INTRODUCTION
Patients presenting to the emergency department (ED) with electrocardiograms (ECGs) indicating wide complex tachycardias (WCTs) are difficult to manage. Furthermore, these ECGs are often challenging to interpret. Patients typically have ongoing chest discomfort, with or without symptoms of dyspnea, lightheadedness, nausea, and diaphoresis. Accurate interpretation of ECGs demonstrating WCTs assists clinicians who must treat patients presenting with this condition. Despite step-wise approaches and numerous criteria suggested to interpret WCTs, physicians (including cardiologists) often fail to agree on a surface electrocardiographic diagnosis. Understanding the etiology of WCTs not only helps identify the cause of this condition, but also prevents its mismanagement, significantly reducing morbidity and mortality.

This is the first of two manuscripts designed to remove the “complex” from wide complex tachycardia identification (part 1) and management (part 2). Information is provided to help clinicians interpret ECGs demonstrating WCTs, including descriptions of the electro- and pathophysiology behind their development. Several examples of WCTs are provided with detailed interpretations. Diagnostic criteria from current literature and their relative accuracy are presented. In part 2, the most recent 2005 American Heart Association (AHA) guidelines for the treatment of WCTs are discussed, and management strategies for various WCTs are described given their underlying etiologies.

Table 1. Etiologies of a wide QRS Complex

| Etiology                                      |
|-----------------------------------------------|
| Nonspecific intraventricular conduction delay  |
| Aberrant ventricular conduction (Bundle Branch Block) |
| Ventricular ectopic beat                      |
| Ventricular excitation syndromes (Wolff-Parkinson-White Syndrome) |
| Left ventricular hypertrophy                  |
| Hyperkalemia                                  |
| Hypermagnesemia                               |
| Pacemaker-generated beat                      |
| Hypothermia                                   |
| Drug Toxicities (tricyclic antidepressants, cocaine, phenothiazines, lithium, diphenhydramine, or other drugs having sodium channel blockade or quinidine-like effects) |

* Any of these causes of a wide QRS complex may result in a WCT, given concomitant tachycardia.
Table 2. Electrophysiologic and/or metabolic causes of WCTs

| Condition                                                                                   |
|---------------------------------------------------------------------------------------------|
| Supraventricular tachycardia (SVT) with pre-existing BBB                                     |
| Supraventricular tachycardia (SVT) with rate-related BBB                                    |
| Supraventricular tachycardia (SVT) with aberrant conduction                               |
| Atrial fibrillation (AFIB) with Wolff-Parkinson-White Syndrome (WPWS)                      |
| Ventricular tachycardia (VT)                                                                |
| Polymorphic VT (Torsade de pointes)                                                        |
| Pacemaker-mediated tachycardia (PMT)                                                        |
| Drug overdose (digitalis, TCAs, lithium, cocaine, diphenhydramine)                         |
| Sodium channel blocking agents                                                             |
| Hyperkalemia                                                                                |
| Post-resuscitation                                                                         |
| Malingering and/or ECG artifact                                                            |

Before reviewing diagnostic criteria recommended for the interpretation of ECGs demonstrating WCTs, clinicians must first recognize the importance of an individual’s presenting symptoms associated with this WCT. The initial assessment of the airway, breathing, and circulation (ABCs) is critical, as these determine the approach taken in treating the patient. However, these may or may not assist the clinician’s ability to identify the underlying condition responsible for the WCT. Oxygen and intravenous access should be started as soon as possible. Cardiac rhythm and blood pressure monitoring should be initiated immediately. A focused history and physical examination must be performed during this initial period, not only to gather additional information about this individual, but also to allow consideration for alternative diagnostic possibilities.

