since they do not relate well to clocks, and much confusion resulted.

Reminder: Axis deviation is in the frontal plane.
Axis rotation is in the horizontal plane.

**Note:** Please observe the simplified technique for determining Axis by turning to page 340. A quick review of the methodology may be found in the Personal Quick Reference Sheets on page 334.
Hypertrophy usually pertains to an increase in size, but when relating to muscle as in myocardium, this term refers to increase in muscle mass.

Note: The photo above is the arm of a weight-lifter. I had contemplated using a photo of my own arm, but I soon abandoned the idea because then I would have to title this section “hypotrophy” (if there is such a word).
Hypertrophy of a chamber of the heart implies an increase in the thickness of the chamber wall, but some dilation is always present also.

Hypertrophy of a chamber of the heart means that the muscular wall of that chamber has dilated and thickened beyond ______ thickness.

Hypertrophy may increase the volume that the ________ contains, and the wall of that chamber is thicker than normal.

The increase in the muscular thickness of the wall of a hypertrophic chamber, as well as dilation of a chamber of the heart may be diagnosed on ____.
Since the P wave represents the depolarization and contraction of both atria, we examine the P wave for evidence of atrial enlargement. (See Note.)

The depolarization of both atria causes their simultaneous contraction.

The depolarization of both atria is recorded on EKG as a ___ wave. P

Signs of atrial enlargement can be detected by examining the P wave on the twelve lead EKG.

Note: Although the designation “atrial hypertrophy” is commonly used, enlargement of an atrium is usually due to dilation of the atrium. Therefore, the general term atrial enlargement is preferred, since it includes both dilation and hypertrophy. Whereas when referring to the ventricles, “ventricular hypertrophy” predominates.
Lead $V_1$ is directly over the atria, so the P wave in $V_1$ is our best source of information about atrial enlargement.

The chest electrode that records lead $V_1$ is considered _________ (positive or negative).

When lead $V_1$ is recording, the electrode is positioned just to the right of the sternum in the 4th interspace; this places the electrode directly over the ______.

Because the $V_1$ electrode is close to the atria, the P wave in lead $V_1$ gives us the most accurate information about atrial _________.

enlargement
Diphasic P Wave

With atrial enlargement, the P wave is usually *diphasic* (both positive and negative).

A wave that has both positive and negative portions is called a _________ wave (two phased wave).

A diphasic P wave has deflections above and below the _________.

The diphasic P wave is characteristic of atrial enlargement, but we want to know which of the two ________ is enlarged.

---

diphasic
baseline
atria
Right Atrial Hypertrophy

If the initial component of a diphasic P wave (in lead V₁) is the larger, then this is Right Atrial Enlargement.

If the P wave in lead V₁ is __________, then we know that one of the atria is enlarged.

If the initial portion of the diphasic P wave is the __________ of the two phases, then there is Right Atrial Enlargement.

A diphasic P wave in V₁ with a large, often peaked, initial component tells us that this patient’s ______ atrium is probably thicker and more dilated than the left.

**Note:** If the height of the P wave in any of the limb leads exceeds 2.5 mm (even if it’s not diphasic), suspect Right Atrial Enlargement.
Chapter 8: Hypertrophy

Left Atrial Hypertrophy

If the terminal portion of a diphasic P wave in $V_1$ is large and wide, there is *Left Atrial Enlargement*.

A patient who has enlargement of the left atrium, because the mitral valve is stenosed*, will have a diphasic P wave in lead $V_1$.

This patient’s diphasic P wave in lead $V_1$ has a small initial component and a larger ______ component.

The terminal component of a diphasic P wave in lead $V_1$ is usually ______ (positive or negative).

* *Mitral stenosis* (narrowed mitral valve opening) can cause left atrial enlargement, but systemic hypertension is the most common cause.
Normal QRS in Lead V₁

Now let's consider the QRS complex in V₁. Normally the S wave is much larger than the R wave in this lead.

The QRS complex represents ventricular depolarization, so we would expect the QRS to reflect the presence of ventricular hypertrophy.

In lead V₁ the QRS complex is mainly negative, and therefore the R wave is usually very short.

**Note:** The V₁ electrode is positive. Ventricular depolarization moves downward to the patient’s left side and also posteriorly (the thicker left ventricle is more posteriorly located). Because ventricular depolarization is moving away from the (positive) V₁ electrode, the QRS in V₁ is usually mainly negative. Remember that the positive wave of depolarization moving toward a positive electrode records a positive deflection on EKG. By the same token, depolarization moving away from a positive electrode records negatively.
Right Ventricular Hypertrophy

However, with Right Ventricular Hypertrophy (RVH) there is a large R wave in $V_1$.

In Right Ventricular Hypertrophy there is a large ___ wave in lead $V_1$.

**Note:** With Right Ventricular Hypertrophy, the wall of the right ventricle is very thick, so there is much more (positive) depolarization (and more vectors) toward the positive $V_1$ electrode. We would therefore expect the QRS in lead $V_1$ to be more positive (taller) than usual.

The S wave in lead $V_1$ is smaller than the ___ wave in Right Ventricular Hypertrophy. (See illustration).
With Right Ventricular Hypertrophy, the large R wave of $V_1$ gets progressively smaller from $V_2$ to $V_3$ to $V_4$ etc.

When Right Ventricular Hypertrophy is present, there is a large R wave in lead ___ that becomes progressively smaller in chest leads $V_2$, $V_3$, and $V_4$.

The progressive decrease in the height of the ___ wave is gradual, proceeding from the right chest leads to the left chest leads.

**Note:** The enlarged right ventricle adds more vectors toward the right side, so there is Right Axis Deviation (in the frontal plane), and in the horizontal plane there is rightward rotation of the (Mean QRS) Vector. Visualize the reasons for these (Mean QRS) Vector shifts and the criteria will become very logical.
Left Ventricular Hypertrophy

With Left Ventricular Hypertrophy (LVH), the left ventricular wall is very thick, causing great QRS deflections in the chest leads.

The heart chamber with the thickest muscular walls is the ______ ventricle.

Hypertrophy of the left ventricle produces QRS complexes that are exaggerated amplitude, both in height and in depth, especially in the ______ leads.

**Note:** Normally the S wave in V₁ is deep. But with Left Ventricular Hypertrophy, even more depolarization is going downward to the patient's left - away from the positive V₁ electrode. Therefore, the S wave is even deeper in V₁. There is Left Axis Deviation, and often the Vector is displaced in a leftward direction in the horizontal plane. Visualize and understand the reason for these shifts of the Vector. Lasting knowledge results from understanding.
With Left Ventricular Hypertrophy there is a large S in V₁ and a large R in V₅.

With Left Ventricular Hypertrophy there is a very tall ___ wave in lead V₅.

**Note:** Lead V₅ is over the left ventricle, so the increased depolarization is going toward the electrode of V₅ when there is L.V.H. This results in more (positive) depolarization going toward the (positive) electrode of V₅ which produces a very tall R wave in that lead.

In Left Ventricular Hypertrophy, there is a very tall R wave in lead ___, and this excessive depolarization moving away from the V₁ electrode produces a deep S wave in lead V₁.
Left Ventricular Hypertrophy

\[ V_1 \quad \text{mm of S in } V_1 \]

+ 

\[ V_5 \quad \text{mm of R in } V_5 \]

**Total**

(if more than 35 mm there is L.V.H.)

Depth (in mm) of S in V₁ plus the height of R in V₅... if greater than 35 mm, there is Left Ventricular Hypertrophy.

To check an EKG for Left Ventricular Hypertrophy, just add the depth of the S wave in V₁ to the height of the ___ wave in V₅.

If the depth (in mm) of the S wave in V₁ added to the height (in mm) of the R wave in V₅ is greater than 35 (mm), then ____ Ventricular Hypertrophy is present.

**Note:** The sum of the S in V₁ plus the R in V₅ should be routinely checked (mere observation will usually suffice) with every twelve lead EKG. When providing a written EKG interpretation, however, one should measure and document the amplitude of these waves in millimeters.
The T wave may show “Left Ventricular Hypertrophy” characteristics. Often there is T wave inversion with T wave asymmetry.

There is a characteristic T wave that is commonly associated with _____ Ventricular Hypertrophy.

Since the left chest leads (V₅ or V₆) are over the left __________, these are ideal leads to check for this characteristic T wave that we find with LVH.

With LVH, the inverted T wave has a gradual downslope and a very steep return to the __________, making it asymmetrical.
Ventricular hypertrophy may be associated with a strain pattern. With ventricular strain, the ST segment becomes depressed and humped.

Ventricular strain is characterized by depression of the ST segment.

Note: Strain is usually associated with ventricular hypertrophy, which is logical, since a ventricle that is straining against some kind of resistance (e.g., increased resistance from a narrowed valve or from hypertension) will become hypertrophied in its attempt to compensate.

Ventricular strain depresses the ST segment, which generally humps upward in the middle of the segment.
Note that lead $V_1$ provides most of the information concerning hypertrophy of the heart's chambers.

When routinely reading a 12 lead EKG, you should check to see if there is _________ of any of the chambers.

First, check lead $V_1$ to see if the P waves are _________.

Second, check the R wave in $V_1$... and then check the S wave in $V_1$ and the ___ wave in $V_5$.

**Note:** You may now review *Hypertrophy* by turning to the Personal Quick Reference Sheets on page 341 and relate this to the simplified methodology that is summarized on page 334.
Myocardial Infarction (M.I.) results from the complete occlusion of a coronary artery. The infarcted area of myocardium becomes necrotic (dead), so it can’t depolarize or contract.

Note: Although the heart’s chambers are filled with blood, the myocardium’s own blood supply is provided exclusively by the coronary arteries. A coronary artery can be gradually narrowed by lipid deposits that become atheromatous plaque beneath the intima lining of the vessel. The intima may eventually rupture, exposing the plaque to the blood within the artery. This initiates the immediate formation of a clot (thrombus). The vessel, already narrowed by the plaque, becomes totally occluded by the thrombus. Instantly the infarcted area of the ventricle (without a blood supply) becomes necrotic. Ventricular foci in the hypoxic area around the infarct become very irritable; this can produce deadly ventricular arrhythmias.

Note: Myocardial Infarction implies the complete occlusion of a coronary artery, which we can diagnose with the EKG. The electrocardiogram will also tell us which coronary artery (or coronary branch) is occluded, and it can even reveal any blocks in the ventricular conduction caused by the infarction. By careful interpretation of the EKG, we can also determine if a coronary vessel is narrowed, rendering a decreased blood supply to the heart. Practical lifesaving knowledge. Let me show you...
Myocardial infarction is due to the occlusion of a coronary artery supplying the left ventricle, so an area of the heart* is without a blood supply and suffers necrosis.

The terms "myocardial __________," "coronary occlusion," and "heart attack" refer to the same serious problem.

The heart derives its own blood supply from the __________ arteries, so when a coronary artery or one of its major branches is occluded, an area of the myocardium is without blood supply.

The infarcted (necrotic) area is primarily in the ____ ventricle; deadly arrhythmias may result.

Note: We understand that the coronary arteries also supply the right ventricle, so there is often some involvement of the right ventricle. But since most of the critical problems originate in left ventricular infarcts, myocardial infarction is usually conceptualized in terms of the left ventricle.

* In this illustration, the pulmonary artery has been "surgically" removed to show the origin of the coronary arteries at the base of the aorta.
Commonly, it is the thick left ventricle that suffers myocardial infarction.

The left ventricle is the thickest chamber of the heart; so if the coronary arteries are narrowed, the left ventricle (which uses the greatest blood supply) is the first to suffer from an obstructed coronary artery.

Blood is pumped to all parts of the body by the powerful, thick, left ventricle.

Note: When we describe infarcts by location, we are speaking of an area within the left ventricle. Coronary arteries to the left ventricle usually send smaller branches to other regions of the heart, so an infarction of the left ventricle can include a small portion of another chamber.
The necrotic infarcted area of the left ventricle (that has no blood supply) is electrically dead and cannot depolarize.

Infarctions usually involve an area of the wall of the left _________.

An area of infarction cannot be depolarized because the cells there are without a ________ supply, so they are necrotic (functionally dead).

