Patients present to the emergency department with either an ongoing tachycardia or a history suspicious of a tachycardia. Either way, the tachycardia needs to be documented, preferably on a 12-lead electrocardiogram (ECG) for diagnosis and management. If a tachycardia is not documented, a careful history of the palpitations should be taken to see if further monitoring and investigations are required. If a tachycardia is confirmed on an ECG, the clinician needs to classify it according to two variables: (i) regularity of the rhythm; and (ii) QRS width. This will allow a differential diagnosis to be made.

A QRS is narrow because ventricular activation occurs via the normal His-Purkinje system. SVT includes a wide range of arrhythmias that originate above the bifurcation of the bundle of His, such as sinus tachycardia, atrial flutter (AF), atrial tachycardia (AT), atrioventricular junctional re-entrant tachycardia (AVJRT) and junctional ectopic tachycardia (JET). AVJRT can be due to atrioventricular nodal re-entrant tachycardia (AVNRT), where the antegrade limb is the slow pathway of the atrioventricular (AV) node and the retrograde limb is the fast pathway, or atrioventricular re-entrant tachycardia (AVRT), where the antegrade limb is the AV node and the retrograde limb is the accessory pathway (orthodromic AVRT). A WCT has a QRS duration ≥120 ms and may be caused by one of the following four mechanisms:

- VT: VT must be the default diagnosis in any patient with a WCT and accounts for 80% of all cases of WCT.[1] The presence of structural heart disease and a previous myocardial infarction (MI) increases the likelihood that the WCT is VT to >90%.[2] VT usually occurs on the basis of a scar re-entrant mechanism (commonly due to a prior MI, cardiac surgery or cardiomyopathy). Some idiopathic VTs may occur in a structurally normal heart and be due to triggered activity (e.g. right ventricular outflow tract VT).
- SVT with a right or left BBB or an intraventricular conduction abnormality. This

### Mechanisms of tachycardia

An SVT usually has a QRS duration of <120 ms, unless there is bundle branch block (BBB) or an intraventricular conduction abnormality. The
accounts for 15 – 20% of WCTs. The BBB conducts antegradely over an accessory pathway (Wolff-Parkinson-White syndrome) will produce a WCT. In ‘antidromic’ AVRT, the antegrade limb is the accessory pathway and the retrograde limb the AV node.

- Pacemaker tachycardia. Usually with RV apical pacing the morphology of the paced QRS complexes will have an atypical left bundle branch block (LBBB) pattern in V1 - V3, with left axis deviation. This may be seen in dual-chamber pacemakers (DDD) when there is tracking of the atrial rhythm (e.g. sinus tachycardia, atrial tachycardia). Pacemaker tachycardia may also occur in single-chamber pacemakers (VVIR) when the lower rate is increased by the sensor in the pacemaker.

**Diagnostic evaluation**

The evaluation of a patient with a tachycardia should begin with a detailed medical history (including medication history) and physical examination. The clinical context often helps in the interpretation of the 12-lead ECG.

**History**

In a patient in whom a tachycardia is suspected but not confirmed, one should focus on the history of palpitations. In the emergency department, palpitations are most commonly due to sinus tachycardia. The following characteristics are important when taking a history, which may raise the suspicion of an underlying arrhythmia:

- Rate and regularity. The clinician should ask the patient to tap out the rhythm with his/her fingers. In this way, the clinician can determine the nature of the palpitations, e.g. rapid regular, rapid irregular, slow regular, slow irregular, ectopic beats, missed beats or strong beats. Rapid regular beats are suggestive of an SVT or a VT. Rapid irregular beats are suggestive of AF or AFL with variable AV block. Ectopic beats or missed beats suggest premature ventricular complexes (PVCs) or premature atrial complexes (PACs).

- Mode of onset. Palpitations with an abrupt onset are suggestive of an SVT or a VT. A gradual onset or ‘warm-up’ is suggestive of sinus tachycardia.