There are several aspects of the medical history that contribute to a clinician’s ability to identify the etiology behind a WCT. Individuals with previous myocardial infarction (MI) or known coronary artery disease (CAD) are approximately four times more likely to present with ventricular rather than supraventricular etiologies of their WCT. Stated another way, approximately 80% of all patients presenting with WCT will be diagnosed having VT as the cause. This figure may be higher than that seen in patients presenting to the ED, due to referral bias at specialty centers and the repeated citation of older literature. One small retrospective study found that histories of prior myocardial infarction (MI), congestive heart failure (CHF), and recent angina pectoris all had positive predictive values for VT greater than 95%, but sensitivities were low to moderate. None of these clinical characteristics was strongly predictive for SVT; the best was age less than or equal to 35 years (positive predictive value of 70%). In fact, this study concluded that for patients presenting with WCTs, there are no ECG findings or clinical variables highly suggestive of SVT with aberration. Previous Holter or electrophysiologically-proven VT, or medications used to treat VT may be helpful historically, as they make the diagnosis of VT more likely. However, neither is conclusive for VT as the etiology of the presenting WCT. Previous electrophysiologically-proven SVT does not exclude the diagnosis of VT, nor does it confirm the diagnosis of VT, as individuals with a previous episode of SVT may present with an episode of VT. Positive responses to medications with prior episodes of “palpitations” or WCTs may not determine the etiology, as several medications work on blocking cardiac conduction both above and below the ventricle, and the etiology for the present rhythm disturbance may differ. Finally, previous adverse responses to cardiac medications may identify which medications to avoid in the management of an individual, regardless of the etiology of their presenting symptoms or ECG.

During a careful physical examination, identifying signs of atrio-ventricular (AV) dissociation strongly supports the
diagnosis of VT. AV dissociation occurs due to independent activity of the atria and ventricles, a characteristic occurrence in VT. Physical signs include irregular cannon A waves, varying intensity of the S1 heart sound, and beat-to-beat variability in systolic blood pressure (SBP). Irregular cannon A waves occur when the atria contract against AV valves that are closed because the ventricles are simultaneously contracting. These waves are the result of brisk blood backflow in the jugular veins. They do not occur with each cardiac cycle, and may be variable in quality because atrial contraction may occur when the AV valves are open or in differing stages of closure. This results in irregularity and varying quality of the cannon A wave intensity, when and if these waves occur. Varying intensity of the S1 heart sound is related to the timing of AV valve closure at the onset of ventricular systole. The beat-to-beat variability of the first heart sound’s loudness is due to the varying positions of the AV valves at the onset of ventricular systole. Beat-to-beat variability in SBP is due to the changing amount of blood in the ventricle during ventricular contraction, the result of varying diastolic filling times. On the ECG or rhythm strip tracings, the P waves and QRS complexes classically lack any relationship (i.e., there is no association between them), as demonstrated in Figure 2. This may be difficult to identify on ECGs demonstrating WCT, due in part to the tachycardia itself, as well as the morphology of the QRS complex. The absence of AV dissociation neither excludes the possibility that the WCT is ventricular in origin (VT), nor confirms that the WCT is supraventricular in origin. Vagal maneuvers, such as cautious carotid massage, gagging or coughing, and valsava can slow the ventricular response in many tachycardias of supraventricular origin, allowing atrial activity to be more apparent. These maneuvers tend to be ineffective in tachycardias of ventricular origin.

The most important point to consider when evaluating and treating individuals presenting with WCTs is their hemodynamic status. An individual’s hemodynamic status does not help determine the etiology of the WCT. A survey of physicians published in JAMA in 1985 demonstrated this fact was commonly misunderstood. The determination of whether or not a WCT is unstable is based on the effect it has on end-organ perfusion. Neither the patient’s heart rate nor QRS width determines hemodynamic stability.

Symptoms providing evidence of unstable WCTs include hypotension, pulmonary...

Figure 2: Lead II rhythm strip from a 12-lead ECG demonstrating AV dissociation in a patient with confirmed VT. Note that P waves march out independent of the QRS complexes, best seen in beats 2 – 10. Some of these P waves fall within or before the T wave, modifying its appearance. Also identified in lead II are fusion beats (QRS complexes #11, 15, and 20), and a narrow complex capture beat (next to last QRS complex #19).
edema, confusion, and angina. Skin color, temperature, and moisture may also be considered as somewhat reliable predictors of end-organ perfusion. Many individuals with WCTs (or any tachycardias) will exhibit one or more of these symptoms to varying degrees, especially chest discomfort (angina). There is likely to be overlap between one or more of these symptoms as well, making bedside clinical judgment in the management of patients presenting with WCTs essential. An approach in which clinicians err on the side of considering someone hemodynamically “unstable” is most prudent. It is incorrect to consider that only VT can be unstable, or that SVTs of any etiology do not cause hemodynamic compromise.

