CASE 1

CLINICAL PRESENTATION

A 78-year-old Chinese man presented with lower limb swelling. He had a background of hypertension, diabetes mellitus, dyslipidaemia, chronic kidney disease and atrial flutter. He was on atenolol 25 mg for his hypertension and atrial flutter. The patient was admitted for worsening renal function and fluid overload, and was found to be bradycardic. The admission electrocardiogram (ECG) is shown in Fig. 1a. A temporary pacing wire (TPW) was subsequently inserted and the post-TPW insertion ECG is shown in Fig. 1b. What do the ECGs in Figs. 1a and b show?

Fig. 1 Case 1: 12-lead ECG (a) on admission and (b) after insertion of temporary pacing wire.
ECG INTERPRETATION
Fig. 1a shows atrial flutter with a regular R-R interval. There is a narrow-complex escape rhythm with a ventricular rate of only 36 beats per minute. The flutter waves are best appreciated as a sawtooth pattern of p waves in leads II, III, aVF and V1. Fig. 1b shows a paced rhythm after insertion of the TPW. There is a pacing spike followed by a wide-complex QRS with a left bundle branch block (LBBB)-like morphology.

CLINICAL COURSE
After TPW insertion, the patient was closely monitored for resolution of bradycardia while waiting for the atenolol to wash out. He underwent diuresis and received non-invasive ventilation for his fluid overload. The patient was no longer dependent on the TPW by Day 3 of admission and was transferred to the general ward for further management. He was subsequently discharged uneventfully and did not require permanent pacemaker insertion.

CASE 2
CLINICAL PRESENTATION
A 68-year-old Indian woman presented with giddiness and was admitted to rule out a posterior circulation stroke. Her significant past medical history included diabetes mellitus, hypertension, hyperlipidaemia and end-stage renal disease for which she was on regular haemodialysis. An ECG was done (Fig. 2). A TPW was inserted emergently, and ECGs done after insertion are shown in Figs. 3a–c.

ECG INTERPRETATION
Fig. 2 shows complete heart block with regular P-P and R-R intervals. After TPW insertion. Although uncommon, uncomplicated right ventricular apical pacing may show an RBBB morphology.\(1-4\) The placement of leads V1 and V2 one intercostal space lower than standard placement usually eliminates the RBBB appearance and results in inscription of deep QR or rS complexes in V1 and V2. On the other hand, placing the leads one intercostal space higher than the usual placement enhances the height of the R wave.\(5\)

CLINICAL COURSE
Chest radiography was performed following TPW insertion (Fig. 4). The patient subsequently underwent coronary angiography, which revealed triple vessel disease. No coronary angioplasty was done and the patient was advised to undergo coronary artery bypass grafting (CABG). The complete heart block resolved spontaneously by the second day of admission with the patient not requiring any support from the TPW thereafter. The initial complete heart block was attributed to underlying ischaemic heart disease. While awaiting the patient and family’s decision regarding CABG, the patient suffered a left middle cerebral artery infarction. Despite a thrombectomy, the patient had poor neurological recovery. The decision was made to postpone further invasive procedures in view of the poor neurological recovery.

DISCUSSION
A third-degree atrioventricular (AV) block or complete heart block occurs when the atria and ventricles are paced independently of each other (i.e. AV dissociation) due to loss of conduction from the atria to the ventricles. In patients with sinus rhythm, the p waves are present with a regular rate faster than the ventricular rate, while QRS complexes are present with a slow and usually fixed ventricular rate. The p waves bear no relationship to the QRS complexes and the PR intervals are completely variable, as the atria and ventricles are electrically disconnected. In patients with atrial fibrillation or atrial flutter, the ventricular rate is slow and regular, as the heart relies on infranodal backup activity such as the junctional or ventricular escape rhythm, irrespective of the
Fig. 3 Case 2: 12-lead ECG done (a) after emergent insertion of a temporary pacing wire shows a pacing spike followed by a wide-complex QRS with a right bundle branch block-like morphology. 12-lead ECG done (b) one intercostal space down and (c) one intercostal space up.
atrial rates. Such infranodal activity, especially the ventricular escape rhythm, can be unreliable.