- Associated heart disease. Underlying structural heart disease is an important substrate for the development of arrhythmias, e.g. a history of ischaemic heart disease or prior MI strongly suggests VT. Heart disease raises the suspicion of AF, AFL or VT. Lung disease may lead to atrial dilatation and AF, AFL or multifocal atrial tachycardia (MAT). A family history of sudden cardiac death or syncope should be sought when a familial cause is suspected (tordse de pointes with the congenital long QT syndrome and VT or PVCs with arrhythmogenic right ventricular cardiomyopathy (ARVC)).

If a tachycardia is suspected, but has not been previously documented, the clinician has the option to arrange 24-hour Holter monitoring or inpatient telemetry, depending on the frequency of palpitations. In clinical practice, performing an ECG during an episode of future palpitations is likely to have the best yield. An implantable loop recorder can be considered if VT is strongly suspected.

**Physical examination**

The physical examination is useful to identify underlying cardiac disease that may serve as a substrate for arrhythmias. Signs of an underlying cardiomyopathy and heart failure raise the possibility that the palpitations are due to VT or AF/AFL. SVT can occur in patients with normal hearts and in those with underlying heart disease. Sometimes the SVT can cause left ventricular dysfunction, i.e. tachycardia-induced cardiomyopathy. The physical examination during a tachycardia is most helpful if signs of AV dissociation are present, which strongly suggests VT. Intermittent prominent cannon ‘A’ waves in the JVP waveform reflecting simultaneous atrial and ventricular contraction confirm AV dissociation.

**Twelve-lead electrocardiography in sinus rhythm**

The physician should look for clues to the cause of tachycardia by examining the 12-lead ECG in sinus rhythm. Prior ECGs should be sought and QRS morphology should be examined looking for BBB patterns and intraventricular conduction abnormalities.
The tachycardia and sinus rhythm QRS complexes may then be compared. Some ECG abnormalities point towards structural heart disease, which may provide clues as to the most likely cause for an arrhythmia:
- Left atrial enlargement and/or PACs suggest the diagnosis may be AF or AFL.
- The presence of Q waves suggests prior MI and possible VT or AF/ AFL.
- Marked left ventricular hypertrophy with deep septal Q waves in I, aVL and V4 - V6 suggests hypertrophic cardiomyopathy.

Other ECG abnormalities point towards a primary electrical cause for the arrhythmia:
- A short PR interval, delta wave and wide QRS complex are evidence of pre-excitation (Wolff-Parkinson-White syndrome) and suggest a diagnosis of AVRT or pre-excited AF.
- Frequent PVCs with LBBB morphology and an inferior axis suggest right ventricular outflow tract VT as a cause, which may be idiopathic or due to ARVC. T-wave inversion in leads V1 - V3 and the presence of an Epsilon wave are further diagnostic signs of ARVC.
- A long QT interval, Brugada pattern or early repolarisation pattern suggests polymorphic VT (PMVT) or torsade de points.

Twelve-lead electrocardiography of the tachycardia
The initial analysis of the tachycardia is best followed by a methodical approach to ECG interpretation. A high-quality 12-lead ECG is more useful in the clinical setting than individual rhythm strips.

The three main observations to consider in the ECG interpretation of a tachycardia are the following:
- Confirm the tachycardia (i.e. ventricular rate ≥100 bpm). The easiest way to calculate the ventricular rate is to count the number of QRS complexes and multiply by 6 (most ECGs are recorded at 25 mm/s = 10-second recording).
- Are the QRS complexes wide (≥120 ms) or narrow (<120 ms)? Care must be taken to examine all leads, looking for the widest QRS complex. Some WCTs may have narrow QRS complexes in some leads.
- Are the QRS complexes regular or irregular?

The clinician is now able to classify the tachycardia into four groups, based on these three simple observations (Fig. 1):
- Regular, narrow complex tachycardias include sinus tachycardia, AT, AFL, AVJRT (AVNRT/AVRT) and JET.
- Irregular, narrow complex tachycardias include AF (most likely), AFL with variable block or MAT.
- Regular, wide complex tachycardias include monomorphic VT (in 80% of cases), SVT with right BBB (RBBB) or LBBB, pre-excited tachycardia over an accessory pathway, and pacemaker tachycardias.
- Irregular, wide complex tachycardias include PMVT, AF or AFL with RBBB or LBBB and pre-excited AF.