Many patients with known cardiac histories (especially those with rhythm or conduction problems) take antidyssrhythmic medications. These agents have various effects on cellular electrophysiology and are divided accordingly using the Vaughan-Williams international classification system. Most of these agents prolong the duration of various portions of the ECG (including the QRS complex), which may lead to the misinterpretation of WCTs. Furthermore, several of these agents are pro-arrhythmic because of their effects on conduction at the cellular level. Therefore, it is important to obtain a thorough medication history.

The age of an individual presenting with a WCT does not help determine its origin. Although WCTs in older individuals are more likely to be ventricular in origin (due to an increased incidence of previous MI or underlying CAD), the age of an individual is neither sensitive nor specific. Furthermore, an individual’s gender is not useful in determining the etiology of a WCT. Males are, however, more likely than females to have ventricular etiologies for their WCT, likely due to their increased incidence of prior structural cardiac disease and MI.

Neither the heart rate nor the width of the QRS complex in a single patient presenting with a WCT can definitively determine the origin of the rhythm disturbance. Many references state that a faster heart rate is more likely found in WCTs of supraventricular origin than in those of ventricular etiology. Ranges of heart rates that occur in SVT with aberrant conduction or VT are often given in a table, for example. However, presenting this information in this manner gives the illusion that there is a distinct heart rate above which VT does not occur. This is not true, as the range of heart rates found in SVTs and VTs overlap, depending on the underlying condition, medications, and physiological reserve of individuals experiencing these rhythms. Overall, the heart rate of WCTs due to aberrantly-conducted SVTs tend to be slightly faster than the rates due to VTs, but in a single individual at a given time, heart rate is not useful in determining the etiology of the WCT.

For patients with a permanent pacemaker, part of the physical examination diagnostics includes careful placement of a pacemaker magnet over a pacemaker generator. In patients presenting with a WCT, the magnet converts the pacemaker’s pulse generator from the synchronous (demand) to asynchronous (fixed or “magnet” rate) mode of response. This action may allow identification of atrial activity. If this WCT is pacemaker-mediated, the magnet is likely to terminate the abnormal rhythm.21

In addition to historical and physical examination clues, several diagnostic criteria exist using the rhythm strip and 12-lead ECG to help determine the etiology of a WCT (see Tables 3 - 7). In terms of the electrophysiology of the abnormal rhythm, a very regular rhythm tends to favor the diagnosis of SVT with aberrant conduction as the cause of the WCT. However, VT is generally considered as resulting in a regular wide complex rhythm. A markedly irregular rhythm favors atrial fibrillation (AFIB) with aberrant conduction, such as a preexisting Bundle Branch Block (BBB) or Wolff-Parkinson-White Syndrome (WPWS). Previous ECGs may provide essential information, such as prior rhythm patterns (normal sinus, AFIB, SVT, or VT) or previous QRS complex morphologies (BBB patterns, PVCs, fusion or capture beats during prior WCTs, electrical axis, evidence of WPWS, or structural heart disease). AV dissociation, although not commonly identified, and/or ventriculoatral (VA) block, apparent in approximately 1% of ECGs demonstrating WCT, are very sensitive for VT. Fusion and capture beats are helpful in diagnosing VT, but are neither always present nor always reliable. Fusion beats are hybrid QRS complexes resulting from simultaneous supranodal (atrial) and infranodal (ventricular) activation of ventricular tissue. They are therefore intermediate in morphology and width from either a capture or ventricular beat. Capture beats are QRS complexes resulting in ventricular activation originating from supranodal tissue, using electrical conduction pathways above the ventricle. These are therefore narrow and are similar (or identical) to a “normal” QRS complex (see Figure 2). Precordial concordance means that the QRS “direction” on the ECG in all the precordial leads is consistent. Positive precordial QRS concordance may occur during VT or SVT using a left posterior accessory pathway for AV conduction. Therefore, positive precordial concordance does not discriminate between VT and SVT with aberrant conduction. Negative precordial concordance is nearly always VT, because antidromic circus movement tachycardia never has negative precordial concordance. According to Wellens, this is true since accessory pathways over which anterograde conduction leads to completely negative QRS complexes in all the precordial leads do not exist.23 However, a recent report of a 17-year-old male demonstrated that this “rule” is not perfect, as his underlying pectus excavatum and SVT with left BBB resulted in a WCT with negative precordial concordance due to SVT with aberrant conduction. This observation led the authors to conclude that “… no diagnostic technique is 100% correct and that there are always exceptions to the rule.”23,24
Similar to the uncertainty of the history and physical examination in establishing the etiology of WCTs, ECG findings may create doubt as well. The duration (width) of the QRS complex is not very helpful in differentiating between supraventricular and ventricular etiologies of WCTs. A common misconception is that a QRS duration exceeding 0.14 second is present only if the rhythm is ventricular in origin (VT). However, in left BBB (V1-negative) WCTs, the QRS complex duration is often greater than 0.14 second even though the rhythm originates above the ventricles (i.e., supraventricular in origin). Even further misleading is that many antidysrhythmic agents cause QRS widening beyond 0.14 second, even if the rhythm originates above ventricular tissue.