**Note:** This necrotic infarcted area produces an electrical void, while the rest of the heart (with an adequate blood supply) functions as usual. The infarcted region does not depolarize, so it does not contract, thereby impairing the muscular function of the left ventricle. Also, hypoxic ventricular foci nearby are often the source of serious ventricular arrhythmias.
Infarction

- ischemia
- injury
- necrosis

The myocardial infarction triad is "ischemia," "injury," and "necrosis," but any of the three may occur alone.

**Note:** Necrosis (death) of a ventricular region produces dead myocardial cells that cannot depolarize.

The myocardial infarction triad is the basis for recognizing and diagnosing a _________ infarction. myocardial

The word hypoxia means decreased oxygen; in the heart it is usually caused by ischemia, which literally means reduced _______ supply (diminished blood flow). blood

**Note:** Ischemia, injury, and necrosis need not all be present at once in order to establish the diagnosis of myocardial infarction. Routine EKG interpretation requires checking these infarction criteria.
Ischemia (decreased blood supply) is characterized by inverted T waves.

Ischemia means reduced _____ supply (from the coronary arteries); the ischemic area is at the periphery of the infarct.

The characteristic sign of ischemia is the ________ T wave. It may vary from a slightly inverted to a deeply inverted T wave.

Inverted ___ waves may indicate ischemia in the absence of myocardial infarction. Coronary blood flow can decrease without producing an infarction.

**Note:** Cardiac ischemia alone can cause chest pain known as angina, which is usually associated with transient T wave inversion.
Ischemia

The typical ischemia T wave is symmetrically inverted.

**Note:** You should check every EKG that you read for T wave inversion. Since the chest leads are nearest the ventricles, T wave changes are most pronounced in these leads. Always run down $V_1$ to $V_6$ (as well as the limb leads) and check for T wave inversion to see if there is diminished coronary flow.

The T wave of ischemia is both inverted and ________, that is, the right and left sides of the inverted T wave are mirror images.

**Note:** In adults flat (nonexistent) T waves or minimal T wave inversion may be a normal variant in any of the limb leads (frontal plane). However, any T wave inversion in leads $V_2$ through $V_6$ is considered pathological. Marked T wave inversion in leads $V_2$ and $V_3$ the hallmark of Wellens syndrome, alerts us to stenosis of the anterior descending coronary.
Injury: means acute or recent

Injury indicates the acuteness of an infarct. Elevation of the ST segment denotes "injury" sometimes called the "current of injury."

Note: "Acute" means recent or new.

The ST segment is that section of baseline between the QRS complex and the ___ wave. The ST segment contains no waves.

Elevation of the ST segment signifies "injury." The ST segment may be elevated only slightly, or as much as ten or more millimeters above the _________.

ST segment elevation tells us that a myocardial infarction is _____. It is the earliest consistent sign of infarction to record on EKG.

Note: Angina without exertion, "Prinzmetal's" angina, can cause transient ST elevation in the absence of an infarction.
If there is ST elevation, this indicates that the infarction is acute. ST elevation, alone, can indicate an infarction.

**Note:** Once you have made a diagnosis of infarction, it is important to know whether the infarction just occurred and needs immediate treatment, or if the infarction is old — maybe years old.

The ST segment rises above the baseline with an acute infarction, in fact it is usually the earliest EKG sign of an infarction. With time, the ST segment returns to the baseline.

**Note:** If the ST segment is elevated without associated Q waves, this may represent *non-Q wave infarction*, which is usually a small infarction that may herald an impending larger infarct. Significant ST changes require enzyme studies and close scrutiny.

**Note:** A *ventricular aneurysm* (the outward ballooning of the wall of a ventricle) can cause persistent ST elevation in most of the chest leads; but in this case, the ST segment does *not* return to the baseline with time. *Pericarditis* (next page) produces a unique type of ST segment elevation that may also elevate the T wave off the baseline.
Brugada Syndrome

- RBBB pattern QRS with ST elevation in V₁ - V₃
- sudden cardiac arrest (in absence of coronary obstruction)

Brugada syndrome is a hereditary condition that can cause sudden death in individuals without heart disease. It is characterized by Right Bundle Branch Block with ST elevation in leads V₁ to V₃. Look for it; this malady is not rare.

Sudden cardiac death (cardiac arrest) can occur spontaneously in patients with Brugada syndrome.

In Brugada syndrome there is RBBB and ST elevation in leads V₁ to V₃. The elevated ___ segments have a peculiar, peaked downsloping shape, particularly in V₁ and V₂.

Note: Brugada syndrome is a familial condition caused by dysfunctional cardiac Na⁺ (sodium) channels. Prophylaxis against the deadly arrhythmias requires ICD implantation in order to immediately treat cardiac arrest (usually ventricular fibrillation.)

Note: This syndrome is responsible for nearly one-half of the sudden deaths in healthy young individuals without structural heart disease.
With pericarditis, the ST segment is elevated and usually flat or concave. The entire T wave may be elevated off the baseline.

**Note:** Pericarditis is inflammation of the membrane (pericardium) that surrounds the heart. Pericarditis may be caused by a virus, bacteria, cancer, or other sources of inflammation, including myocardial infarction.

Pericarditis can elevate the ___ segment. It usually produces an elevated ST segment that is flat or slightly concave (middle sags downward). This resolves with time.

Pericarditis seems to elevate the entire ___ wave off the baseline; that is, the baseline gradually angles back down (often including the P wave) all the way to the next QRS (illustration on right).

**Note:** The characteristics shown in the left illustration are found in leads in which the QRS is usually mainly negative (like the right chest leads). The pattern shown in the right illustration is seen in leads where the QRS is mainly positive (such as the lateral and inferior limb leads). Sometimes PVC’s are produced.
The ST segment may become depressed under certain circumstances or conditions.

**Note:** During an angina* attack, the ST segment may be temporarily depressed.

A subendocardial infarction, an infarct that does not extend through the full thickness of the _____ ventricular wall, will depress the ST segment.

When a patient with narrowed coronaries exercises, the myocardium demands more blood flow than its arteries can deliver. A stress (or “exercise”) test will record depression of the ___ segment on EKG when such a patient is exercised.

Digitalis can cause ____________ of the ST segment, however it has a unique, unforgettable appearance (see page 317).

* Chest pain caused by diminished coronary blood flow (without infarction).
Subendocardial Infarction

Subendocardial infarction causes flat depression of the ST segment; however, any significant ST depression (in leads where the QRS is upright) indicates compromised coronary blood flow until proven otherwise.

Subendocardial infarction (often referred to as subendocardial injury) is identified by flat ST _________ depression, which _________ segment may be either horizontal or down-sloping.

**Note:** Subendocardial infarction, a type of “non-Q wave infarction” involves only a small area of myocardium just beneath the endocardial lining. Classical myocardial infarction is said to be transmural; that is, the full thickness of the left ventricular wall is damaged in the infarcted area. Even though subendocardial infarction involves only a small area of the myocardium, it must be respected as a true M.I. that requires appropriate care. A subendocardial M.I. may enlarge or extend and become more life-threatening.

**Note:** Any patient with acute ST depression (or elevation), particularly if it persists, should have an immediate, complete workup including cardiac enzymes.
The Q wave indicates **necrosis** (dead tissue) and makes the diagnosis of infarction.

The diagnosis of myocardial infarction is usually based on the presence of significant ___ waves produced by an area of necrosis in the wall of the left ventricle.

**Note:** The Q wave is the first downward stroke of the QRS complex, and it is never preceded by anything in the complex. In the QRS complex, if there is any positive wave — even a tiny spike — before the downward wave, the downward wave is an S wave (and the upward wave preceding it is an R wave).

Significant Q ______ are absent in normal tracings. We use a capital “Q” to designate a significant Q wave, however “q” (small, lower case q) waves are not significant (see next page).
insignificant q waves

The earliest ventricular depolarization is initiated by Left Bundle Branch fibers at mid-septum (and moves left-to-right)...

... producing tiny q's in some leads.

Normally, ventricular depolarization begins midway down the interventricular septum. Septal depolarization (initiated at mid-septum by the Left Bundle Branch) is left-to-right, and this initial rightward ventricular activation may produce tiny, insignificant q (small q) waves in leads where the QRS is usually upright.

The Right Bundle Branch traverses the septum vertically without branching, however the _____ Bundle Branch gives Left off terminal Purkinje filaments at mid-septum.

So this initial mid-septal depolarization moves left-to-right, away from:

- the positive left arm electrode of lateral leads I and AVL, and...
- the positive left foot electrode of inferior leads II, III, and AVF, and...
- the positive chest electrode of left chest leads V₅ and V₆...

... to occasionally record tiny, insignificant __ waves in those leads. q

Note: This mid-septal depolarization is brief since the efficient ventricular conduction system quickly transmits depolarization to the endocardial surface of both ventricles. So brief is this mid-septal depolarization that only a tiny q wave of less than .04 second is recorded. Insignificant q waves are, by definition, less than one millimeter (.04 sec.) in duration.
A significant Q wave is at least one small square wide (0.04 sec.) or one-third of the entire QRS amplitude. Significant Q waves indicate the necrosis of a myocardial infarction.

Significant Q waves are indicative of the necrosis of a myocardial infarction.

A significant Q wave is one small square (one millimeter) or more wide, and therefore is at least __ second or more in duration.

An old, but persistent, criterion of the significant Q wave is when the Q wave is __________ the amplitude (height and depth) of the entire QRS complex.
When looking at an EKG tracing, note which leads have significant Q waves. Omit lead AVR. Keep in mind the leads that make up the lateral, inferior, and chest lead designations.

To check for an infarction, we scan all leads (except AVR) for the presence of ___________ Q waves.

**Note:** Forget about lead AVR, since this lead is positioned in such a way that data regarding Q waves are unreliable. Lead AVR is like an upside-down lead II, so the large Q waves that are commonly seen in lead AVR are really the upside-down R waves from lead II. Even if you don’t understand the logic behind AVR’s phony Q’s, don’t bother to check it for signs of infarction.

When examining a tracing, either a long strip or mounted, write down those _____ in which you find significant Q* waves, leads ST segment elevation (or depression), and inverted T waves.

*For proper documentation, insignificant q waves should be recorded as well.*
Left ventricular depolarization moves in opposite directions (simultaneously) in opposing walls

Purkinje fibers conduct so rapidly that depolarization is initiated in all endocardial surfaces lining the left ventricle nearly simultaneously. So, depolarization passes from endocardium to epicardium in all left ventricular areas at once.

**Note:** Vectors describe the path of myocardial conduction (endocardium to epicardium), so left ventricular depolarization moves in opposite directions in opposing walls simultaneously.

In the left ventricle, depolarization of the lateral wall moves toward the patient’s left, while depolarization of the medial (septal) wall moves toward the ________.

Depolarization of the anterior left ventricular wall moves anteriorly, while simultaneously, depolarization of the posterior left ventricular wall moves in a ___________ direction.
An infarct is necrotic; it cannot depolarize and has no vectors. So, the positive electrode nearest the infarct detects no “toward” vectors, it sees only the “away” vectors from the opposite wall (through the necrotic void). Therefore, a Q wave is inscribed on EKG in the leads which use that positive electrode for recording.

**Note:** Depolarization moving away from a positive electrode records a negative wave (in this case a Q wave) on EKG.

**Note:** Take your time and visualize each sentence as you read it.

In recording the initial left ventricular depolarization:

- with an *anterior* infarct, the positive (chest) electrode detects only the initial “away” vectors from the opposite side, so a Q is inscribed on EKG in leads V₁ - V₄ which use that positive electrode for recording.

- with a *lateral* infarct, the positive left arm electrode detects only the initial “away” vectors from the opposite side, so a Q is inscribed on EKG in leads I and AVL, which use that positive electrode for recording.

- with an *inferior* infarct, the positive left foot electrode detects only the initial “away” vectors from the opposite side, so a Q is inscribed on EKG in leads II, III, and AVF, which use that positive electrode for recording.
Q waves in $V_1$, $V_2$, $V_3$, or $V_4$ signify an anterior infarction. The infarction in the illustration is definitely acute, because of the ST elevation in all four leads.

**Note:** The chest leads are mainly placed anteriorly, so this is a good way to remember the leads for anterior infarction.

The presence of Q waves in $V_1$, $V_2$, $V_3$, or $V_4$ indicates an infarction in the anterior wall of the _____ ventricle.