In patients with SVT, the next step would be to look for underlying P waves, flutter or fibrillatory waves. There may be a P wave hidden in the T wave or QRS complex:
- P-wave morphology: P waves are usually best seen in the inferior leads (II, III and aVF) and V1. The P-wave axis should be calculated (normal: 30 - 70°). P waves should be positive in the inferior leads because right atrial activation starts at the sinus node and propagates to the AV node in an inferior direction. By contrast, AVJRT (AVNRT or AVRT) or AT activates the atria in a superior direction, with negative P waves in the inferior leads (Fig. 4). These P waves tend to be narrow (80 ms) because both atria are depolarised simultaneously from the lower septum.
- Relative position of the P wave in the R-R interval. A common nomenclature used is to classify SVT as long RP (latter half)

![Fig. 3. Typical patterns of right and left BBB.](image)

![Fig. 4. An example of a narrow complex (supraventricular) tachycardia. P waves are visible at the end of the QRS complexes. In V1, the P wave is seen as a pseudo R wave in V1 (arrow). In II, III and AVF the P waves are negative and are seen as pseudo S waves in AVF (arrow). In AVNRT the antegrade limb is the slow pathway and the retrograde limb the fast pathway of the AV node.](image)
or short RP tachycardias (first half), based on the position of the P wave in the R-R interval. This is purely descriptive and does not help the clinician to narrow the differential diagnosis. Examples of a short RP tachycardia include: typical AVNRT, AVRT or AT. Examples of a long RP tachycardia include: atypical AVNRT, AVRT over a slowly conducting accessory pathway (also called permanent junctional reciprocating tachycardia) and AT. JET and AT can present as both types of examples. AVNRT (90%) and AVRT (87%) are usually short RP tachycardias, but ATs are usually long (11%).

- Onset and termination. If the SVT starts with a very long PR interval at the onset of the tachycardia, the clinician should diagnose AVNRT with antegrade slow pathway conduction accounting for the prolonged PR interval. An SVT that starts with a PVC is usually an AVJRT, and AT is improbable. SVT that terminates with a P wave is unlikely to be an AT, as the last atrial beat will probably not cause AV block with the same beat. It is more likely an AVJRT, where block in the AV node caused the tachycardia to terminate. A vagal manoeuvre increases vagal tone and may terminate an AV nodal-dependent tachycardia, but never an AT. For AFL and AT, transient AV block may reveal underlying AVJRT as well as some types of AT (making it less specific than carotid sinus massage). Intravenous verapamil should be avoided in the emergency setting.

The long-term management of SVT is medical or catheter ablation. Patients with recurrent symptomatic AVJRT can be treated with AV nodal blockers (such as beta-blockers or verapamil). Catheter ablation should be offered to patients with AVJRT as the success rate is high (95%), with 5% requiring a second procedure. The decision to pursue a rate versus rhythm control strategy for AF/AFL should be individualised depending on symptoms, rate of tachycardia and whether there is underlying LV dysfunction.

**Management of tachycardia**

**Acute management of SVT**

The golden rule of ABC (airway, breathing, circulation) should be followed in the emergency management of SVT. If the patient is haemodynamically compromised, urgent direct current (DC) cardioversion should be performed. Most patients who present with an SVT are haemodynamically stable, allowing the physician to obtain a thorough history and do a physical examination.

The initial strategy should be to terminate an SVT with a vagal manoeuvre, such as carotid sinus massage, after excluding any carotid artery bruit. This should be performed with the patient in the supine position with a continuous 12-lead ECG. A vagal manoeuvre increases vagal tone and may terminate an AVJRT (AVNRT/AVRT), but never an AT. For AFL and AT, transient AV block may reveal underlying P waves or flutter waves. If these manoeuvres do not terminate the SVT, the next step is intravenous adenosine. Starting doses of 6 mg with doses of 12 mg and 18 mg are normally used. Adenosine has a short half-life (<10 seconds) and produces transient heart block. Adenosine should be avoided in patients with an obvious AFL, as the reflex sympathetic activation may precipitate 1:1 AFL with haemodynamic decompensation. Adenosine will usually terminate AVJRT as well as some types of AT (making it less specific than carotid sinus massage). Intravenous verapamil should be avoided in the emergency setting.

