Modified Cone Reconstruction of the Tricuspid Valve for Ebstein Anomaly as Performed in Siberia

The cone reconstruction technique, first described by da Silva and modified by Dearani and by others, has become the repair method of choice in patients with Ebstein anomaly of the tricuspid valve. This report details the outcome of the modified cone reconstruction technique in 6 children who underwent surgical correction of Ebstein anomaly at the Tomsk Institute of Cardiology in Siberia.

From 2012 through 2015, 4 boys and 2 girls (age range, 11 mo–12 yr) underwent surgery to correct Ebstein anomaly. All had presented with cyanosis, exertional dyspnea, fatigue, or new-onset atrial arrhythmia, and none had undergone previous cardiac surgery. All survived the operation. One patient needed tricuspid valve replacement with a bioprosthesis after early breakdown of the cone reconstruction. As of December 2016, all the patients had no symptoms, tricuspid stenosis, or arrhythmia. This series indicates that cone reconstruction—the most anatomic repair technique for the dysmorphic Ebstein tricuspid valve—can be successfully performed in pediatric heart centers with a large experience.

In 1866, Wilhelm Ebstein described autopsy findings of a strikingly abnormal tricuspid valve (TV).1,2 In the 1950s, this lesion was termed Ebstein anomaly or Ebstein malformation. After many surgeons unsuccessfully attempted to repair Ebstein anomaly through the years, da Silva described the cone reconstruction (CR) technique.3,4 Modifications of this technique have been used by Dearani and by others,5–9 and CR has become the preferred repair method for patients with Ebstein anomaly. We describe the procedure and report the outcomes of the initial 6 patients who underwent modified CR at a major pediatric cardiac center in Tomsk, Russia.

Patients and Methods

The Tomsk Institute of Cardiology, a large referral center in Siberia for congenital heart disease surgery, serves patients from Russia and several former Soviet republics. Consent for treatment and data-sharing was obtained for the patients involved in this report.

Patients. From 2012 through 2015, 4 girls and 2 boys (age range, 11 mo–12 yr) underwent corrective surgery for Ebstein anomaly at the Tomsk Institute of Cardiology. All presented with cyanosis, exertional dyspnea, fatigue, or new-onset atrial arrhythmia. None had undergone prior cardiac surgery. One patient with Wolff-Parkinson-White syndrome needed preoperative radiofrequency catheter ablation. The primary surgeon (EVK) performed CR with use of Dearani modifications8,9 that he had learned at the Mayo Clinic in 2012.

Surgical Technique. The modified CR technique involves surgical delamination (dissection) of all viable leaflet tissue from the myocardium and approximation of the sides of the leaflets to form a cone that completely surrounds the tricuspid orifice in the atrioventricular (AV) junction.5,8,9 The reconstructed valve is anchored at the hinge point of the right AV groove.

The most important aspect of CR is to free all fibrous and muscular attachments between the leaflet bodies and the myocardium while preserving the fibrous attachments of the leaflets’ leading edges, particularly the chordae tendineae. Surgical fenestrations enable improved leaflet mobility while these chordal attachments are maintained.
Delamination is started at the 12 o’clock position of the anterior leaflet of the TV, just below the true anatomic annulus, and is continued clockwise toward the inferior leaflet. After the septal leaflet has been delaminated, the mobilized inferior leaflet is rotated clockwise and approximated to the leading edge of the septal leaflet. This creates a 360° cone of leaflet tissue, which gives the reconstructed TV hinge points at the true anatomic annulus. Care is taken throughout to avoid injuring the right coronary artery, especially during plication of the atrIALIZED right ventricle (RV), and to prevent heart block. Adding a partial or complete annuloplasty ring can reduce postoperative valvular regurgitation.

Results

All 6 patients survived surgery. Intra-atrial communications were closed in each. Five of the 6 underwent successful CR and were discharged from the hospital with excellent results. A 12-year-old girl had unfavorable leaflet anatomy that led to early breakdown of the CR; her TV was replaced with a bioprosthesis.

The longest follow-up duration was 4 years. As of December 2016, the 5 patients with enduring CR had no symptoms, arrhythmias, right-sided heart failure, or tricuspid stenosis, and only mild tricuspid regurgitation. In 2 patients, RV systolic function remained mildly depressed.

Figures 1 and 2 show the preoperative anatomy in a 3-year-old girl from this cohort. Images obtained before her hospital discharge (Figs. 3 and 4) reveal the excellent surgical result: the TV leaflets residing at the level of the true anatomic annulus, minimal residual tricuspid regurgitation, and a large portion of the RV below the level of leaflet coaptation in comparison with the preoperative findings. Qualitative RV systolic function was good in this patient after CR.

Discussion

In correcting Ebstein anomaly, surgical goals include complete or subtotal closure of intra-atrial communications, TV repair or replacement, elimination of arrhythmia, plication of the atrIALIZED RV, right reduction atrioplasty, and repair of associated defects (such as closing ventricular septal defects and relieving RV outflow tract obstruction). Cone reconstruction, the most anatomic repair technique described to date, has revolutionized TV repair in patients with Ebstein anomaly.

In 2007, da Silva and colleagues reported the cases of 40 patients who underwent correction of Ebstein anomaly by means of CR. In 2013, Anderson and colleagues reported that 98% of Mayo Clinic patients younger than 21 years of age who had undergone CR were discharged from the hospital with good results.

To date, reports of postoperative arrhythmias have been rare in this patient population.

No one in our series needed a bidirectional cavopulmonary anastomosis (BDCPA) at the time of CR. A BDCPA is indicated when there is severe RV enlargement or dysfunction, compression of the left ventricle because of a shift of the interventricular septum, or moderate TV stenosis after CR (mean gradient, ≥8 mmHg); or when the right atrial-to-left atrial pressure ratio is >1.5, indicating poor RV function.

In the Mayo Clinic series, approximately 30% of patients who underwent CR also needed a BDCPA, which is not ideal because it can cause pulsatility of the head.
and neck veins, facial swelling, and the development of venovenous collateral vessels and pulmonary artery venous fistulae. In addition, BDCPA procedures preclude access to the heart from the internal jugular vein in future cardiac catheterization or electrophysiologic procedures.\textsuperscript{11,12}

One patient, a 12-year-old girl, underwent TV replacement with a bioprosthesis after the CR failed. Her anterior leaflet was heavily muscularized, and there was no septal leaflet delamination. Patients with Ebstein anomaly and this valvular morphology are challenging candidates for any form of TV repair. However, CR can be viable for other patients who have recurrent, severe tricuspid regurgitation after previously attempted TV repairs.\textsuperscript{8}

This case series indicates that other pediatric cardiac centers can adopt the CR technique for Ebstein anomaly with reproducible results. However, mentoring in the nuances of Ebstein-anomaly repair is essential. Care needs to be given to the right posterior AV groove to avoid injury to the right coronary artery, and one must meticulously avoid causing heart block. The Tomsk Institute’s lead surgeon (EVK) spent considerable time studying CR techniques at the Mayo Clinic. Despite the steep learning curve, CR has greatly improved the treatment of children with Ebstein anomaly, providing the most anatomic repair of the dysmorphic TV.

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**References**