**Note:** The anterior portion of the left ventricle includes part of the interventricular septum. Some cardiologists say that when isolated Q waves appear in $V_1$ and $V_2$, the infarction includes the septum, so it is called an antero-septal infarction. Similarly, isolated Q waves in $V_3$ and $V_4$ (more laterally located chest leads) are said to represent an antero-lateral infarction. Remember that (insignificant) q waves are seen normally in $V_5$ and $V_6$.

**Note:** Statistically, anterior infarctions are very deadly, but fortunately, immediate treatment with intravenous thrombolytic medications or angioplasty with stenting has improved the survival rate substantially.
The positive electrode that is used to record the *lateral* limb leads, I and AVL, is on the left arm. The positive electrode used to record the *inferior* limb leads, II, III, and AVF, is on the left “foot.”

The lateral limb leads are I and AVL; they are recorded by a positive left arm electrode.

Yawn... excuse me.

The inferior leads are II, III, and AVF; they are recorded by a positive electrode on the left foot.

**Note:** Yes, it is necessary to have this page here. You’ll see why in just a few seconds.
If there are Q waves in the lateral leads, I and AVL, there is a lateral infarction.

**Note:** Depolarization moving away from a positive electrode records a negative wave (in this case a Q wave) on EKG.

A lateral infarction involves the lateral portion of the _____ ventricle.

In lateral infarction, the positive left arm electrode senses only the initial “away” vectors from the opposite (septal) wall, so it records a Q wave in the lateral _____, I and AVL.

When a lateral infarction occurs, Q waves appear in the lateral leads, which are leads I and ____; Q waves are produced by the initial “away” vectors recorded by the positive left arm electrode through the void of the necrotic lateral infarct.

**Note:** One might abbreviate Lateral Infarction as L.I., which is diagnosed using leads AVL and I.
Inferior infarction is diagnosed by the presence of Q waves in the inferior leads, II, III, and AVF. Check the ST segments to see if this infarction is acute.

**Note:** Depolarization moving away from a positive electrode records a negative wave (in this case a Q wave) on EKG.

The inferior wall of the left ventricle rests upon the diaphragm, so the alternate term “diaphragmatic” infarction is occasionally used instead of "inferior _________."

In inferior infarction, absent the initial “toward” vectors, the positive left foot electrode senses only the initial “away” vectors from the opposite wall, so it records a Q wave in the inferior ________, II, III, and AVF.

An inferior infarction is identified by significant Q waves in inferior leads II, III, and ____; Q waves are produced by the initial “away” vectors recorded by the positive left foot electrode through the void of the necrotic inferior infarct.

**Note:** Autopsy data show that about one-third of inferior infarctions also include portions of the right ventricle.
Depolarization of the anterior wall and depolarization of the posterior wall of the left ventricle are in opposite directions.

**Note:** Left ventricular depolarization may be said to proceed from the *endocardium* (inner lining) to the *epicardium* (outer surface).

Depolarization of the anterior wall of the left ventricle proceeds from the inner endocardium, which lines the ventricle, through the full thickness of the ventricular wall to the outer ventricular surface (__________).

Similarly, depolarization of the posterior wall of the _____ ventricle proceeds from the endocardium to the epicardium.

So, vectors representing depolarization of the anterior and the posterior portions of the left ventricle point in _________ directions.
Acute Anterior Infarction
(note ST elevation)

If an acute anterior infarction produces Q waves and ST elevation in $V_1$ and $V_2$ then a posterior infarction would appear the opposite.

An acute anterior infarction produces significant Q waves with ST ____________ in the first few chest leads.

Considering only $V_1$ and $V_2$ the appearance of significant Q waves and ST elevation indicates an acute ____________ infarction.

**Note:** Acute posterior infarction of the left ventricle would produce the exact opposite to the pattern of acute anterior infarction, because the anterior and posterior walls of the left ventricle depolarize in opposite directions. This will be clarified on the next page.
In acute *Posterior Infarction* there is a large R wave (the opposite of a Q wave) in $V_1$ and $V_2$.

**Note:** In lead $V_1$ a Q wave turned upside-down would look like an R wave (and as you recall, R waves in lead $V_1$ are normally very tiny).

A significant "Q wave" from an infarction in the posterior portion of the ______ ventricle will cause a large R (positive deflection) wave to appear in lead $V_1$.

Suspect a true posterior infarction when you see a large ___ wave in $V_1$ or $V_2$ — even though Right Ventricular Hypertrophy can also produce a large R in $V_1$. 
In acute posterior infarction, there is ST depression (the opposite of the usual ST elevation of injury) in V₁ or V₂.

Acute anterior infarction produces Q waves in the chest leads and the ST segments are elevated.

**Note:** Since the posterior wall of the left ventricle depolarizes in a direction opposite to that of the anterior wall, an acute infarction of the posterior wall causes ST depression in V₁ or V₂.
In summary, acute posterior infarction is characterized by a large R wave and ST depression in V₁ or V₂ (sometimes even in V₃).

**Note:** Always be suspicious of ST segment depression in the right chest leads, for it could indicate an acute posterior infarction. If you do not remember those things that can cause ST depression, look back at page 270. For instance, the diagnosis of an "anterior subendocardial infarction" (because of depressed ST segments in chest leads, see page 271) should be made only with extreme caution, because this ST depression may actually represent an acute true posterior infarct.
If you suspect an acute posterior infarction (large R wave and ST depression in V₁ or V₂), then try “reversed trans-illumination” or the “mirror test.” You must follow the instructions for each test precisely.

**Note:** If acute posterior infarction is suspected because of tall R waves and ST depression in V₁ or V₂ — try reversed trans-illumination or the mirror test. Both of these tests require that you invert the tracing first, then hold the blank (unprinted) back side towards your face.

- **Reversed trans-illumination:** First, invert the EKG tracing, then hold the inverted tracing so that it faces a strong light. Observe the back side of the tracing to check for “Q waves and ST elevation” in the inverted V₁ and V₂ leads.

- **Mirror Test:** First, invert the EKG tracing, then observe it in a mirror. If there is an acute posterior infarction, you will see the classic signs of “Q waves and ST elevation” in the reflection of the inverted V₁ and V₂ leads.

**Note:** With either test, remember to first invert the tracing. Then face the tracing toward a mirror for the mirror test; or for reversed trans-illumination, place the tracing in front of a strong light, viewing the EKG through its back side.
Always Check $V_1$ and $V_2$ for:

1. ST elevation and Q waves
   (Anterior Infarct)

2. ST depression and large R waves
   (Posterior Infarct)

Although posterior infarctions are severe, they are easy to overlook.

When making your routine reading of an EKG, pay special attention to leads $V_1$ and ___ while looking for signs of infarction.

**Note:** ST changes in $V_1$ and $V_2$ are always significant and important… both depression and elevation.

Check for Q waves in $V_1$ and $V_2$ and be sure to observe the height of the ___ waves.

**Note:** And remember how important T wave inversion can be in all leads.
The EKG diagnosis of infarction is generally not valid in the presence of Left Bundle Branch Block.

In Left Bundle Branch Block, the left ventricle (generally, the main chamber to suffer infarction) depolarizes after the \underline{right} ventricle depolarizes.

So any Q wave originating in the left ventricle could not appear at the beginning of the QRS \underline{complex} (with Left BBB); rather, it would fall somewhere in the middle of the QRS complex. In this instance it would be difficult to detect significant Q waves.

\textbf{Note:} One special exception is possible. The right and left ventricles share the interventricular septum in common. So an infarct in the septal area would be shared by the right ventricle, which depolarizes first in Left BBB. This would produce Q waves at the beginning of the wide QRS. Therefore, even in the presence of Left BBB, Q waves in the chest leads might suggest (but not confirm) septal (anterior) infarction.
Locating an infarct is important because treatment modalities and prognosis depend on the location of the infarction.

There are four general locations within the ventricle where infarctions commonly occur.

**Note:** More than one area of the left ventricle may infarct. One infarction may be very old, while another is very recent (acute). So correlate the ST elevation with the appropriate leads both to locate, and to determine the acuteness of each infarct. If ST elevation is present in leads *without* Q waves, “non-Q wave infarction” must be ruled out.

Be careful about diagnosing an infarction in the presence of Bundle Branch Block.

**Note:** Isolated areas of Ischemia (T wave inversion) or ST elevation without Q’s (for non-Q wave infarction) can also be “located” by using the same location criteria.
It is common practice to determine the location of an infarction, but with a little anatomical knowledge of the heart’s coronary blood supply*, we can make a far more sophisticated diagnosis.

There are two coronary arteries that provide the heart with a continuous supply of oxygenated blood.

Quickly review the illustration.
The Left Coronary Artery has two major branches; they are the Circumflex branch and the _______ Descending branch.

The Right Coronary Artery curves around the right ventricle.

* The pulmonary artery has been “surgically” removed in this illustration to show the origin of the coronary arteries at the base of the aorta.
A lateral infarction is caused by an occlusion of the Circumflex branch of the Left Coronary Artery. An anterior infarction is due to an occlusion of the Anterior Descending branch of the Left Coronary Artery.

The Circumflex branch of the Left Coronary Artery distributes blood to the ________ portion of the left ventricle.

The Anterior Descending branch of the Left Coronary Artery supplies blood to the anterior portion of the _____ ventricle.

The Circumflex and the Anterior Descending are the two main branches of the _____ Coronary Artery.
True posterior infarctions are generally caused by an occlusion of the Right Coronary Artery or one of its branches.

The Right Coronary Artery wraps around the right ventricle posteriorly to supply the _________ portion of the left ventricle. posterior

So, a posterior infarction usually is caused by an occlusion of a branch of the _______ Coronary Artery. Right

**Note:** For a long time the Right Coronary Artery was thought to play only a minor role in supplying blood to the heart. Sophisticated techniques of cardiac catheterization and coronary angiography have shown that the Right Coronary Artery usually provides the blood supply to the SA Node, the AV Node, and the His Bundle. It is no wonder that acute posterior infarction is often associated with serious arrhythmias. Wise health care providers treat posterior infarction with concern and respect.
The base of the left ventricle receives its blood supply from branches of either the Right or the Left Coronary Artery, depending on which artery is “dominant.”

Inferior (“diaphragmatic”) infarctions are caused by an occluded terminal branch of either the Right or the _____ Coronary Artery.

So the diagnosis of inferior infarction does not necessarily identify the artery branch that is occluded, unless you have a previous coronary arteriogram (an x-ray highlighting the coronary arteries) to identify which _________ artery supplies the inferior portion of that patient’s left ventricle.

**Note:** Left or Right Coronary “dominance” denotes which coronary artery is the major source of blood supply to the base of the left ventricle. Right Coronary dominance is by far most common in humans.
Hemiblocks are presented in this section (Infarction) because they commonly occur with infarction and an associated diminished blood supply to one of the two divisions of the Left Bundle Branch.

**Note:** The Left Bundle Branch subdivides into two divisions.

The hemiblocks are blocks of either the Anterior or the Posterior Division of the _____ Bundle Branch.

Hemiblocks are commonly due to loss of blood supply to either the Anterior or the Posterior ________ of the Left Bundle Branch.

**Note:** The Right Bundle Branch does not have consistent, named subdivisions of either clinical or electrocardiographic importance (yet).
To understand hemiblocks, you should be familiar with the blood supply to the AV Node and the ventricular conduction system. Follow text and illustration closely.

The Right Coronary Artery usually renders a blood supply* to the AV Node, Bundle of His and a variable twig to the Posterior Division of the Left Bundle Branch.

The Left Coronary Artery also sends a variable twig to the Posterior Division of the Left Bundle Branch.

A total occlusion of the Anterior Descending branch of the Left Coronary Artery may cause a subsequent Right Bundle Branch Block with an Anterior Hemiblock (a block of the Anterior Division of the Left Bundle Branch). Study the illustration carefully.

**Note:** The key to knowing hemiblocks is understanding that an infarction may be due to an occlusion of a vessel at any of numerous locations, and therefore may cause a variety of blocks of the Bundle Branch system. There can be single blocks of a bundle or division, or combinations of these blocks, that spare one or more branches. A coronary obstruction that is not quite complete may cause an *intermittent* block.

* Let’s not forget that the SA Node is usually dependent on the right coronary artery.
Anterior Hemiblock

- LAD - usually assoc. with an M.I. (or other heart disease)
- Normal or slightly widened QRS
- \( Q_1 S_3 \)

*Anterior Hemiblock* refers to a block of the Anterior Division of the Left Bundle Branch, and the above criteria are used in the diagnosis.