Transesophageal atrial “pill” electrodes are used by electrophysiologists in an attempt to determine the etiology of underlying wide complex conduction abnormalities. These pills are “easily” swallowed by the patient, are secured adjacent to the atria, and may prove diagnostic. These electrodes record atrial activity (contraction) as large P wave spikes, which allows for easier identification of P waves and their relationship to QRS complexes. This may help identify AV dissociation or 1:1 VA conduction. A combination Esophageal Pill Electrode and Pacing Electrode may have a role in terminating certain tachydysrhythmias, such as atrial flutter, by overdrive pacing. The pill electrode has an unlikely role in the emergency department identification and management of WCTs due to its cost, availability, ease of use, safety, and lack of emergency physician experience using them. These pills are not indicated for patients experiencing hemodynamic compromise. The role of transesophageal electrocardiography in the diagnosis of WCT has received much attention (although little in the EM literature) as a safe, easy method for evaluating a WCTs’ mechanism.

One case report describes the ease of bedside placement of a transesophageal lead in the ED in a patient with an uncertain WCT unresponsive to a number of medications. A diagnosis of VT was confirmed using this recording device, which demonstrated marked disparity between atrial and ventricular rates, with the ventricular rate exceeding that of the atrial rate.

Research is ongoing using signal-averaging ECG technology, which may gain future acceptance assisting with the electrophysiological diagnosis of WCTs. There are reports of transcardiac lead systems, modified precordial lead use, and ice mapping of the AV node (reversible cooling of this tissue to test function prior to ablation) for identification of the etiologies of WCTs in electrophysiology laboratories. Presently, these technologies have limited roles in the ED.

**Bedside ECG Diagnostic Criteria**

Not all individuals presenting in a WCT are hemodynamically unstable. This is for the clinician to determine at the bedside. Once an individual has been deemed clinically stable, the ECG and rhythm strip should be reviewed closely for diagnostic clues to the etiology of the WCT. The “final” determination of the WCT should be considered an “academic exercise” and not of the utmost importance even for a stable patient in the ED. It is perfectly acceptable for an emergency physician to provide a final interpretation of the abnormal ECG demonstrating a WCT as “wide complex tachycardia of uncertain (undetermined) etiology.” In fact, this may result in better patient care, as the focus of a patient presenting in a WCT should be on the patient’s hemodynamic status and management, not the “correct” interpretation of the ECG.

A WCT can occur in any individual presenting with tachycardia for any reason, such as fever or dehydration, if he or she has a pre-existing wide QRS complex. Other tachycardias that create or result in wide QRS complexes will by definition be WCTs. The best approach when considering the etiology of a WCT is to determine whether or not the rhythm and its associated tachycardia originate from above the ventricle (supraventricular), the ventricle itself, or is due to metabolic causes.