The patient in Case 1 had atrial flutter and complete heart block. Under normal circumstances, the AV node in patients with atrial flutter is constantly inundated with flutter waves, resulting in a typical 2:1 or 4:1 conduction with a rapid ventricular rate. Variable blocks may also occur, resulting in an irregular heart rate. Therefore, whenever a patient with atrial flutter presents with very slow ventricular rate and regularisation of the R-R interval, the clinician should examine the ECG closely for the presence of underlying high-grade AV block (high-grade second-degree heart block or complete heart block). In patients with atrial flutter, complete heart block is suspected if the heart rate becomes very slow, as demonstrated in Case 1. Similarly, the presence of complete heart block should be suspected in patients with atrial fibrillation who present with regularisation of the R-R interval.

While drugs such as beta-adrenergic agonists (e.g. dopamine, epinephrine) are able to exert direct effects to enhance the junctional and infranodal escape rhythm in the setting of complete AV block, they may also result in adverse effects such as elicitation of ventricular arrhythmias and induction of coronary ischaemia, particularly in the setting of unstable coronary artery disease (as in Case 2). In addition, atropine is unlikely to improve AV block at the His bundle or His-Purkinje level. Therefore, TPW insertion was chosen as the interim measure of choice over pharmacological measures in both cases.

A thorough evaluation should be done to look for transient or reversible causes (Box 1) in all patients presenting with new AV block, for which treatment or resolution may make permanent pacing unnecessary. In the interim, this group of patients should have optimal medical and supportive care, including temporary transvenous pacing if necessary, before determination of the need for permanent pacing. In Case 1, the patient was no longer pacing-dependent after atenolol was washed out and did not require a permanent pacemaker. It is worth noting that atenolol is almost exclusively excreted in the kidneys and was previously reported to be one of the most frequently used beta blockers in patients aged above 60 years with symptomatic bradyarrhythmia. It should therefore be used with caution in elderly patients with renal impairment. In Case 2, the complete heart block was attributed to underlying ischaemic heart disease. No coronary intervention was performed and the complete heart block subsequently resolved spontaneously. No further interventions were carried out for the patient due to her poor neurological status.

Serious complications can arise from a misplaced TPW, including systemic thromboembolism, arrhythmia and perforation. In Case 2, the development of a cerebrovascular accident after TPW insertion is a major red flag and should cue the clinician to urgently determine the position of the TPW. To confirm its placement, a left anterior oblique (LAO) projection on fluoroscopy immediately after TPW insertion is useful in confirming the position of the lead. Chest radiographs (posteroanterior and lateral), echocardiograms and ECGs are other useful adjuncts in detecting pacemaker lead malposition.

Conventionally, the insertion of a transvenous TPW should yield an LBBB QRS pattern, as evidenced in Case 1. When an RBBB configuration appears after insertion of TPW, such as that observed in Case 2, it is imperative to rule out inadvertent left ventricular pacing through intracardiac defects such as a patent foramen ovale, a ventricular septal defect or complications such as ventricular perforation or malposition of the pacing lead. In addition, an RBBB pacing pattern can also be seen if the lead is placed in the middle cardiac vein or branches of the coronary sinus.

Klein et al previously reported that eight patients whose pacing leads were located in the right ventricle apex had an RBBB pattern in leads V1 and V2. The pseudo-RBBB pattern was suggested to be a result of a superior vector with depolarisation of the right ventricle preceding activation of the left ventricle, and not left-to-right septal activation. Coman et al reported seven similar cases with an RBBB pattern during permanent right ventricular pacing, with each case having pacing leads located in the distal right ventricle septum or apex. In our patient (Case 2), moving the leads one intercostal space up resulted in an increase in amplitude of the R wave at V1 and V2. However, moving the lead tips back into the right ventricle apex resulted in the typical pseudo-LBBB pattern. The pseudo-LBBB pattern was therefore interpreted as an artifact due to the placement of the pacing leads.
V1 and V2 leads one intercostal space down did not eliminate the RBBB appearance.