The slight delay of conduction to the antero-lateral and superior area of the left ventricle causes (late) unopposed depolarization upward and leftward, recognized on EKG as Left ____ Deviation. Acute LAD is usually what makes you suspect Anterior Hemiblock.

With pure Anterior Hemiblock, the QRS is widened only .10 to .12 sec., but association with other blocks of the Bundle ______ system will widen the QRS more.

Anterior Hemiblock is a block of the Anterior Division of the Left Bundle Branch. Finding a Q in I and a wide and/or deep ___ in III ("\( Q_1 S_3 \)") helps to confirm the diagnosis of \( S \) Anterior Hemiblock.

**Note:** The patient’s previous EKG’s are essential in making a diagnosis of Anterior (or any) Hemiblock. You *must* always rule out pre-existing sources of Left Axis Deviation, e.g., Left Ventricular Hypertrophy, "horizontal heart," or Inferior Infarction.
An occlusion of the Anterior Descending coronary artery produces an Anterior Infarction, and about one-half of these patients develop Anterior Hemiblock. Study the illustration on page 296.

Anterior Hemiblock is a block of the Anterior Division of the Left Bundle Branch, which produces unopposed, late superior-leftward depolarization in the left ventricle, resulting in Left Axis ______.  

Deviation

An occlusion of the Anterior Descending coronary artery will produce an Anterior Infarction, which often causes Anterior ______. (That’s easy to remember.)

Hemiblock

If a patient with an acute Anterior Infarction has an associated Axis change from normal to -60°, you should suspect Anterior ______ (and look for Q, S, Q).  

Hemiblock

But if a patient with an Inferior Infarction develops Left Axis Deviation, don’t jump to hasty conclusions! Inferior Infarction can cause LAD, so ______ Hemiblock may not be the culprit.  

Anterior
An infarction of the anterior wall of the left ventricle (due to an occluded Anterior Descending branch of the Left Coronary Artery) may cause Anterior Hemiblock and Right Bundle Branch Block. Review the illustration on page 296.

**Note:** Don’t forget that the Anterior Descending also renders blood supply to the Right Bundle Branch, so Anterior Infarction may have an associated Right Bundle Branch Block, depending on the location of occlusion.

With Right Bundle Branch Block, the Mean QRS Vector is within the normal range or shows minimal Right Axis ___________.

However, when a patient develops a Right Bundle Branch Block with Left Axis Deviation as well, this is probably caused by Anterior Hemiblock, particularly if there is an acute Anterior ___________.

Infarction
Posterior Hemiblock

- RAD - usually assoc. with an M.I. (or other heart disease)
- Normal or slightly widened QRS
- $S_Q$$_3$

Pure, isolated *Posterior Hemiblock* is rare because the posterior division is short, thick, and commonly has a dual blood supply. See the illustration on page 296.

An inferior infarction may impair the blood supply to the Posterior division of the Left Bundle ________

Posterior Hemiblocks cause Right Axis ________ due to the late, unopposed depolarization forces toward the right.

When Posterior Hemiblock is suspected, look for a deep or unusually wide S in I and Q in III (known as $S_Q$$_3$) to help confirm the ________ of Posterior Hemiblock.
Posterior Hemiblock is always to be respected, and all Inferior Infarctions should be scrutinized to rule it out.

A lateral infarction, either recent or old, can cause Right Axis Deviation, which can be confused with Posterior Hemiblock. In the presence of a ______ M.I., the EKG diagnosis of Posterior Hemiblock is equivocal.

Make certain that by history and previous EKG's, chronic Right Axis Deviation due to slender body build ("vertical heart"), ______ Ventricular Hypertrophy, ______ Right and pulmonary disease, etc. are ruled out.

**Note:** Posterior Hemiblock is serious, and when associated with Right Bundle Branch Block, this combination is considered very dangerous because of the tendency to progress into AV Blocks.

**Important!!** AV Block refers to "atrio-ventricular block", that is, a block between atrial depolarization and ventricular depolarization, so we commonly think of a block in the AV Node or in the His Bundle. However, simultaneous blocks of both Bundle Branches can block AV conduction. Also, RBBB in association with the simultaneous blocks of both divisions of the Left Bundle Branch can produce a block of AV conduction. Please contemplate that for a while.
Yes, you see an *ant* in the Left Axis Deviation quadrant, a memory tool for Left axis shift suggests Anterior Hemiblock. And, that's a "rad" (short for "radiation") sign on a *post* that represents a shift to Right Axis Deviation suggesting Posterior Hemiblock.

For some reason this silly illustration will stick in your mind.

When a patient with a normal axis shifts into an abnormal axis, particularly when associated with a serious cardiac event, we suspect _________.

A shift from normal axis to Right Axis Deviation (remember the "rad post") is characteristic of Posterior _________.

A shift from normal axis to Left Axis Deviation (get that @#$&%!* insect off my illustration!) is characteristic of Anterior _________.

* I wonder how the Japanese translators will deal with that?
Bifascicular Blocks

The word “fascicle” means bundle (bundle of Purkinje fibers), so any main division of the ventricular conduction system is a fascicle. Both Bundle Branches as well as both divisions of the Left Bundle Branch are fascicles.

**Note:** Previously, “Bundle” implied only the Right or the Left Bundle Branch. But now, to avoid confusion for combinations of blocks (e.g., Hemiblock + Bundle Branch Block) we use a more inclusive term, “fascicular” block, to denote a Bundle Branch Block with a Hemiblock.

**Note:** “Bifascicular” block means that two fascicles are blocked. Anterior Hemiblock plus Posterior Hemiblock is clinically the same as Left Bundle Branch Block. So Bifascicular Block generally refers to Right Bundle Branch Block together with a block of either the Anterior Division or the Posterior Division (of the Left Bundle Branch).

**Note:** A block of both the Right and the Left Bundle Branch is a Complete AV Block. Right BBB plus a block of both the Anterior and Posterior Divisions (of the Left Bundle Branch) is also a Complete AV Block. Complete AV Block is very serious since only a ventricular focus remains to slowly pace the ventricles... so slowly that syncope often occurs (airway!), and a patient’s life is at stake.

**Note:** When Bundle Branch Blocks or fascicular blocks are intermittent, we don’t see them continuously on monitor or EKG tracing; just occasionally.
Intermittent Block

Intermittent block of one fascicle:
continuous normal EKG pattern –
• with intermittent wide QRS pattern characteristic
  of the type of intermittent Bundle Branch Block
  present.
• or with intermittent change of QRS Axis
  (i.e., QRS orientation changes intermittently)
  typical of the type of intermittent hemiblock
  present.

Permanent block + Intermittent block:
• continuous EKG signs of one permanent block
  with intermittent EKG signs of another block, as
  long as a third fascicle conducts normally.

Fortunately, combinations of (fascicular) blocks are often intermittent, making them
quite obvious. Intermittent change in QRS axis (e.g., upright QRS’s that transiently
change to downward QRS’s) usually indicates intermittent hemiblock, and a steady
rhythm with transiently widened QRS’s suggests intermittent BBB.

Intermittent block may exist in more than one fascicle in the same
patient, producing a variety of transient changes of ____ Axis
(intermittent [anterior or posterior] hemiblock) or...

...transiently widened QRS’s typical of intermittent (left or right) ____
on EKG or cardiac monitor*. Don’t ignore these intermittent changes;
document them and give proper notification.

Note: Like a failing light bulb that occasionally flickers, sick fascicles
may suffer intermittent block. As a failing, flickering light bulb
eventually burns out, similarly, intermittent fascicular blocks often
warn of impending permanent block of the fascicle. With a pre-existing
permanent block of another fascicle, intermittent fascicular block can
be a timely warning (the only warning!) of an imminent complete block
(that’s why the first word on this page is “fortunately”). In most cases,
permanent block plus intermittent block is an indication for an artificial
pacemaker.

* It is important and challenging to differentiate between intermittent anterior and posterior
hemiblock, as well as intermittent right and left Bundle Branch Block. You know how
already, but a little review wouldn’t hurt.
Considering the three pathways of ventricular depolarization, it becomes apparent that one fascicle must remain functional at least intermittently to provide AV conduction. Early detection allows for early intervention (see page 199).

"Trifascicular" blocks are diagnosed only when one or more of the fascicular blocks is intermittent, otherwise there would be no AV conduction.

Also, the diagnosis of "bilateral" (Left and Right) Bundle Branch Block is made only if one of the Bundles has an intermittent block (or there would be no AV conduction).

Note: If all fascicles are permanently blocked except one that has an intermittent block, then an intermittent Mobitz pattern (occasional non-conduction to the ventricles) will emerge. If that Mobitz pattern becomes more frequent in the tracing, or if a continuous Mobitz 2:1 pattern begins, or worse yet, if there is a continuous high ratio Mobitz block, there is a strong likelihood that complete AV block is imminent and an implantable pacemaker is needed. Knowledge plus vigilance saves lives.

Warning! With Mobitz, every cycle missing its QRS has a regular, punctual P wave — but never a premature P' wave (see Note, page 128). This distinction is critical!

Note: Quickly review from page 295 to this page.
Patients with acute myocardial infarctions are placed in coronary care units and monitored continuously. In most hospitals patients with symptoms (only) of myocardial infarction receive the same cautious care. Patients with no physical symptoms of infarction but with definite EKG criteria of acute infarction ("silent infarction") require admission and monitoring also.

**Note:** Just as medical treatment of arrhythmias changes with the times, so do the attitudes toward indications for artificial pacemakers, angioplasty with stenting, coronary bypass procedures, and thrombolytic treatment. Keep up with the changing standards in your local medical community, read the current literature, and **always know the basics.**

You should always know how to determine the location of an infarction and the vessel(s) involved, as well as their association with ___________.

In patients with myocardial infarction, be alert for subtle changes of Axis (change of QRS orientation in the same lead), and also rhythm changes that may indicate impending ___________ AV block. Vigilance is critical.
Remember that the patient's history and clinical diagnosis are still the most valuable tools you have (using your knowledge and judgment) in determining infarction and infarction-related problems.

The EKG has never become obsolete because it provides more _______ information than any other diagnostic modality.

There is no substitute for obtaining an accurate ________, even if it is volunteered by witnesses to an event.

Although the laboratory provides much useful information, the _____ is an immediate diagnostic gift for those skilled in its interpretation.

**Note:** The value of an EKG increases multifold when it is compared to a patient's previous tracings — get them as soon as possible! Incidentally, is this a photo of Dr. Paul Dudley White, and who is his examining physician with the Elvis sideburns?

**Note:** Review Infarction by turning to the Personal Quick Reference Sheets on pages 342 and 343, and again, look at your simplified methodology (page 334).
You now have the knowledge and certainly the interest and enthusiasm to interpret EKG’s, but always do it methodically. Begin with Rate, then Rhythm, Axis, Hypertrophy, and Infarction. Get accustomed to this routine.

**Note:** In the excitement of an emergency you may be tempted to hunt for Q waves. By breaking the routine you will inevitably miss important diagnostic information - valuable information necessary for the proper treatment of the patient. Keep a cool head and read every EKG properly. Your patients will benefit from your thoroughness.

**Note:** Take one final look at page 334 and review each step of the entire methodology. Then (please) slowly review all of the PQRS pages from 334 to 343. But before you close this book there is some very helpful information in the Miscellaneous chapter. It is next. No, you aren’t done yet.
Chapter 10: Miscellaneous

Before you begin, look at this chapter’s summary on pages 344 to 346.

Miscellaneous Effects

- Pulmonary
- Electrolytes
- Medications
- Artificial Pacemakers
- Heart Transplants

The above effects are common to, but not necessarily diagnostic of, certain conditions or situations that can produce recognizable changes on the EKG.

**Note:** Certain effects may be recognized by their characteristic appearance on the electrocardiogram or on cardiac monitor. For most of the conditions to be discussed in this section, the electrocardiographic signs merely alert us to be aware of existing conditions, certain pathology, or drug or electrolyte effects. But to confirm your suspicion, you should review the medical history, carry out a detailed physical exam, and obtain proper diagnostic tests. Rarely is a diagnosis based entirely on any of the following EKG findings, however they are exceptionally helpful.
Three Important Syndromes
detection can save a life

**Brugada Syndrome**
- RBBB with ST elevation in V₁, V₂, and V₃ (see page 268)
- Susceptible to deadly arrhythmias

**Wellens Syndrome**
- Marked T wave inversion in V₂ and V₃
- Ant. Descending Coronary stenosis

**Long QT Syndrome**
- QT interval longer than 1/3 of the cardiac cycle
- Predisposed to ventricular arrhythmias

Each of these perilous syndromes are easily detected in relatively asymptomatic patients. Routine examination of all EKG’s for these innocuous-looking hallmarks can avoid an inevitable demise. Conventional treatment offers the patient reasonable longevity. The satisfaction of saving a human life is your reward for your vigilance.