The golden rule of ABC should be followed in the emergency setting of a WCT. If the patient is haemodynamically compromised, urgent DC cardioversion should be performed. It is not uncommon for a WCT to present in an unstable way, and does not help the clinician to narrow the differential diagnosis. Examples of a short RP tachycardia include: typical AVNRT, AVRT or AT. Examples of a long RP tachycardia include: atypical AVNRT, AVRT over a slowly conducting accessory pathway (also called permanent junctional reciprocating tachycardia) and AT. JET and AT can present as both types of examples. AVNRT (90%) and AVRT (87%) are usually short RP tachycardias, but ATs are usually long (11%).

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**Acute management of WCT**

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**Acute management of WCT**

The golden rule of ABC should be followed in the emergency setting of a WCT. If the patient is haemodynamically compromised, urgent DC cardioversion should be performed. It is not uncommon for a WCT to present in an unstable way,
irrespective of the cause. VT may degenerate into pulseless electrical activity and ventricular fibrillation (VF). As described above, a wide complex tachycardia should always be considered VT until proven otherwise. Haemodynamic stability does not exclude VT.

The treatment of choice for stable and unstable monomorphic VT is cardioversion after sedation has been administered. Amiodarone IV is recommended in some guidelines as an initial strategy. Amiodarone was only 25% effective in terminating VT, with a side-effect profile of hypotension that may worsen the haemodynamic status. Therefore, cardioversion should be readily available if amiodarone is chosen as an initial treatment strategy. Beta-blockers and verapamil are contraindicated in the acute setting and can cause severe haemodynamic deterioration and even VF or cardiac arrest. Patients with VT or WCT of uncertain origin should be referred to a cardiologist for further investigation and consideration of an implantable cardioverter defibrillator (ICD).

The treatment of PMVT or torsade de points due to a long QT interval (either congenital or more often acquired) is different from that of monomorphic VT. Electrical cardioversion is the treatment of choice. As the underlying cause of torsade de points is a long QT interval, most anti-arrhythmic drugs (including amiodarone) will exacerbate this and are contraindicated. Once the patient has been stabilised, the underlying cause should be sought and treated (correct the hypokalaemia and hypomagnesaemia and discontinue all long QT-prolonging drugs). Acute ischaemia is a common cause of PMVT and urgent coronary angiography is indicated. Other causes of PMVT include channelopathies such as Brugada syndrome and catecholaminergic PMVT.

Some idiopathic outflow tract VTs may be adenosine sensitive. Adenosine should generally be avoided unless an SVT with BBB is suspected, as the reflex sympathetic activation may worsen VT. Verapamil must be avoided in all WCTs, with the rare exception of a patient with a known history of left fascicular VT proven to be verapamil sensitive in the past. Patients with idiopathic VT should be referred for ablation.

A WCT may suggest the diagnosis of pre-excited AF (Fig. 6). There is no role for medical therapy in the treatment of pre-excited AF, as beta-blockers, calcium channel blockers, adenosine and digoxin do not block and may enhance conduction over an accessory pathway, resulting in very rapid ventricular response with cardiac arrest. These patients should receive immediate electrical cardioversion and be referred for urgent ablation of the accessory pathway.

Carotid sinus massage or adenosine may terminate an antidromic AVRT. Pre-excitation should be looked for on the sinus rhythm ECG to confirm the diagnosis.

Conclusion

ECG documentation of a suspected tachycardia is crucial in the management of a patient with a suspected tachycardia. The clinician needs to decide whether the tachycardia is narrow or wide and whether the rhythm is regular or irregular. When an SVT is suspected, the clinician should look for P waves and use vagal manoeuvres (such as carotid sinus massage) before administering adenosine, to differentiate AVRT from AT and AFL. A WCT should always be considered to be VT until proven otherwise. QRS morphology is the most useful feature to distinguish between VT and SVT with BBB. Electrical cardioversion is the treatment of choice for stable and unstable VT.

References


Fig. 6. An example of a wide complex irregular tachycardia. This is pre-excited AFL over a left lateral accessory pathway. Note that the QRS complexes are wide and vary in morphology because of variable degrees of fusion over the accessory pathway and the AV node.