Several diagnostic criteria exist to assist clinician determination of the WCTs’ etiology. Much research has been published debating or, in some cases, modifying these criteria, in an effort to prove or discredit them as being the most accurate, reproducible, or easiest to use. Some researchers claim that the QRS axis helps determine the etiology of a WCT. ECG leads placed on the body create Einthoven’s triangle, which can be used to determine the overall electrical axis of a patient. The quadrant in which the QRS axis is found may provide additional support that the WCT pattern is due to VT (see Table 3).

**Table 3. Electrocardiographic axis and WCTs**

| Criteria | Interpretation |
|----------|----------------|
| 1. Abnormal QRS axis supports VT (particularly if newly abnormal) |
| 2. Northwest axis (-90 degrees to +/- 180 degrees) strongly suggests VT |
| 3. In V1-negative WCT, RAD strongly supports VT |

The three most well-accepted criteria for the interpretation of WCTs based on the 12-lead ECG are the Wellens’ criteria (Table 4), Kindwall’s 4 electrophysiologic criteria for VT in LBBB (Table 5), and Brugada’s 4-step approach to regular tachycardias with wide QRS complexes (Table 6). Also included is a table using morphologic criteria of the ECG and its associated bundle branch pattern (positive and negative deflections) of the QRS (Table 7). The classification schemes based on QRS morphology are essentially the same; both are included because some individuals prefer classifying WCT ECGs as RBBB-like or LBBB-like, while others prefer to
Table 4. Wellens' Criteria (VT favored in the presence of)\textsuperscript{38}

1. AV Dissociation
2. Left Axis Deviation
3. Capture or Fusion Beats
4. QRS generally greater than 140 msec
5. Precordial QRS concordance
6. RSR' in V1, mono- or biphasic QRS in V1, or monophasic QS in V6

Table 5. Kindwall's ECG criteria for VT in LBBB\textsuperscript{39}

1. R wave in V1 or V2 of >30 ms duration
2. Any Q wave in V6
3. Duration of >60 ms from the onset of the QRS to the nadir of the S wave in V1 or V2
4. Notching on the downstroke of the S wave in V1 or V2

Table 6. Brugada's 4-step Algorithm Approach\textsuperscript{40,41}

This step-wise approach is performed as a series of questions. If the answer to any of these questions is “YES,” VT is identified and no further steps are made. If the criteria are not met for that step, the next question is asked.

1. If RS complex absent from all precordial leads, then VT
2. If RS present, and the longest precordial RS interval > 100 msec in one or more precordial lead(s), then VT
3. If atrioventricular dissociation present, then VT*
4. If morphological criteria for VT present both in precordial leads V1-2 and V6, then VT. If morphologic criteria for VT not present, then the diagnosis of SVT with aberrant conduction is made by exclusion.

*Step (Question) 3 was modified in a paper published three years later by many of the same authors at the same research facility to “more QRS complexes than P waves” (if yes, then VT).

consider WCT ECGs according to V1(V2)-positive or negative classification schemes. The Brugada criteria are perhaps slightly favored by clinicians and/or cardiologists, although this may be because they are the most recent. All of these papers have generated further research and commentaries suggesting modification of these criteria, supported by examples demonstrating inappropriately classified WCTs when using these criteria. None of these approaches is favored, as each diagnostic set has inherent limitations resulting in ECG misidentifications.\textsuperscript{36,37} It is therefore prudent to be comfortable with more than one of these classification schemes, although many feel it is better to be very confident with just one approach.