Patients with *Brugada Syndrome*, a familial condition, may succumb to deadly arrhythmias; implantation with an ICD can prevent sudden death.

*Wellens Syndrome*, caused by a stenosed anterior descending coronary artery, is easily recognized. Angioplasty with stenting or a coronary bypass graft can remove the imminent peril of impending myocardial infarction.

There are six known forms of (hereditary) *Long QT Syndrome*; these patients are predisposed to dangerous ventricular arrhythmias. A long QT interval exceeds one-half of the cardiac cycle.

**Note:** If everyone who reads this book becomes familiar with, and routinely looks for, these important diagnostic signs, it will serve humanity immensely. A glance at the right chest leads and observing the QT interval is sufficient.
**Chronic Obstructive Pulmonary Disease** (COPD) often produces low voltage amplitude in all leads, and there is usually Right Axis Deviation.

Chronic Obstructive Pulmonary Disease (COPD) commonly produces QRS complexes of small amplitude* in all leads. In fact, all waves in the EKG are minimized in _______.

With COPD, the right ventricle works against considerable resistance, so there is usually some degree of Right Ventricular Hypertrophy and therefore associated _______ Axis Deviation (notice negative QRS’s in lead I).

**Note:** Multifocal Atrial Tachycardia (MAT) is also seen with COPD.

* Low voltage in all leads also appears with hypothyroidism and chronic constrictive pericarditis.
Pulmonary Embolus

- large S wave in lead I
- ST depression in II
- large Q wave in III (with T wave inversion)

With *Pulmonary Embolus* we usually see a large S wave in lead I, and a Q wave and an inverted T wave in lead III ($S_1Q_3L_3$)*.

*S1Q3L3* syndrome characterizes acute *cor pulmonale* resulting from pulmonary embolus. It is called $S_1Q_3L_3$ because of the large S wave in lead I, and there is a Q wave and an inverted T wave in lead III.

**Note:** Notice the typical tendency toward Right Axis Deviation (lead I).

There is usually ST segment __________ in lead II. depression

* Don’t be confused by the inverted T in the printed text. It’s a great memory tool, even if the publisher dislikes it.
Also with pulmonary embolus, there is usually T wave inversion in V₁ through V₄. Often there is Right Bundle Branch Block.

T wave inversion in the chest leads (particularly in leads V₁ through V₄) is a very important diagnostic sign of pulmonary embolus.

Pulmonary embolus may cause Right Bundle Branch Block. This block often subsides after the patient improves.

We can recognize the presence of Right Bundle Branch Block by the R,R’ in the right chest leads.

Note: Occasionally the Right Bundle Branch Block may be "incomplete" (QRS of normal width, but R,R’ is present).
With elevated serum potassium the P wave flattens down, the QRS complex widens, and the T wave becomes peaked.

**Note:** The potassium ion (K⁺) plays an extremely important role in cardiac electrophysiology. The range of normal serum K⁺ concentration is very narrow. In medical parlance we add the suffix “-emia” to the end of the ion name to denote its presence in the blood... but it sounds funny with “potassium.” So its chemical symbol, K, is pronounced verbally, and the prefix “hyper” for increased, or the prefix “hypo” for decreased, is added to communicate deviations from normal. Now you’ll understand both hyper- and hypo- kalemia* (pronounced “kay-LEE-mia”). And that’s right, it’s written “kalemia.” That should help you and also your friends who might be perplexed...

The most striking and classic feature of elevated serum potassium is the ______ T wave.  

The P wave widens and flattens with increased serum potassium, and with extreme hyperkalemia the ___ wave nearly disappears.

When a patient has hyperkalemia, ventricular depolarization takes longer, so the QRS complex ______.  

* The “l” is added to enhance the phonics, so you don’t have to get the “l” out of there (chuckle!).
Potassium

Hypo K⁺

As the serum potassium drops below normal levels, the T wave becomes flat (or inverted) and a U wave appears.

With hypokalemia, as the serum potassium concentration drops, the _wave flattens out, and if the K⁺ concentration drops lower, the T wave inverts.  

**Note:** I always think of the T wave as a tent housing potassium ions. When there is an increase in potassium ions, the tent peaks up, but lowering of potassium ions lowers the height of the tent.

With hypokalemia a _wave appears. This wave becomes more pronounced as the loss of potassium becomes more severe.

**Note:** Potassium is not just “one of those serum electrolytes.” Potassium plays a critical role in repolarization and also in maintaining a precise resting potential. A decrease in potassium makes ventricular automaticity foci extremely irritable. In fact, low potassium can initiate Torsades de Pointes, and it can also evoke dangerous ventricular tachyarrhythmias. Hypokalemia also enhances the toxic effects of digitalis excess.
With hypercalcemia, the QT interval shortens; however hypocalcemia prolongs the QT interval.

**Note:** Since you already understand "hyper-" and "hypo-", I only need mention that "-calcemia" is pronounced "cal-SEE-mia".

Hypocalcemia will prolong the ___ interval.  

**Note:** The QT interval is measured from the beginning of the QRS complex to the end of the T wave. Normally, the QT interval should be less than half of the cycle length.

An increase in calcium (Ca++) ions accelerates both ventricular depolarization and ventricular repolarization. This is manifested as a short QT _________.  

**Note:**
Digitalis causes a gradual downward curve of the ST segment, to give it the appearance of Salvador Dali's mustache. Notice that the lowest portion of the ST segment is depressed below the baseline.

Digitalis produces a unique, gradual downward curve of the ___ segment; this is the classical "digitalis effect."

**Note:** To identify the classical pattern of digitalis effect, you should observe a lead with no demonstrable S wave. The downward portion of the R wave gradually thickens as it curves down into the ST segment, which is usually depressed. The downward limb of the R wave has a gentle, curving slope that gradually blends into the depressed ST segment. Look for it the next time you have a patient on a digitalis preparation.

**Note:** Digitalis in therapeutic doses has a parasympathetic effect. With a Sinus Rhythm, digitalis slows the SA Node pacing rate. Conduction through the AV Node is slowed, and digitalis also inhibits the AV Node's receptiveness to multiple stimuli, allowing fewer stimuli to reach the ventricles (necessary with Atrial Flutter and Atrial Fibrillation) to permit a more physiological and more efficient ventricular response rate. Digitalis has a very narrow range of therapeutic effectiveness, and should this therapeutic range be exceeded, a multitude of undesirable effects can result. See the next two pages...
Excess Digitalis

- atrial & Junctional premature beats
- PAT with block
- Sinus block
- AV blocks

Excess digitalis tends to cause AV Blocks of many varieties, and may even induce Sinus (SA) Block.

**Note:** Supraventricular (particularly atrial) foci are exceptionally sensitive to digitalis, so premature atrial beats (PAB's) are often the earliest warning sign that your patient has elevated levels of digitalis. Atrial automaticity foci are very effective digitalis sensors.

Digitalis in excess may cause transient Sinus _______. Block

Digitalis retards conduction of depolarization through the AV Node; and in excess, it can cause various types of ___ Block, particularly rate-dependent AV Block. AV

Automaticity foci of the atria and the AV Junction can become irritable when __________ preparations are present in excessive concentrations in your patient.

**Note:** Low serum potassium can enhance the toxicity of digitalis, so that digitalis, even in therapeutic concentrations, can produce undesirable signs of toxicity if the serum potassium is low.


**Digitalis Toxicity**

- atrial & Junctional tachy-arrhythmias
- PVC's
- Ventricular Bigeminy, Trigeminy
- Ventricular Tachycardia
- Ventricular Fibrillation

Atrial and Junctional automaticity foci are very likely to become irritable in the presence of excessive digitalis. In fact, marked digitalis toxicity can even provoke ventricular foci into rapid and dangerous rhythms.

The foci of the atria and AV Junction are most sensitive to excessive digitalis, but with marked digitalis toxicity, even ventricular foci may become so irritable that they spontaneously emit PVC's.

Marked digitalis toxicity can make ventricular foci so irritable that they may suddenly fire multiple discharges that initiate dangerous ventricular tachy-arrhythmias.

**Note:** Digitalis preparations have been used medicinally by civilized people since the thirteenth century. But like most other cardiac medications, in certain circumstances or in high concentrations, digitalis can induce deadly arrhythmias.
Quinidine causes widening of the P wave and widening of the QRS complex. There is often ST depression with a prolonged QT. The presence of U waves is typical as well.

**Note:** Quinidine retards depolarization and repolarization through both the atrial and the ventricular myocardium. Most of the effects of quinidine that we see on EKG relate to its pharmacological effects on sodium and potassium ion channels.

Quinidine causes a wide, notched ___ wave on EKG, and the QRS complex is also widened.

Quinidine prolongs the ___ interval, and depresses the ST segment. Look for U waves (which represent delayed repolarization of the ventricular conduction system).

**Note:** Episodes of Torsades de Pointes – a rapid and dangerous ventricular rhythm can result from quinidine toxicity (see page 158).
Artificial pacemakers have a pulse generator with a long-lasting lithium battery. The pacemaking stimuli are designed for ventricular or atrial (or both) pacing modalities, and a wide variety of sensing features are available.

**Note:** Artificial pacemakers are surgically implanted as a permanent pacemaking source. Originally, they were designed to counter the bradycardia that attends Complete AV Block and Sick Sinus Syndrome. Now, the uses and variety of pacemaker types is well beyond the scope of this book, so we will review only basic principles of artificial cardiac pacing. In most cases the electrode lead wire is passed transvenously into the right side of the heart; however, sometimes the stimulating electrode is surgically attached to the epicardial surface of the heart.

The pacemaker generator emits regular pacing stimuli, which record on the EKG as a narrow vertical spike.

The pacemaker emits regular, paced electrical stimuli and each stimulus should "capture" (i.e., depolarize) the myocardial tissue in contact with the electrode. The depolarization stimulus then conducts through the myocardium.
The demand feature of many artificial pacemakers is designed to imitate the physiological mechanisms of an automaticity focus (great idea!). The demand pacemaker is programmed with an "inherent rate" that is overdrive-suppressed by normal Sinus pacing.

**Note:** The illustration depicts the EKG of a demand pacemaker with a ventricular sensing electrode and a ventricular pacing electrode.

A demand pacemaker is *overdrive-suppressed* by normal Sinus pacing, but should the Sinus rate drop below the pacemaker’s programmed inherent rate, the pacemaker, no longer overdrive-suppressed, escapes to assume pacemaking responsibility at its inherent rate.

But if the SA Node resumes pacing at a normal rate (which is faster than the inherent rate of the demand pacemaker) the demand pacemaker is overdrive-suppressed and stops pacing.

The demand pacemaker is designed to *reset* just like an automaticity focus. When the demand pacemaker senses a PVC, it resets its pacing (at the cycle length of its inherent rate) in step with the PVC. This provides for uninterrupted cardiac function (clever engineers design to imitate Nature).
Contemporary pacemakers offer many features that can be used to treat many types of cardiac dysfunction and pathology.

With failure of the SA Node, Atrial Pacing can be used when the AV Node and ventricular conduction system function normally, so the artificially paced atrial stimuli are properly conducted from the atria to the ventricles.

A complete AV block prevents normal Sinus pacing from conducting to the ventricles and may require P wave triggered pacing*, which senses the patient's ___ wave, then after a brief pause (imitating normal AV conduction) it generates a stimulus for ventricular depolarization.

SA Node malfunction combined with complete AV block sometimes necessitates A-V sequential pacing, which provides a stimulus for atrial depolarization followed by a brief pause, then the ventricles are depolarized.

**Note:** Modern pacemakers are computerized wonders that can detect and respond to physiological needs such as decreased rate during sleep and increased rate during exercise.

* Also called “atrial synchronous” or “atrial tracking” pacing.
Possible pacemaker electrode positions in the right ventricle

Usually a right ventricular electrode is used for cardiac pacemaking; the electrode tip of the lead is positioned within the cavity of the right ventricle. Three possible catheter lead positions are shown with the way they record on EKG.

