Surgical correction of cor triatriatum sinister in the paediatric population: mid-term results in 15 cases

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Abstract

OBJECTIVES: Cor triatriatum sinister (CTS) is a rare developmental cardiac abnormality resulting in left ventricular inflow obstruction. In this report, we aimed to present our mid-term results of early childhood patients operated for CTS and associated cardiac abnormalities.

METHODS: We enrolled 15 patients with CTS who were operated by a single surgeon between 2001 and 2013. A retrospective analysis was performed in order to determine the demographics, operative and postoperative results of the patients. The median age was 14 months and the median body weight was 8.2 kg at the time of operation.

RESULTS: Fourteen patients had concomitant cardiac pathology. Three of the patients had atrial septal defect and 1 of the patients had partial abnormal pulmonary venous connection, whereas 4 of the patients had both. In 2 cases of complete atrioventricular septal defect, 1 case with ventricular septal defect, 1 case with patent ductus arteriosus, 1 case with double outflow right ventricle and another case with tetralogy of Fallot, complete repair was performed together with membrane resection in the left atrium (LA). The mean preoperative left atrial gradient was 13.3 mmHg, whereas the mean LA pressure at the first postoperative year was 4.2 mmHg. There was 1 case with early mortality due to septic multiorgan failure secondary to pneumonia.

CONCLUSIONS: CTS is a rare congenital cardiac anomaly in which the results of the corrective surgery are encouraging. Early and long-term outcomes may be variable according to the associated congenital heart defects.

Keywords: Cor triatriatum sinister • Congenital heart defects • Cardiac surgery

INTRODUCTION

Congenital obstructions of the left ventricle inflow can be classified into four subtypes: (i) congenital mitral stenosis, (ii) cor triatriatum sinister (CTS), (iii) pulmonary venous (PV) obstruction and (iv) supramitral ring [1]. CTS is encountered with an incidence of 0.1–0.4%, in congenital heart diseases (CHDs) [2].

The first post-mortem case of an abnormal membrane in the left atrium (LA) was described by Church in 1868 [3]. CTS is characterized by the presence of a fibromuscular membrane that subdivides the LA [4]. The postero-superior proximal cavity receives the pulmonary veins, whereas the anteroinferior chamber (true LA) contains mitral valve and left atrial appendage (LAA) [5]. Rarely, this membrane may divide the LA into multiple chambers leading to cor polyatriatum [6].

Clinical presentation and symptoms of patients with CTS are mainly due to obstruction at the PV-LA junction, which leads to a pressure overload at the right side of the heart [7]. The severity of the clinical symptoms depends on the size and the number of fenestrations on the obstructing membrane as well as the presence and location of the atrial septal defect (ASD) and associated CHD [8]. The blood flow between the proximal chamber and the distal left atrial cavity can be through three paths: from the proximal chamber to the right atrium (RA), from the RA to the distal LA or through the fenestrations on the membrane. Figure 1 schematically demonstrates the fibromuscular membrane dividing the atrial cavity into two separate spaces and possible paths of interchamber communications.

Most of the cases with CTS are diagnosed in infancy, whereas in cases with wide orifice and non-significant pressure drop through the membrane, the diagnosis of CTS may be challenging until late adulthood due to missing symptoms [9–11]. Since calcification at the fenestrations may hinder the blood flow between the proximal and distal chambers, CTS could be detected in adulthood with atrial fibrillation and symptoms of mitral stenosis [12, 13].
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through it. For cases where both cardiac catheterization and trans-
reserved for detection of the associated CHD, not all patients went
transthoracic echocardiography. Since cardiac catheterization was

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months (range 1 month to 7 years) and median body weight was

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33 kg). Eleven of the patients (73%) were infants.

MATERIALS AND METHODS

A retrospective analysis was performed in order to determine the
patients with a history of surgical intervention for CTS following
the approval of the ethical committee. Clinical presentations, pre-
operative echocardiographic findings, perioperative course and
follow-up records were evaluated.

Fifteen patients (7 male and 8 female) with CTS were operated
by a single surgeon between 2001 and 2013. Median age was 14
months (range 1 month to 7 years) and median body weight was

8.2 (range 3–33 kg). Eleven of the patients (73%) were infants.

RESULTS

We encountered 14 cases with classical CTS (Type A) where the
proximal chamber was receiving the pulmonary venous return.
Seven of these patients had ASD, all of which were located
between the proximal chamber and RA. One patient had pulmonary
venous connection of left-sided pulmonary veins into coronary
sinus (CTS, Type B).