In both our cases, the Q waves in lead III were larger than the Q waves in lead II. Lead III is a right-sided lead, whereas lead II is left-sided. A pacing wire placed in the right ventricle would thus cause a larger Q wave in lead III than in lead II, as the wavefront travels from right to left. In patients whose TPW is placed in the middle cardiac vein, lead II would have a larger Q wave than lead III, as the wavefront travels in a left-to-right direction.

An RBBB pattern with maximal QRS vector oriented to the right, inferior and posterior may be a warning sign of perforation of the right ventricle, whereas an RBBB pattern in right ventricular pacing with maximal QRS vector oriented to the left, superior and anterior may indicate uncomplicated right ventricular pacing. Coman et al developed an algorithm to differentiate left and right ventricular RBBB pacing morphologies using frontal axis and precordial transition. When a frontal plane axis occurs between −90° and −180°, it is more likely to be left ventricular pacing, whereas if the frontal axis plane is between 0° and −90° and precordial transition occurs by V3, uncomplicated right ventricular apical pacing is said to be present with a sensitivity of 86% and specificity of 99%. In addition, a transition at V4 may indicate middle cardiac vein pacing with a sensitivity of 72% and specificity of 100%. In our patient in Case 2, the frontal axis was between 0° and −90° with a precordial transition occurring by V3. By applying the criteria of Coman et al and Klein et al, as well as arranging for a chest radiograph, we determined that our pacing lead was satisfactorily placed at the right ventricular apex. The lead placement at the right ventricular apex was subsequently further confirmed on echocardiography and on computed tomography pulmonary angiography (done to rule out pulmonary embolism).

The exact mechanism of RBBB QRS morphology in right ventricular apical pacing remains unknown. However, several hypotheses have been proposed. Lister et al postulated that the left ventricle may be activated first through numerous abnormal pathways when the right ventricle is paced. Mower et al suggested that the pacemaker stimulus may enter the right bundle branch and then travel in a retrograde direction to the AV junction and down the left bundle branch. An alternative explanation is that the anatomical right ventricular part of the interventricular septum acts both functionally and electrically as part of the left ventricle. Meanwhile, Barold et al suggested that the RBBB pattern could be due to a combination of right ventricular activation delay due to severe disease of the right ventricular conduction system and early penetration of the electrical impulse into the left ventricular conduction system.

In conclusion, an RBBB pattern on ECG following TPW insertion does not necessarily point towards lead perforation or malposition. An LAO view of the TPW on fluoroscopy after insertion, along with careful analysis of the ECG, chest radiograph and bedside echocardiogram, can facilitate recognition of the lead position in cases of doubt.

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SINGAPORE MEDICAL COUNCIL CATEGORY 3B CME PROGRAMME  
(Code SMJ 202010B)

Question 1. The following are features suggestive of complete heart block:  
(a) Atrioventricular dissociation.  
(b) 1:1 atrioventricular conduction.  
(c) Regularisation of R-R interval in patients with atrial fibrillation.  
(d) Ventricular rate of > 100 beats per minute.  

Question 2. Transient or reversible causes of atrioventricular block include the following:  
(a) Acidosis.  
(b) Myocardial ischaemia.  
(c) Hypothyroidism.  
(d) Congenital atrioventricular block.  

Question 3. Drugs commonly implicated in increased risk of atrioventricular block include:  
(a) Atenolol.  
(b) Diltiazem.  
(c) Atropine.  
(d) Digoxin.  

Question 4. The following are indication(s) for inserting a temporary pacing wire (TPW):  
(a) First-degree heart block.  
(b) Syncope in the setting of high-grade atrioventricular block.  
(c) Haemodynamic compromise in the setting of high-grade atrioventricular block.  
(d) Atrial fibrillation with rapid ventricular rate.  

Question 5. A right bundle branch block morphology after TPW insertion may indicate:  
(a) Left ventricular pacing through intracardiac defects.  
(b) Complications of TPW insertion such as ventricular perforation.  
(c) Malposition of the pacing lead (e.g. in the coronary sinus).  
(d) Normal right ventricular pacing.

Doctor’s particulars:  
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