**Note:** The ideal location of the tip electrode of a right ventricular pacemaker, is in the apex of the right ventricle. The resultant QRS complex has a Left Bundle Branch Block pattern with Left Axis Deviation.

When a paced QRS shows a LBBBB pattern with a normal axis, the electrode tip is in the mid-inflow tract of the right ________.

But if you notice a paced QRS with a LBBBB pattern and Right Axis Deviation, the tip of the ________ is just below the pulmonic valves.

**Note:** Certain cardiac patients may have a surgically implanted “pacemaker”, called an *Implantable Cardioverter Defibrillator* (ICD, see next page) that can pace, detect and interpret rhythm disturbances, and treat tachyarrhythmias by overdrive pacing or cardioversion, even defibrillate in the event of ventricular fibrillation. Oh, Brave New World!
The Implantable Cardioverter Defibrillator (ICD) is a self-contained, computerized device that can instantly analyze and treat most dangerous cardiac arrhythmias. It can simulate normal sinus pacing, institute overdrive (suppression) pacing to treat ventricular tachycardia, provide cardioversion, and even defibrillate VF.

The ICD can detect and treat certain arrhythmias with cardioversion (a precisely timed electrical shock), and...

it can diagnose ventricular tachycardia and respond with overdrive pacing to suppress the causative ventricular ______. focus

The ICD can detect ventricular fibrillation and instantly defibrillate the heart, and...

should the SA Node perform sluggishly after defibrillation, the ICD will provide pacemaking stimuli at a physiological _____.

Note: The ICD is a technological masterpiece!
There is an external non-invasive pacemaking device that effectively delivers pacing stimuli to the heart through intact skin in emergency situations.

Sophisticated pacemakers are available that can painlessly pace the heart through the intact skin. These external, non-invasive pacemakers are ideal for temporary pacing.

Pacing the heart through the body surface requires an impulse of longer duration than that of intracardiac pacemakers so each pacing spike is wide with a flat end.

Note: Another externally applied emergency device, the Automated External Defibrillator (AED), records and analyses the patient’s EKG, and then automatically defibrillates the patient if a deadly arrhythmia is detected. The AED is very accurate in its computerized recognition of Ventricular Fibrillation and high rate Ventricular Tachycardia; it is easily operated by moderately trained personnel. Numerous trials and studies have proven the AED to be a very effective method of defibrillation in a non-hospital setting. See page 170.
A heart transplant procedure leaves portions of the recipient patient’s “native” atria in place. These portions of atria contain the patient’s own SA Node, so the transplant patient has his native SA Node, plus the SA Node of the donor heart.

**Note:** To expedite these procedures, the portions of the native atria that contain the large vessel orifices are left behind to be sutured to the atria of the transplanted heart. So the recipient patient retains the native SA Node, and the donor heart that the patient receives also has a functioning SA Node.

Transplant patients therefore have two SA Nodes, each producing ___ waves.

The native SA Node produces depolarizations (P$_n$) that do not pass beyond the suture line, so they do not depolarize the donor _____.

The transplanted “donor” heart has its own functional SA Node that remains the dominant pacemaker, so all of its P waves (P$_d$) are followed by ___ complexes.
A heterotopic heart transplant is a procedure that leaves the native heart in place, while a temporary, donor heart is surgically attached to assist the pumping effort.

In order to assist in pumping, a heterotopic heart transplant gives the patient (temporarily) two ________.

So the EKG in this temporary, emergency situation displays the simultaneous recording of the electrical activity of two separate ________.

**Note:** With great advances in medical technology and increasing sophistication of biomechanical engineering, attempts are constantly being made to devise an efficient artificial heart. It is unlikely that a totally artificial heart will ever approach the efficacy and safety of that Designed by Nature.

Let me know if your understanding was a kind of ecstacy. 
(It has been for me.) –DD
Cardiac Monitor Displays

Cardiac monitors display the same information as recorded on a standard 12 lead EKG. Some initial apprehension may arise because of lack of familiarity with the display. The EKG tracing is in bright green on a black background, and the amplitude of waves (height and depth) is increased. Because the "leads" of a cardiac monitor are modifications of standard leads with exaggerated amplitudes to aid in visualization at a distance, voltage (height and depth) criteria can not be utilized. But don't despair, this is just another method of displaying the heart's electrical activity... and familiarity eventually breeds content.

1° AV block

3:2 Wenckebach (2° AV block)

3:1 Mobitz (2° AV block)

multifocal PVC's

ventricular bigeminy

run of 3 PVC's (Ventricular Tachycardia)

Ventricular Tachycardia

Ventricular Fibrillation
Electrocardiography was your challenge; knowledge, your achievement.

Now that you are certainly pleased with your understanding of basic electrocardiography, and proud of your ability to interpret the information on EKG's and cardiac monitors, you realize how logical and marvelously designed is the heart.

You're probably ready for Ion Adventure in the Heartland, Dr. Dubin's highly-acclaimed, entertaining text. For your giant leap into the 21st century, your knowledge needs to be on a molecular level, so let Dr. Dubin be your guide and simplify your understanding.

**Ion Adventure in the Heartland** is an exciting full-color expedition deep into the secret molecular wonderland of cardiac physiology with a splash of biochemistry. We will explore the vivid inner world of the magnificent "ion movers," a dynamic microcosm of exotic ion channels, ion pumps, ion exchangers, the mysterious connexons, and the fast moving ions that they control. You will be immersed in this never-before-seen, living wonderland that generates the heart's electrical energy and power in response to physiological demands. What a performance to behold, as we expose the private activities of these ion movers, exquisitely orchestrated by the autonomic nervous system. The book is narrated in Dr. Dubin's entertaining, easy to understand style. You will discover what really makes the healthy heart tick, yet falter with stress and disease. Though this adventure is not likely to become a great movie, it is an easy to understand, illustrated story of the intimate lifestyles of the ions and their movers as recorded by the surface EKG.

To learn more:

[www.IonAdventure.com](http://www.IonAdventure.com)

COVER Publishing Company
P.O. Box 1092, Tampa, FL 33601
U. S. A.
Ion Adventure in the Heartland, is an exciting adventure in living color, providing vital knowledge for the medical profession in millennium 2000.

Scientists and researchers in the twentieth century found the microstructure of the cells of the heart to be an engineering wonder. Research continues to reveal intriguing information, while raising many new questions. Current concepts may seem complex—even intimidating—to medical professionals, although, in reality, they are easy to understand.

The key to cardiac function is at the ionic-molecular level, where autonomic control occurs, and where medications work. All the electrical and mechanical properties of the heart are due to the movement of only three types of ions...yes, three little ions!

Let me have them come forward to introduce them by name:

Sodium ion (Na⁺)  Calcium ion (Ca⁺⁺)  Potassium ion (K⁺)
Ion-moving ("ion-kinetic") structures of the cell membrane (and cell interior) produce ion movement. Most of these structures are sophisticated molecular portals that employ precision mechanisms to control and regulate the movement of Na⁺, Ca²⁺, and K⁺ ions. Each variety of ion-kinetic structure has its own unique behavior.

We are launching an expedition to explore this incredible ionic-molecular microcosm to learn just how these mechanisms move Na⁺, Ca²⁺, and K⁺ ions to govern the heart's function. We would love to have you join us on this fascinating adventure.

Our fantastic journey is narrated by a five year old boy named Dale, so certainly anyone who has read *Rapid Interpretation of EKG’s* can easily master this vital medical knowledge, which is so useful and necessary in millennium 2000.

Hurry... your knowledge is needed!
Personal Quick Reference Sheets

(pages 333 to 346)

from: Rapid Interpretation of EKG's

by Dale Dubin, MD
COVER Publishing Co., P.O. Box 1092, Tampa, FL 33601, USA

The owner of this book may remove pages 333 through 346 to carry as a personal quick reference. The entire text and all illustrations of Rapid Interpretation of EKG's are fully protected by domestic United States copyright as well as the Universal Copyright Convention, and all rights of absolute imprimatur are enforced by COVER Publishing Co.

RAPID INTERPRETATION OF EKG's

Dr. Dubin's classic, simplified methodology for understanding EKG's

6th Ed.

Dale Dubin, MD

May humanity benefit from your knowledge.

Dale Dubin

Learning Web Sites:

Physicians and medical students: www.theMDsite.com
Nurses and nurses in training: www.CardiacMonitors.com
Emergency medical personnel: www.EmergencyEKG.com
1. RATE (pages 65-96)
   Say "300, 150, 100" ... "75, 60, 50"
   - but for bradycardia:
     rate = cycles/6 sec. strip x 10

2. RHYTHM (pages 97-202)
   Identify the basic rhythm, then scan tracing for prematurity, pauses, irregularity, and abnormal waves.
   - Check for: P before each QRS.
     QRS after each P.
   - Check: PR intervals (for AV Blocks).
     QRS interval (for BBB).
   - If Axis Deviation, rule out Hemiblock.

3. AXIS (pages 203-242)
   - QRS above or below baseline for Axis Quadrant
     (for Normal vs. R. or L. Axis Deviation).
     For Axis in degrees, find isoelectric QRS in a limb lead
     of Axis Quadrant using the "Axis in Degrees" chart.
   - Axis rotation in the horizontal plane: (chest leads)
     find "transitional" (isoelectric) QRS.

4. HYPERTROPHY (pages 243-258)
   Check:
   \[
   \begin{align*}
   V_1 & : P \text{ wave for atrial hypertrophy.} \\
        & R \text{ wave for Right Ventricular Hypertrophy.} \\
   \end{align*}
   \]
   S wave depth in \( V_1 \) ...
   + R wave height in \( V_5 \) for Left Ventricular Hypertrophy.

5. INFARCTION (pages 259-308)
   Scan all leads for:
   - Q waves
   - Inverted T waves
   - ST segment elevation or depression
   Find the location of the pathology (in the Left ventricle),
   and then identify the occluded coronary artery.
Determine Rate by Observation (pages 78-88)

Fine division/rate association: reference (page 89)

Bradycardia (slow rates) (pages 90-96)
- Cycles/6 second strip × 10 = Rate
- When there are 10 large squares between similar waves, the rate is 30/minute.

Sinus Rhythm: origin is the SA Node ("Sinus Node"),
normal sinus rate is 60 to 100/minute.
- Rate more than 100/min. = Sinus Tachycardia (page 68).
- Rate less than 60/min. = Sinus Bradycardia (page 67).

Determine any co-existing, independent (atrial/ventricular) rates:
- Dissociated Rhythms: (pages 155, 157, 186-189)
  A Sinus Rhythm (or atrial rhythms) may co-exist with an independent rhythm from an automaticity focus of a lower level. Determine rate of each.

Irregular Rhythms: (pages 107-111)
- With Irregular Rhythms (such as Atrial Fibrillation) always note the general (average) ventricular rate (QRS's per 6-sec. strip × 10) or take the patient's pulse.
Rhythm (pages 97 to 111)

from: Rapid Interpretation of EKG's
by Dale Dubin, MD
COVER Publishing Co., P.O. Box 1092, Tampa, FL 33601, USA

★ Identify basic rhythm...
...then scan entire tracing for pauses, premature beats, irregularity, and abnormal waves.

★ Always:
- Check for: P before each QRS.
  QRS after each P.
- Check: PR intervals (for AV Blocks).
  QRS interval (for BBB).
- Has QRS vector shifted outside normal range? (to rule out Hemiblock).

Irregular Rhythms (pages 107-111)

Sinus Arrhythmia (page 100)
Irregular rhythm that varies with respiration.
All P waves are identical.
Considered normal.

Wandering Pacemaker (page 108)
Irregular rhythm. P waves change shape as pacemaker location varies.
Rate under 100/minute...

...but if the rate exceeds 100/minute, then it is called

Multifocal Atrial Tachycardia
(page 109)

Atrial Fibrillation (pages 110, 164-166)
Irregular ventricular rhythm.
Erratic atrial spikes (no P waves) from multiple atrial automaticity foci. Atrial discharges may be difficult to see.
**Escape** (pages 112-121) – the heart's response to a pause in pacing

- An unhealthy Sinus (SA) Node may fail to emit a pacing stimulus ("Sinus Block"); this pause may evoke an escape beat from an automaticity focus.
- But a sick Sinus (SA) Node may cause an automaticity focus to "escape" to assume pacemaker status.
- The SA Node usually resumes pacing.

