INTRODUCTION

Ebstein’s anomaly is a rare congenital heart defect characterized by inferior displacement of the septal tricuspid leaflets and atrialization of the right ventricle (RV), and accounts for <1% of all newly diagnosed congenital disorders. It encompasses a wide anatomical and physiological spectrum and the disorder can present itself either as cyanosis in the neonatal life and early childhood or exercise intolerance in the older adult.

The heterogeneous spectrum of presentation of its structural anomalies, and associated cardiomyopathy, accounts for a widely varied clinical and hemodynamic manifestation of the pathology and its impact on timing of diagnosis and prognosis. There is a wide spectrum of pathology with an infinite variety of combinations of severity of the involved structures. Neonatal Ebstein’s anomaly is characterized by reduced survival, while an average risk of mortality per year of 1%–1.4% has been reported in infancy and adulthood. Accessory pathway-related arrhythmia, atrial flutter, and atrial fibrillation are frequent complications; it is clear that atrial flutter and atrial fibrillation (AF) are uncommon in the neonatal period. Medical management and a careful clinical and echocardiographic follow-up are advisable for all asymptomatic patients and those with minimal abnormalities. Radiofrequency catheter ablation (RFCA) is a standard treatment, and depending on the patient’s size, first-line treatment for supraventricular tachycardia supported by pathways and atrioventricular (AV) nodal reentry tachycardia. RFCA is not the treatment of choice for atrial fibrillation unless it is a secondary arrhythmia due to frequent pathway-induced supraventricular tachycardia. Pulmonary vein trigger and left atrial substrate is probably a relatively rare pathogenesis of AF in patients with Ebstein’s anomaly. For the worst spectrum of Ebstein’s anomaly surgical palliation and staging to one or one and a half ventricle repair, but biventricular repair is often not applied. Patients often face revision surgery. Surgical correction is also recommended in the presence of progressive dilation of the right atrium (RA) and RV, development of RV dysfunction, or episodes of paradoxical embolization, reduced exercise capacity, or significant desaturation. RFCA or cryoablation is recommended as first-line treatment for supraventricular and ventricular tachycardia. Prosthetic valve replacement and repair of the tricuspid valve are both common strategies in the correction of tricuspid regurgitation. Mortality is not limited only to the immediate postsurgical period. Only with ample knowledge, this complex and diverse group of patients can be correctly treated in order to improve not only duration, but also quality of life (Geerdink & Kapusta, 2014).
2 | ETIOLOGY

Ebstein’s anomaly is a rare (1–5.2 per 200,000 live births), complex congenital anomaly with a broad pathologic-anatomical and clinical spectrum accounting for <1% of all congenital heart diseases (Bjornard, Riehle-Colarusso, Gilboa, & Correa, 2013; Correa-Villasenor, Ferencz, Neill, Wilson, & Boughman, 1994). Anatomical components of Ebstein’s anomaly (Figure 1).

In 1988, Carpentier et al. proposed the following classification of Ebstein’s anomaly: type A, the volume of the true RV is adequate; type B, a large atrialized component of the RV exists, but the anterior leaflet of the tricuspid valve moves freely; type C, the anterior leaflet is severely restricted in its movement and may cause significant obstruction of the right ventricular outflow tract; and type D, almost complete atrialization of the ventricle except for a small infundibular component.

Celermajer et al. (1992) described an echocardiographic grading score for neonates with Ebstein’s anomaly, extended Glasgow Outcome Scale, with grades 1–4. The ratio of the combined area of the RA and atrialized RV is compared with that of the functional RV and left heart (ratio <0.5, grade 1; ratio of 0.5–0.99, grade 2; ratio of 1.0–1.49, grade 3; ratio ≥1.5, grade 4).

3 | ECG FEATURES IN EBSTEIN’S ANOMALY

3.1 | P wave

Tall P waves (≥2.5 mm) occur in about a third of the patients and are attributable to right atrioventricular atrophy. In close to 50% of cases, the P-wave amplitude is >3 mm. A prolonged P-wave duration is occasionally registered. This pattern has been ascribed to prolonged aberrant conduction in the enlarged RA (Kastor et al., 1975; Macruz et al., 1968). In some cases, giant tall peaked P waves (called “Himalayan” P waves; Kaushik, Sharma, & Kashyap, 2007), abnormal in height (≥5 mm), duration, and shape may be observed.

An electrocardiographic characteristic very suggestive of Ebstein’s anomaly is constituted by the contrast between the giant P wave as opposed to the corresponding atypical right bundle branch block (RBBB) pattern of V1.

3.2 | PR interval

The PR interval can be: (a) short when ventricular pre-excitation is associated with or without delta (δ) wave without a history of paroxysmal tachycardia; (b) prolonged (20% of cases): PR interval duration and the width of the P wave correlate with prolonged conduction in the large RA (Makous & Vander Veer, 1966). First degree AV block occurs in 42% of patients (Ho et al., 2000). Another possibility is intra-atrial conduction disturbance, AV nodal conduction prolongation, or infranodal conduction disturbance (intra His or infra His with RBBB; 3) normal.