Table 7. Morphological criteria favoring VT

A. RBBB-like QRS:
   - monophasic R, QR, or RS in V1
   - R/S ratio less than 1.0, QS or QR in V6
   * triphasic QRS in V1 or V6 supports SVT with aberrant conduction

B. LBBB-like QRS:
   - R > 30 msec, >60 msec to nadir S, or notched S in V1 or V2
   - QR or QS in V6
   *monophasic R in V6 not helpful

C. Using V1(V2)-positive and V1-negative QRS morphology characteristics:
   1. V1(V2)-positive:
      - V1: mono- or biphasic QRS = VT
      - Rabbit ear sign with first peak > second (L > R) = VT
      - rSR' (triphasic) = SVT + RBBB
      - V6: QS or deep S (R/S ratio < 1.0) = VT
      - qRS (triphasic) with R/S ratio > 1.0 = SVT + RBBB
   2. V1-negative:
      - V1, V2: broad r > 0.04 sec and/or slurred or notched S resulting in prolonged interval from beginning QRS to S nadir = VT
      *narrow r wave and quick S wave downstroke = SVT + LBBB
      - V6: any q wave = VT

An interesting commentary published in 1989 following an article describing yet another set of criteria for identifying WCTs stated that “… knowing the specific criteria for diagnosis may not be that important.” The academic emergency physician who wrote this recommended instead that appropriate therapy should be dictated by a three-step protocol (Table 8).\textsuperscript{42} This simplified three-step protocol focuses on clinical criteria rather than electrophysiologic ones, which more closely resembles 2005 AHA guidelines. When in doubt, VT should be considered the default diagnosis in WCTs.\textsuperscript{43}

Table 8. Three-step clinical protocol to WCTs\textsuperscript{42}

1. If the patient is asymptomatic, or minimally symptomatic and hemodynamically stable, … call an experienced electrocardiographer or look up the criteria … while observing the patient….
2. If the patient is hemodynamically unstable, immediate synchronized graded cardioversion … is indicated. Subsequent pharmacologic therapy can be guided by experienced physicians.
3. If the patient is symptomatic … but is otherwise hemodynamically stable, then controlled graded cardioversion or pharmacologic therapy [sic] … may be tried.
UNDERSTANDING THE ECG IN WCTs

Understanding the electrophysiology behind ECGs with WCT assists clinicians with decision analysis and the explanation for abnormal conduction. Situations often exist during which a “normal” pathway of cardiac conduction does not occur. When this happens, the QRS complexes may appear different, having an increased duration or width. One such situation occurs during pre-excitation of the ventricle. Here, the normal electrical impulse originating from the atria reaches ventricular tissue earlier than expected by means of an accessory pathway. This impulse bypasses the AV node, which eliminates AV nodal delay and allows for earlier activation of the ventricle. The three most common accessory pathways identified are the Kent bundle (AV bypass tracts), responsible for the Wolff-Parkinson-White Syndrome (WPWS), the James’ fibers (atrio-His fibers), responsible for the syndrome known as Lown-Ganong-Levine, and the Mahaim bundles (nodoventricular fibers). These accessory pathways each result in ventricular tissue receiving activation from a supraventricular source earlier than expected. This premature ventricular activation results in a shortened PR interval (the time from the beginning of atrial depolarization to the onset of ventricular depolarization). WPWS, which uses the Kent accessory bundle, is important because the QRS complex in individuals with this accessory pathway is widened to more than 0.10 seconds when this pathway is used.\(^4\)\(^4\)\(^5\) This QRS complex widening occurs due to premature activation of the ventricle, not because of delayed activation (which occurs in BBB). WPWS has a slurred initial upstroke of the QRS known as a delta wave, the result of partial premature activation of ventricular tissue from the Kent accessory bundle combined with the majority of ventricular activation via the normal conduction pathway.\(^4\)\(^6\) This delta wave is therefore analogous to a fusion beat.