**Premature Beats** (pages 122-145) – from an irritable automaticity focus

- An irritable automaticity focus may suddenly discharge, producing a premature atrial beat (pages 124-130).
- Premature junctional beat (pages 131-133).
- Premature ventricular contraction (pages 135-141). PVC's may be: multiple, multifocal, in runs, or coupled with normal cycles.
Tachyarrhythmias (pages 146-172), “focus” = automaticity focus

**Paroxysmal Tachycardia**
- **Paroxysmal Atrial Tachycardia**
  - An irritable atrial focus discharging at 150-250/min. produces a normal wave sequence, if P waves are visible. (page 149)
  - *P.A.T. with block*
    - Same as P.A.T. but only every second (or more) P wave produces a QRS. (page 150)
- **Paroxysmal Junctional Tachycardia**
  - AV Junctional focus produces a rapid sequence of QRS-T cycles at 150-250/min. QRS may be slightly widened. (pages 151-153)
- **Paroxysmal Ventricular Tachycardia**
  - Ventricular focus produces a rapid (150-250/min.) sequence of (PVC-like) wide ventricular complexes. (pages 154-158)

**Flutter**
- **Atrial Flutter**
  - A continuous (“saw tooth”) rapid sequence of atrial complexes from a single rapid-firing atrial focus. Many flutter waves needed to produce a ventricular response. (pages 159, 160)
- **Ventricular Flutter** (pages 161, 162) also see “Torsades de Pointes” (pages 158, 345)
  - A rapid series of smooth sine waves from a single rapid-firing ventricular focus; usually in a short burst leading to Ventricular Fibrillation.

**Fibrillation**
- **Atrial Fibrillation** (pages 110, 164-166)
  - Multiple atrial foci rapidly discharging produce a jagged baseline of tiny spikes. Ventricular (QRS) response is irregular.
- **Ventricular Fibrillation** (pages 167-170)
  - Multiple ventricular foci rapidly discharging produce a totally erratic ventricular rhythm with unidentifiable waves. Needs immediate treatment.
Personal Quick Reference Sheets

Rhythm: ("heart") blocks (pages 173 to 202)

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Sinus (SA) Block (page 174)
An unhealthy Sinus (SA) Node misses one or more cycles (sinus pause)...
the Sinus Node usually resumes pacing, but the pause may evoke an "escape" response from an automaticity focus. (pages 119-121)

AV Block (pages 176-189)
Blocks that delay or prevent atrial impulses from reaching the ventricles.

1o AV Block ...prolonged PR interval (pages 176-178).
PR interval is prolonged to greater than .2 sec (one large square).

2o AV Block ... some P waves without QRS response (page 179-185)
Wenckebach ... PR gradually lengthens with each cycle until the last P wave in the series does not produce a QRS.
Mobitz ... some P waves don't produce a QRS response. If "intermittent," an occasional QRS is dropped.
More advanced Mobitz block may produce a 3:1 (AV) pattern or even higher AV ratio (page 181).

2:1 AV Block ... may be Mobitz or Wenckebach.
(pages 182, 183) PR length and QRS width or vagal maneuvers help differentiate.

3o ("complete") AV Block ... no P wave produces a QRS response (pages 186-190)

Bundle Branch Block ... find R', R" in right or left chest leads (pages 191-202)

Right BBB (pages 194-196)

Left BBB (pages 194-197)

Caution: With Left BBB infarction is difficult to determine on EKG.

Hemiblock ... block of Anterior or Posterior fascicle of the Left Bundle Branch.

Anterior Hemiblock
Axis shifts Leftward → L.A.D.
look for Q, S,
(pages 297-299)

Posterior Hemiblock
Axis shifts Rightward → R.A.D.
look for S, Q,
(pages 300-302)
**Axis** (pages 203 to 242)

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### General Determination of Electrical Axis (pages 203-242)

Is QRS positive (\(\uparrow\)) or negative (\(\downarrow\)) in leads I and AVF?

**Is Axis Normal?** (page 227)

<table>
<thead>
<tr>
<th>QRS in lead I (pages 215-222)</th>
<th>First Determine Axis Quadrant (pages 214-231)</th>
</tr>
</thead>
<tbody>
<tr>
<td>...if the QRS is Positive (mainly above baseline), then the Vector points to positive (patient's left) side.</td>
<td></td>
</tr>
<tr>
<td><strong>Normal:</strong> QRS upright in I and AVF</td>
<td></td>
</tr>
<tr>
<td><strong>“two thumbs-up” sign</strong></td>
<td></td>
</tr>
<tr>
<td>QRS in lead AVF (pages 223-226)</td>
<td></td>
</tr>
<tr>
<td>...if the QRS is mainly Positive, then the Vector must point downward to positive half of the sphere.</td>
<td></td>
</tr>
</tbody>
</table>

---

### Axis in Degrees (pages 233, 234) (Frontal Plane)

After locating Axis Quadrant, find limb lead where QRS is most isoelectric:

<table>
<thead>
<tr>
<th>Extreme Right Axis Deviation</th>
<th>Left Axis Deviation</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>lead</strong></td>
<td><strong>Axis</strong></td>
</tr>
<tr>
<td>I</td>
<td>(-90^\circ)</td>
</tr>
<tr>
<td>AVL</td>
<td>(-120^\circ)</td>
</tr>
<tr>
<td>III</td>
<td>(-150^\circ)</td>
</tr>
<tr>
<td>AVF</td>
<td>(-180^\circ)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Right Axis Deviation</th>
<th>Normal Range</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>lead</strong></td>
<td><strong>Axis</strong></td>
</tr>
<tr>
<td>AVF</td>
<td>0(^\circ)</td>
</tr>
<tr>
<td>II</td>
<td>(+30^\circ)</td>
</tr>
<tr>
<td>AVR</td>
<td>(+120^\circ)</td>
</tr>
<tr>
<td>I</td>
<td>(+90^\circ)</td>
</tr>
</tbody>
</table>

---

### Axis Rotation (left/right) in the Horizontal Plane (pages 236-242)

Find transitional (iselectric) QRS in a chest lead.

- **transitional QRS is “isoelectric”**
- **Rightward rotation**
- **Normal Range**
- **Leftward rotation**
- **Patient's Right**
- **Normal Range**
- **Patient's Left**
Hypertrophy (pages 243 to 258)

from: Rapid Interpretation of EKG’s
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Atrial Hypertrophy (pages 245-249)

Right Atrial Hypertrophy (page 248)
- large, diphasic P wave with tall initial component

Left Atrial Hypertrophy (page 249)
- large, diphasic P wave with wide terminal component

Ventricular Hypertrophy (pages 250-258)

Right Ventricular Hypertrophy (pages 250-252)
- R wave greater than S in V1, but R wave gets progressively smaller from V1 - V6.
- S wave persists in V5 and V6.
- R.A.D. with slightly widened QRS.
- Rightward rotation in the horizontal plane.

Left Ventricular Hypertrophy (pages 253-257)

\[
\begin{align*}
S \text{ wave in } V_1 \text{ (in mm.)} \\
+ \quad R \text{ wave in } V_6 \text{ (in mm.)} \\
\text{Sum in mm. is more than 35 mm. with L.V.H.}
\end{align*}
\]

- L.A.D. with slightly widened QRS.
- Leftward rotation in the horizontal plane.

Inverted T wave:
- slants downward gradually,
- but up rapidly.
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**Infarction** (pages 259 to 308)

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**Q wave = Necrosis (significant Q’s only)** (pages 272-284)

- Significant Q wave is one millimeter (one small square) wide, which is .04 sec. in duration...
  
  ... or is a Q wave 1/3 the amplitude (or more) of the QRS complex.

- Note those leads (omit AVR) where significant Q’s are present...
  
  ... see next page to determine infarct location, and to identify the coronary vessel involved.

- Old infarcts: significant Q waves (like infarct damage) remain for a lifetime. To determine if an infarct is acute, see below.

---

**ST (segment) elevation = (acute) Injury (pages 266-271) (also Depression)**

- Signifies an acute process, ST segment returns to baseline with time.

- ST elevation associated with significant Q waves indicates an acute (or recent) infarct.

- A tiny “non-Q wave infarction” appears as significant ST segment elevation without associated Q’s. Locate by identifying leads in which ST elevation occurs (next page).

- ST depression (persistent) may represent “subendocardial infarction,” which involves a small, shallow area just beneath the endocardium lining the left ventricle. This is also a variety of “non-Q wave infarction.” Locate in the same manner as for infarction location (next page).

---

**T wave inversion = Ischemia (pages 264, 265)**

- Inverted T wave (of ischemia) is symmetrical (left half and right half are mirror images). Normally T wave is upright when QRS is upright, and vice versa.

- Usually in the same leads that demonstrate signs of acute infarction (Q waves and ST elevation).

- Isolated (non-infarction) ischemia may also be located; note those leads where T wave inversion occurs, then identify which coronary vessel is narrowed (next page).

---

NOTE: Always obtain patient’s previous EKG’s for comparison!
Infarction Location — and — Coronary Vessel Involvement (pages 259 to 308)

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Coronary Artery Anatomy (page 291)

Right Coronary Artery

Left Coronary Artery

circumflex

anterior descending

Infarction Location/Coronary Vessel Involvement (pages 278-294)

**Posterior**
- large R with ST depression in V₁ & V₂
- mirror test or reversed transillumination test (Right Coronary Artery) (pages 282-286)

**Laterali**
Q's in lateral leads I and AVL (Circumflex Coronary Artery) (pages 280, 292)

**Inferior**
(diaphragmatic)
Q's in inferior leads II, III, and AVF (R. or L. Coronary Artery) (pages 281, 294)

**Anterior**
Q's in V₅, V₆, V₇, and V₈ (Anterior Descending Coronary Artery) (pages 278, 292)
Pulmonary Embolism (pages 312, 313)
- $S_1Q_3L_5$ - wide $S$ in I, large $Q$ and inverted $T$ in III
- acute Right BBB (transient, often incomplete)
- R.A.D. and clockwise rotation
- inverted $T$ waves $V_1 \rightarrow V_6$ and ST depression in II

Artificial Pacemakers (pages 321-326)
Modern artificial pacemakers have sensing capabilities and also provide a regular pacing stimulus. This electrical stimulus records on EKG as a tiny vertical spike that appears just before the "captured" cardiac response.

Demand Pacemakers:
- are "triggered" (activated) when the patient's own rhythm ceases or slows markedly.
- are "inhibited" (cease pacing) if the patient's own rhythm resumes at a reasonable rate.
- will "reset" pacing (at same rate) to synchronize with a premature beat.

Pacemaker Impulse (delivery modes)

- Ventricular Pacemaker (page 323) (electrode in Right Ventricle)
- (Asynchronous) Epicardial Pacemaker
  Ventricular impulse not linked to atrial activity.

- Atrial Pacemaker (page 323)

- Atrial Synchronous Pacemaker (page 323)
P wave sensed, then after a brief delay, ventricular impulse is delivered.

- Dual Chamber (AV sequential) Pacemaker

- External Non-invasive Pacemaker (page 326)
**Electrolytes**

*Potassium* (pages 314, 315)
- Increased K⁺ (page 314) (hyperkalemia)
- Decreased K⁺ (pages 315) (hypokalemia)

*Calcium* (page 316)
- Hyper Ca++
- Hypo Ca++

**Digitalis** (pages 317-319)
- EKG appearance with digitalis ("digitalis effect")
  - remember Salvador Dali.
  - T waves depressed or inverted.
  - QT interval shortened.

**Digitalis Excess** → **Digitalis Toxicity**
- SA Block
- P.A.T. with Block
- AV Blocks
- AV Dissociation
- (irritable foci firing rapidly)
- Atrial Fibrillation
- Junctional or Ventricular Tachycardia
- multiple PVC's
- Ventricular Fibrillation

**Quinidine** (page 320)
- EKG appearance with quinidine (page 320)
- Excess quinidine or other medications that block potassium channels (or even low serum potassium) may initiate Torsades de Pointes (page 158)
Personal Quick Reference Sheets

Practical Tips

from: Rapid Interpretation of EKG’s
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Dubin’s Quickie Conversion
—for—
Patient’s Weight from Pounds to Kilograms

Patient wt. in kg. = Half of patient’s wt. (in lb.) minus 1/10 of that value.