3.3 | QRS duration: prolonged

QRS prolongation is the result of prolonged activation of the atrialized RV and is less fully manifest in infants. The conduction disturbance is always distal to the right bundle branch (Kastor et al., 1975) and is sometimes present despite a septal accessory pathway (AP). RBBB is often present after anteroseptal pulmonary artery ablation. One-third of patients with Ebstein’s anomaly and symptomatic tachyarrhythmias have minimal or absent ECG features of ventricular pre-excitation. In these patients, the absence of RBBB pattern is a strong predictor of an AP (Iturralde et al., 2006). Atypical RBBB...
is observed in ~60% of cases and nonspecific intraventricular conduction delay in ~15% in neonates (Barbara, Edwards, Connolly, & Dearani, 2008).

### 3.4 | QRS axis (SAQRS)

It is generally inferior and to the right between +90 and +130°, but the range varies widely from −30 to −170°. The QRS axis is inferior, although a splintered polyphasic QRS makes the axis difficult to determine. On the other hand, when associated with right anterior and inferior AP, the QRS axis is located in the left superior quadrant.

### 3.5 | QRS patterns

Incomplete or complete RBBB is registered in approximately 44% of cases and is thought to result mainly from a paucity of conduction fibers in the atrialized RV with abnormally low R (>7 mm) and S waves over the right precordium. Incomplete or complete RBBB patterns result from infra-Hisian conduction disturbance and abnormal activation of the atrialized RV (Hebe, 2000).

It is frequent to record tri- or tetraphasic QRS patterns (Figure 2). RBBB is observed in ~85% of cases (Figure 3).

Pre-excitation and WPW syndrome are more frequently seen in Ebstein’s anomaly than in any other congenital heart diseases. APs are predominantly right-sided anterior and inferior, manifest, and localize to the lower half of the anatomic tricuspid annulus (Wei et al., 2014; Figure 4). APs are revealed in 6%-30% of patients with Ebstein’s anomaly and can lead to both supraventricular and ventricular tachyarrhythmias. The most common supraventricular tachyarrhythmias are AP-mediated reciprocating tachycardia, AV nodal reentrant tachycardia, and atrial flutter or AF. Tachyarrhythmias cause palpitation, syncope, or sometimes cardiac arrest and are more poorly tolerated due to the hemodynamic and anatomic abnormalities associated with Ebstein’s anomaly. Mahaim nodoventricular fibers are likely to be present when left bundle branch block (LBBB) pattern occurs during sinus rhythm or during an episode of tachycardia (Smith et al., 1982).

### 3.6 | Fragmented QRS

Abnormal conduction through the atrIALIZED RV leads to fragmented QRS (fQRS) on the ECG. Its presence suggests a more severe abnormality and a higher risk of arrhythmia (Park et al., 2013). fQRS disappears after corrective surgery with resection of the atrialized RV (Acharya, Ang, & Gitler, 2017).

Deep Q waves in the right and middle precordial leads (from V1 to V3) are recorded in 50% of the cases. These distinctive Q waves occur because the precordial surface leads record RV intracavitary potentials and usually far leftward as a result of the large size of the RA (Bialostozky, Medrano, Munoz, & Contreras, 1972; Sodi-Pallares, Bisteni, & Herrmann, 1952). This explains the indirect sign of atrIALIZED RV enlargement: qR in V1 (Sodi-Pallares sign). The volumetric increase of the RA gets it closer to the exploring electrode V1, recording initial QRS negativity in this lead, because this electrode records the epicardial morphology of the RA (Sodi-Pallares et al., 1952).

### 3.7 | Arrhythmias

In ~30% of the cases, paroxysmal supraventricular tachycardia (orthodromic or antidromic), atrial flutter, AF, and ventricular tachycardia are registered. Permanent atrial standstill has been reported.
in familial Ebstein's anomaly. (Pierard, Henrard, & Demoulin, 1985). The persistent atrial standstill with slow escape rhythm was most probably a consequence of the abnormalities in both the atrial wall and the His bundle which, together with the abnormal attachment of the tricuspid valve, may be features of the same congenital cardiac anomaly.