Reentry may be defined as an alternative pathway for cardiac conduction, with pre-excitation being one of its manifestations. These alternative pathways generally result in faster activation of ventricular tissue. They have varying periods of refractoriness, which is important when considering reentrant tachycardias. The Kent bundle, for example, is composed of faster conducting tissue than the AV node, but it has a longer refractory period. AV Nodal Reentry Tachycardias (AVNRTs) are tachycardias that use two distinct pathways within the AV node, having different refractory periods and conduction velocities. Circus Movement Tachycardias (CMTs), on the other hand, are reciprocating tachycardias that demonstrate either narrow or wide QRS complexes depending on the direction of reentry. An orthodromic reentry tachycardia means that AV conduction occurs in the normal direction (over the normal pathway). In antidromic CMTs, impulse conduction over the “expected” pathway occurs in the reverse direction. Here, an accessory pathway is used for anterograde (forward) conduction and the AV node or other accessory pathway is used in the retrograde (backward) direction as part of the conduction loop (see Figure 1). This results in wide QRS complexes, and therefore a WCT if the heart rate is greater than 100 beats per minute. Anterograde and retrograde are descriptive terms of convention; by themselves, they do not provide information about the configuration of the QRS complex nor the etiology of the WCT.

WPWS is the most common form of pre-excitation, using the Kent bundle as the accessory pathway for conduction. Left free wall pathways account for the majority of locations of the Kent bundle, although other locations for this accessory pathway exist in WPWS. Common ECG features of WPWS include short PR intervals (< 0.12 seconds), delta waves, and prolonged QRS intervals (> 0.10 seconds). Inverted T waves and increased precordial voltage may also be seen. Three types of WPWS are determined based on the location of the accessory pathway; however, they have little clinical significance other than identifying this syndrome. Type A WPWS has its accessory pathway in the inferoposterior portion of the LV, which results in a dominant R wave in lead V1 and Q wave in the inferior leads (the ECG configuration is therefore RBBB-like). Type B WPWS has its accessory pathway in the inferoposterior region of the RV. This results in an rS or QS pattern in lead V1 (the ECG configuration is therefore LBBB-like). Type C WPWS has its bundle in the posterolateral region of the LV, which results in negative or isoelectric delta waves in leads V5 or V6. The ECG exhibits small Q waves in leads I and aVL, commonly referred to as the pseudoinfarct pattern of WPWS.

WPWS is important because greater than 50% of individuals with these accessory pathways for AV conduction are predisposed to supraventricular tachydysrhythmias via reentry. Atrial fibrillation occurring in individuals with WPWS may be extremely dangerous, as the Kent bundle may allow fast, chaotic atrial impulses to pass to ventricular tissue without delay at the AV node. Ventricular rates (identified as R-R intervals) may reach as high as 300 beats/minute in this situation. A clue to this is the presence of rapid, wide, and irregularly-spaced QRS complexes on the ECG.\(^4\)\(^7\)\(^8\)\(^9\) In this situation, clinicians must not administer agents that slow conduction through the AV node, as this makes conduction via the accessory pathway more likely. Therefore, antidysrhythmic agents that block AV nodal conduction or preferentially increase conduction through an accessory pathway in the situation of a WCT due to AFIB and WPWS can result in immediate hemodynamic collapse, ventricular fibrillation, or death.
Figure 3. ECG demonstrating AV Dissociation (see lead II rhythm strip, in which P waves march independently of wide QRS complexes). This RBBB-like patterned ECG with delayed ("notched") upsloping of the S wave was subsequently confirmed to be VT (image courtesy Amal Mattu, MD). Used with permission, ACEP.

Figure 4. ECG demonstrating WCT (rate just under 200 bpm). It appears quite regular. The QRS duration is 155 msec, and its morphology is V1-positive (RBBB-like). Although positive precordial concordance is present, this WCT is due to SVT with aberrant conduction. An AV nodal reentry mechanism was confirmed during subsequent EP testing.

Figure 5. WCT with a rate of 140 bpm. The QRS duration is just less than 200 msec, with a V1-negative (LBBB) QRS morphology. There is a very narrow R wave (less than 40 msec) in V1 and V2, with a quick, near-vertical downward deflection to the S wave nadir. The RS duration is less than 80 msec. This ECG demonstrates SVT with LBBB conduction, a pattern present during old ECG tracings. This tracing does not meet any of Brugada’s 4-step algorithm for VT.