Examples:
- 180 lb. patient (becomes 90 minus 9) is 81 kg
- 160 lb. patient (becomes 80 minus 8) is 72 kg
- 140 lb. patient (becomes 70 minus 7) is 63 kg.

Modified Leads
—for—
Cardiac Monitoring

Locations are approximate. Some minor adjustment of electrode positions may be necessary to obtain the best tracing. Identify the specific lead on each strip placed in the patient’s record.

<table>
<thead>
<tr>
<th>Sensor Electrode</th>
<th>Identification</th>
</tr>
</thead>
<tbody>
<tr>
<td>+</td>
<td>Letter</td>
</tr>
<tr>
<td>-</td>
<td>Color (inconsistent)</td>
</tr>
<tr>
<td>G*</td>
<td>R (or RA)</td>
</tr>
<tr>
<td></td>
<td>red</td>
</tr>
<tr>
<td></td>
<td>L (or LA)</td>
</tr>
<tr>
<td></td>
<td>white</td>
</tr>
<tr>
<td></td>
<td>G (or RL)</td>
</tr>
<tr>
<td></td>
<td>variable</td>
</tr>
</tbody>
</table>

* Ground, Neutral or Reference

Modified Lead I

Modified Lead II

Conventional Lead

MCI, To make this MCI, move * electrode to same (mirror) position on the patient’s left chest.
This section contains EKG tracings (and their interpretation) from various patients. The tracings and interpretations are provided so that you can see how this method of reading EKG’s actually works. Try these few examples so that you grow accustomed to this systematic approach. Once you learn how to read an EKG systematically, you will soon become very skilled at routine EKG interpretations.
Patient D.D. is a 29 year old white male known to be a hypochondriac with numerous complaints.
EKG Tracings

EKG Interpretation

Patient: D.D.
Rate: about 70/minute
Rhythm: Regular Sinus Rhythm
PR less than .2 sec. (No AV Block).
QRS less than .12 sec. (No BBB).
...but note the R',R' in III suggesting incomplete
Bundle Branch Block.
Axis: Normal Range (about +30°).
Rightward rotation in the horizontal plane.
Hypertrophy: No atrial hypertrophy.
No ventricular hypertrophy.
Infarction: No significant Q waves.
ST segments: not elevated, except for V₄ and V₉ where
ST is elevated 1/2 mm. due to “early repolarization.”
T waves: generally upright.
Comment: This is an essentially normal tracing. This is the author’s own EKG,
however he is no longer 29 years old.

* Early repolarization is characterized by (minimal) ST elevation in the left chest leads, often with
rightward rotation (horizontal plane). It is a normal finding in young athletic males.
This is the followup EKG on patient D.D., 30 years after his last EKG (see previous page); he has had labile hypertension for the last 25 years. This EKG clearly demonstrates the value of obtaining the patient’s prior EKG’s for comparison.
**EKG Interpretation**

Patient:  
D.D.

Rate:  
about 58/minute

Rhythm:  
Sinus Bradycardia  
*PR* less than .2 sec. (No AV Block)  
*QRS* greater than .12 sec. reveals Bundle Branch Block.  
Leads V₁ and V₂ show R, R' complexes typical of Right Bundle Branch Block.

Axis:  
Left Axis Deviation of about -25° (in lead I, R wave greater than S wave) and Q, S, indicate probable Anterior Hemiblock.  
Axis rotation in the horizontal plane difficult to assess due to RBBB.

Hypertrophy:  
Left atrial enlargement.  
Left ventricular hypertrophy verified by other tests  
(difficult to assess on EKG with RBBB present).

Infarction:  
No significant *Q* waves.  
*ST segments*: depressed in V₁ and V₂ as related to RBBB  
*T waves*: generally upright; some T inversion in inferior leads, possibly distorted by RBBB.

Comment:  
Compared to his previous, normal EKG (see page 348), this patient has developed significant changes. There is a sinus bradycardia. The new Right Bundle Branch Block (previously incomplete) and Anterior Hemiblock have occurred in the absence of infarction. After 25 years of poorly compensated hypertension, the patient has developed left ventricular hypertrophy. The associated left atrial enlargement may stretch irritable atrial foci in the ostia of the pulmonary veins, initiating atrial fibrillation. A few days later, the patient's rhythm strip of lead I (below), shows just that. All EKG's are, unfortunately, authentic.
Patient R.C. is a 45 year old black male with a history of coronary vascular disease. Blood pressure was 210/100 on admission.
Patient: R.C.

Rate: Atrial rate of 300/minute
Ventricular rate generally 60/min. but occasionally slower.

Rhythm: Atrial Flutter (with inconsistent ventricular response, i.e.,
o no fixed AV ratio).
PR is variable.
QRS is less than .12 sec. (No BBB).

Axis: Left Axis Deviation (-30\(^\circ\)).
Leftward rotation in the horizontal plane.

Hypertrophy: Atrial hypertrophy difficult to determine.
No ventricular hypertrophy.

Infarction: \(Q\) waves: Q in lead I (also note large S in Lead III).
ST segments are generally isoelectric.
\(T\) waves are inverted in I and AVL (look closely)
and the mid-to-left chest leads.

Comment: The most obvious problem is Atrial Flutter with an atrial rate of
300/min. and a variable irregular ventricular rate (average 60/min.)
caused by the variable AV conduction ratio between 3:1 and 7:1.
An old occlusion of the Left Circumflex Coronary artery is evidenced
by the old lateral infarction. New involvement of the Anterior
Descending Coronary artery is suggested by anterior ischemia (T wave
inversion in \(V_4, V_5, V_6\), as well as by the probable Anterior Hemiblock
(shift to Left Axis Deviation with QS, configuration; previously R.A.D.
with his old lateral M.I.). Note that if one scrutinizes the T wave regions
(somewhat obscured by flutter waves) in the limb leads, the flutter
waves dip lower (suggestive of negative T waves) rather than higher
(if superimposed on upright T waves) in all but AVR, indicating a
generalized cardiac ischemia, as well as the obvious compromise of
both branches of the Left coronary Artery.
Patient K.T. is a 61 year old obese, black male who was brought into the emergency department by his family. This patient had a sudden episode of severe left chest pain. Blood pressure was 95/65.
Patient: K.T.

Rate: about 75/minute

Rhythm: Generally regular Sinus Rhythm with occasional PVC’s. 
*PR* is exactly .2 sec. so we will have to say there is a 
borderline first degree AV Block. 
*QRS* is less than .12 sec. (No BBB).

Axis: Left Axis Deviation (nearly -90°). 
No rotation in the horizontal plane.

Hypertrophy: Probable left atrial hypertrophy. 
Left ventricular hypertrophy.

Infarction: 
*Significant Q waves* in I and AVL. 
*ST segments* are elevated in I and AVL. ST segments 
are depressed in *V₁, V₂, V₃, and V₄*.
*T waves* are flat or inverted in II, III, and AVF and all chest leads.

Comment: This patient has a classical acute lateral infarction caused by an occlusion of the Left Circumflex Coronary Artery. Coincident with this is a probable occlusion of the Right Coronary Artery characterized by prominent R waves with ST depression in the (*V₁ to V₄*) chest leads. Also, T wave inversion in II, III, and AVF suggests Right Coronary compromise. T wave inversion in all chest leads is indicative of ischemia of the Anterior Descending Coronary Artery. Note also the tall, peaked T waves in I and AVL known as “hyperacute T waves,” which, although uncommon, characterize a very acute M.I. The Left Axis Deviation appeared in this patient’s previous EKG’s and is most likely related to his left ventricular hypertrophy rather than implicating Anterior Hemiblock (also, the Bundle Branch System appears to conduct normally). Occasional PVC’s caused by the ischemia, depending on frequency and multiplicity of origin, may forebode more serious arrhythmias.
Patient G.G. is a 45 year old Asian male who was doing heavy work when he was overcome by severe, crushing, anterior chest pain. Blood pressure was 110/40 on admission to the hospital.
EKG Interpretation

Patient: G.G.

Rate: about 100/minute but variable.

Rhythm: Sinus Rhythm, somewhat irregular due to Sinus Arrhythmia. 
PR less than .2 sec. (No AV Block).
QRS less than .12 sec. (No BBB).

Axis: Left Axis Deviation (-30° to -60°).  
Leftward rotation in the horizontal plane.

Hypertrophy: No atrial hypertrophy.
No ventricular hypertrophy.

Infarction: Significant Q waves in II, III, and AVF.
There are also very large Q waves in Vp, Vs, and Vq.
ST segments are elevated in Vp, Vs, and Vq.
T waves are difficult to distinguish, but inverted T waves
are noted in Vq, Vs, and Vp.

Comment: This patient has an acute antero-septal infarction, probably representing
an occlusion of the Anterior Descending branch of the Left Coronary.
Generalized ischemia of the myocardium is evident by the flat-to-
inverted T waves in nearly every lead. The old inferior infarction
demonstrated on this EKG was noted on the patient’s previous hospital
record and is the documented etiology of his Left Axis Deviation (no Hemiblock). Note that the QRS becomes isoelectric between Vp and
Vq but this is not within the normal (Vp, Vq) range; this represents
minimal leftward rotation away from the septal infarction. Old EKG’s
showed no anterior involvement on his previous admission.
Patient E.M. is a 65 year old Hispanic female. She was admitted to the hospital because of constant left chest pain for twelve hours. Blood pressure on admission was 110/75.
EKG Interpretation

Patient: E.M.

Rate: 60/minute

Rhythm: Sinus Bradycardia
PR is about .2 sec. so there is probably a first degree AV Block.
QRS is more than .12 sec. (it is .16 sec. wide). R,R' is present in V₃ and V₅, so there is a Left Bundle Branch Block.

Axis: Suggestive of Left Axis Deviation, but not reliable because of the presence of Bundle Branch Block.

Hypertrophy: No atrial hypertrophy.
Ventricular hypertrophy is difficult to determine because of Bundle Branch Block.

Infarction: Q Waves: not a reliable criterion of infarction in the presence of Left Bundle Branch Block.
ST segments: not reliable in the presence of Left Bundle Branch Block.
T Waves are flat in V₃, V₅, and V₆, but not reliable with Left Bundle Branch Block.

Comment: Enzyme studies confirmed a presumptive diagnosis of myocardial infarction. The patient’s chest pain made us suspicious.
Patient M.A. is a 75 year old black female with a long history of marked hypertension.
EKG Interpretation

Patient: M.A.
Rate: about 125/minute
Rhythm: Sinus Tachycardia
PR is less than .2 sec. (No AV Block).
QRS is less than .12 sec. (No BBB).
Axis: Left Axis Deviation (minimal amplitude of QRS in limb leads
make exact axis determination difficult).
No rotation in the horizontal plane.
Hypertrophy: Left atrial hypertrophy.
Left ventricular hypertrophy with strain.
Infarction: Q waves are present in II, III, and AVF.
ST segments: generally isoelectric (on baseline),
but V₅ and V₆ show strain pattern.
Twaves are inverted in I and AVL, and also in V₅, V₆.

Comment: This patient has hypertrophy of both the left atrium and left ventricle
with a left ventricular strain pattern. The patient also had an old inferior
infarction. The Left Axis Deviation is caused by the Mean QRS Vector
pointing away from the (old) inferior M.I. and toward the thickened
left ventricle. It does not represent Hemiblock. There is currently
(lateral) ischemia in the distribution of the Left Circumflex Coronary
Artery.
R.M., an anxious, obese, 57 year old white male whose law practice was failing, complained of “tight, squeezing” pain in his anterior chest. An electrocardiogram was quickly taken by an Emergency Medical Technician.
EKG Tracings

EKG Interpretation

Patient: R.M.

Rate: 75/minute

Rhythm: Sinus Rhythm
PR .16 sec. (No AV Block).
QRS .08 sec. (No BBB).

Axis: about +45° (Normal).
No rotation in the horizontal plane.

Hypertrophy: Possible minimal left atrial hypertrophy.
No ventricular hypertrophy.

Infarction: 
Q waves: no significant Q waves.
ST segments: elevated 2+ mm. in I and AVL.
T waves: inverted in I and AVL.

Comment: It is interesting that in this innocuous appearing EKG there is a subtle non-Q wave infarction in the lateral left ventricle, which very soon developed into a serious lateral infarction. Symptomatology suggestive of M.I. always must be investigated and scrutinized.
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