4 MAHAIM ACCESSORY PATHWAYS

4.1 Mahaim fibers

The term Mahaim conduction is used to describe decremental conduction connections between the RA or between the AV node and the RV or near the right branch. Although these conductions are rare, their unique electrophysiological properties make their diagnosis and treatment complicated. The true nature of these still obscure pathways makes them particularly interesting from the electrophysiological point of view (Katritsis et al., 2017). In 1941, Mahaim and Winston described the histology of abnormal connections originating from the AV node with RV distal insertion. This was the first description of the Mahaim Node-Ventricular Pathways or "Mahaim Accessory Pathways" (MAPs). Ueshima et al. reported a case of a 7-year-old girl with atriofascicular Mahaim (AFM) pathway concomitant with Ebstein's anomaly. The ECG showed LBBB. Holter showed PR interval prolongation and QRS morphological change during sinus tachycardia. An electrophysiological study demonstrated that the distal His potential appeared earlier than the proximal His potential, which suggested retrograde His conduction through the AV node. Conduction from the Mahaim fiber to the His bundle was faster than that from the AV-node toward the His bundle. The findings allowed a differential diagnosis between AFM and WPW syndrome (Ueshima, Nakamura, Takeno, Miyake, & Takemura, 2017). Ebstein's anomaly is often accompanied by either WPW syndrome or AFM. Therefore, it is important to differentiate between them with regard to treatment and associated risks. AFM pathway indicates AP that demonstrate decremental conduction property without ventriculoatrial conduction (Tchou, Lehmann, Jazayeri, & Akhtar, 1988). Differential diagnosis between AFM and WPW syndrome is very important because both of them cause antidromic AV reciprocating tachycardia; sometimes it can only be diagnosed in the electrophysiology laboratory. One of the points to consider differentiating AFM from WPW syndrome is the change of PR interval. In WPW syndrome, PR interval during sinus rhythm is stable because APs do not exhibit decremental conduction property. Meanwhile in AFM, both PR interval and QRS waveform often change due to decremental conduction property in the Mahaim fiber. In this case, the Holter monitoring demonstrated a change in the PR interval and narrowing of the QRS complex, which supported the diagnosis of AFM rather than WPW syndrome.

Main electrophysiological features in Ebstein's anomaly:

- Intra-atrial conduction disturbance: (RA) P-wave abnormalities, PR interval prolongation
- AV nodal conduction: PR prolongation
- Infranodal conduction: intra-His or infra-His conduction abnormalities
- RBBB distal to the right bundle branch
- Right-sided anterior and inferior AP
- Supraventricular tachycardias
- Atrial fibrillation
- Atrial flutter
- Arrhythmias originating from the atrialized portion of the RV
- Mahaim fibers-mediated arrhythmias
- Deep Q waves from V1 to V4 (it is an indirect signal of RA enlargement)
5 | VECTORCARDIOGRAM

The Frank VCG in Ebstein’s anomaly shows prolongation of the QRS loop/complex owing to RBBB. In patients between 1 day and 31 years old, with the mean age of 8.6 years, the QRS duration was increased in all patients, averaging 103 ms (Ellison & Restieux, 1972). The delay is secondary to RBBB because delayed conduction being primarily in the terminal of the QRS loop/complex. In patients with right-sided anterior and inferior AP, QRS prolongation is a consequence of delayed initial forces. In infants from 1 to 3 days of age there is prolongation of QRS (between 80–100 ms). In normal conditions, the newborns have a QRS duration average of 50 ms.

RV voltages are reduced as shown by decrease in the right maximum spatial voltage (RMSV; 0.6 ± 0.4 mV) and 10–20 ms vectors, and by the predominantly leftward orientation of the HP and FP QRS loops. The RMSV is usually of greater magnitude in patients with little or without tricuspid regurgitation than in those with severe tricuspid insufficiency. The QRS axis in the majority of patients shows a leftward inferior frontal axis. In presence of right-sided anterior and inferior AP, the QRS axis has extreme left axis deviation.

HP: QRS loop rotation is counterclockwise (CCW) or eight-shaped, and the loop is oriented to the left and predominantly posteriorly. This reflects the reduction in the RV potentials.

In the presence of right-sided anterior and inferior AP, initial conduction delay is observed (6 loop) and the early forces directed to the left.

FP: QRS axis is located in the left inferior quadrant, and the QRS loop rotation is clockwise or eight-shaped. In the presence of right-sided anterior and inferior AP, the QRS axis has extreme left axis deviation.

LSP: Rotation is predominantly CCW and the reduction in anterior forces could be seen. P loop is oriented in the normal direction, to the left, inferiorly and anteriorly. Clear right atrial enlargement loop is the rule. The T loops are directed predominantly to the left and posteriorly; 50% directed inferiorly and 50% directed superiorly.

6 | CONCLUSIONS

Ebstein’s anomaly encompasses a wide anatomical and physiological spectrum. Accessory pathway-related arrhythmia, atrial flutter, and atrial fibrillation are frequent complications, although that atrial flutter and atrial fibrillation are uncommon in the neonatal period. The ECG is of great value to raise the suspicion of Ebstein’s anomaly, and it is very useful to guide future diagnostic imaging methods, both noninvasive and invasive. The association of giant P waves with prolonged PR interval and atypical, low voltage RBBB in the right precordial leads constitute the hallmark of this entity. Pre-excitation and WPW syndrome are more frequently seen in Ebstein’s anomaly than in any other congenital heart diseases, and APs can lead to both supraventricular and ventricular tachyarrhythmias. Ebstein’s anomaly is also often accompanied by AFM, and certain ECG features help to distinguish between the WPW syndrome and AFM. The VCG is a relevant additional diagnostic tool, because it allows for the diagnosis of ventricular pre-excitation (right-sided anterior and inferior AP) with the presence of comets very close to one another in the initial part of the QRS loop.

CONFLICTS OF INTEREST

None.

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