Figure 6. WCT due to SVT with a preexisting BBB in an 81 year-old male. This patient has known structural heart disease, including prior LBBB and 1st degree AV block, which makes the diagnosis of VT more likely. Previous ECGs demonstrated a LBBB pattern with identical QRS morphology. Note the PVC has different QRS morphology from the other beats, best seen in the lead II tracing. Due to its originating location, it appears similar to the QRS complexes in leads V1 and V2. However, this differing appearance of this PVC does not provide conclusive evidence that the WCT is being generated above the ventricle.
Figure 7. 12-lead ECG demonstrates classic VT.

Figure 8. Another example of VT, although this ECG’s axis differs from the previous example. Previous ECGs in this older patient with a prior MI demonstrate PVCs identical in morphology to these QRS complexes. Lead V5 has baseline wander due to a poorly adherent lead.

Figure 9a. Episodic VT following an exercise treadmill test in a 44 year-old male with atypical chest pain. It is unusual to capture both VT and sinus activity on a 12-lead ECG. AV dissociation is noted in first part of lead II and the lead II rhythm tracing. A post-conversion ECG (next image) demonstrates normal sinus rhythm and PVCs, without significant ischemic changes (image courtesy of Eleanor Levin, MD).

Figure 9b. Same patient as previous ECG, following spontaneous conversion of exercise-induced VT. Several PVCs are present throughout the tracing; the PVC in lead V2 is nearly identical to that seen in lead V2 in the previous tracing.
Figure 10a. ECG incorrectly interpreted by the computer as VT. This is a regular WCT with a RBBB/LPFB configuration. In this 88 year-old patient, the correct clinical interpretation, however, is VT. In most tracings with a rapid heart rate, it becomes difficult to appreciate irregularity in the rhythm, or it may disappear completely. A subsequent ECG tracing (next image) demonstrates AFIB with the identical QRS morphology, including RBBB and LPFB.

Figure 10b. Subsequent ECG from the previous patient demonstrates AFIB with an irregularly irregular wide complex QRS having RBBB and LPFB morphologies. An isolated PVC (beat #12) has a different morphological quality than the QRS complex. This does not exclude VT, although VT is more likely if PVCs are identical to the QRS morphology in a previous (or subsequent) WCT.

Figure 11a. 19 year-old patient with a WCT ECG, incorrectly interpreted by the computer as VT. On close inspection, the R-R intervals are irregularly spaced, several approaching 300 msec (best seen in beats 3-4 in lead I and beats 3-5 in lead V2, V3). When the R-R interval is that narrow, atrial fibrillation in a patient with an accessory pathway (bypass tract) must be considered. The QRS morphology in lead V6 gives the impression of a delta wave (slurred upstroke). The diagnosis is an irregularly irregular WCT, which is due to AFIB and WPWS (image courtesy of Charlie Young, MD).

Figure 11b. The same 19 year-old patient with an irregular WCT that returned more than one hour later. In this tracing, it is somewhat easier to see the irregular irregularity of the QRS complexes, and to identify the delta waves in lead V6 that have a slightly different appearance from the previous tracing. The computer correctly identified this WCT.
Figure 11c: Previous patient post-conversion in normal sinus rhythm, demonstrating WPWS (Type B). There are negative QRS complexes over the right anterior precordium (leads V1-V3), resembling a LBBB pattern, and some associated ST-T wave abnormalities. A right posterolateral AV accessory pathway was identified on EP studies, mapped near the tricuspid annulus.

Figure 12: ECG representing Wolff-Parkinson-White syndrome (WPWS), Type A. There is a dominant R wave in V1 (RBBB pattern). Note the classic delta waves (slurred upstroke of the QRS complex) and the short PR intervals. The delta wave causes a slightly widened QRS morphology. The cause of this patient’s pre-excitation pattern was determined to be due to a left-sided accessory AV pathway during EP studies.

SUMMARY

WCTs are particularly challenging for clinicians to correctly identify. This is especially true given that most patients presenting to an emergency department with ECGs demonstrating WCTs are uncomfortable or have some hemodynamic distress. Correct interpretation of the WCT should not be the primary concern of emergency physicians; treating patients without doing harm takes precedence. Appropriate treatment approaches, including a detailed discussion of pharmacologic therapies, as well as additional (miscellaneous) causes of WCTs will be described in a forthcoming article.

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