In our patient population, 14 of the patients (93%) had com-
comitant cardiac pathology. Three of the patients had ASD and 1
of the patients had partial abnormal pulmonary venous connec-
tion (PAPVC), whereas 4 of the patients had both ASD and PAPVC.
In cases with PAPVC, the abnormal pulmonary veins were draining
into the RA (right superior PV) or coronary sinus (left superior and
inferior PV). All the ASDs were between the proximal chamber
and RA in our patients. There were 2 patients with concomitant
complete atrioventricular septal defect (AVSD—Rastelli type C), 1
with ventricular septal defect, 1 with patent ductus arteriosus
(PDA), 1 with double outlet right ventricle (DORV) and another
patient with tetralogy of Fallot (TOF).

The mean preoperative LA gradient that was 13.3 mmHg (range
6–21), which was calculated by either cardiac catheterization or
transthoracic echocardiography. Since cardiac catheterization was
reserved for detection of the associated CHD, not all patients went
through it. For cases where both cardiac catheterization and trans-
thoracic echocardiography were implemented, we took the
transmembrane pull-back gradient results of the cardiac catheteriza-
tion into account regardless of the echocardiographic calculations.

The preoperative symptoms ranged from mild respiratory dis-
tress to critical congestive heart failure. The surgical indication for
intervention was predominantly determined by the accompany-
ing symptoms and the concomitant CHD. The only case where
the surgical intervention was deemed necessary based merely on
symptoms (exertional dyspnoea and fatigue) was a 7-year-old child
with isolated CTS, whose pressure gradient across the mem-
brane was 21 mmHg at echocardiographic evaluation.

Eight of the patients (53%) had preoperative pulmonary hyperten-
sion, which was either detected by echocardiography or cardiac
catheterization. The pulmonary hypertension was defined as
follows: where (i) the mean pulmonary artery pressure (PAP) was
over 25 mmHg, if measured by cardiac catheterization, or (ii) the
ratio of systolic PAP to systolic systemic arterial pressure was more
than 0.4 as a result of transthoracic echocardiographic evaluation.
The PAPs were within normal limits in the follow-up period for all
patients, except for a 4-year-old patient with PDA and CTS, who
had moderate pulmonary hypertension that was medically managed.

Cardiopulmonary bypass (CPB) with mild-to-moderate hypo-
thermia was implemented in all cases. Mean aortic cross-clamping
time and total CPB time were 55 ± 24 and 79 ± 41 min, respective-
ly. Following the cross-clamping of the aorta, cold antegrade
blood cardioplegia was administered where intermittent doses
were applied every 15 min.

Our surgical procedure incorporated resection of the mem-
brane at the left atrial cavity and a complete correction of the ac-
companying CHD. There was one exception to this approach, in
which the membrane at the LA was resected with palliative pulmo-

nary artery banding in a newborn with AVSD. All patients presented
with ASD and/or PAPVC underwent atrial resection. In all cases
with PAPVC, the pulmonary venous return was rerouted to the LA.
A patient with DORV and another with TOF underwent complete
repair together with membrane resection at the LA. Two patients
with AVSD (both Rastelli type C) were successfully operated. One
patient underwent complete repair with a modified single patch
along with LA membrane resection, whereas LA membrane resec-
tion with pulmonary artery banding was the choice of treatment
modality for a newborn patient with unbalanced ventricles. We
operated 1 patient with ventricular septal defect and another with a
PDA with concomitant membrane resection in the LA.

A right atrial approach was preferred in 13 cases, all of which
included concomitant CHD. In a young patient with isolated CTS,
the membrane was excised with a left atriotomy, whereas in
another case with a concomitant persistent left superior vena cava
(PLSVC), a combined bialtrial approach was performed in order to
facilitate the surgical exposure.

The early postoperative results were evaluated using transthora-
oric echocardiography at the first postoperative day, before dis-
charge and at the third postoperative month. Afterwards, annual
transthoracic echocardiographic evaluations were recorded. The
mean postoperative first year LA pressure was 4.2 mmHg (range
2–6). The preoperative LA gradients and follow-up LA pressures
are shown in Figure 2. None of the patients needed postoperative
reintervention following the initial repair.

Early postoperative mortality (8%) was encountered in a patient
with VSD and CTS due to septic multiorgan failure secondary to
pneumonia 32 days after surgery. The postoperative transthoracic
echocardiography evaluation of this patient revealed an intact
ventricular septal patch without any residual LA gradient. We did
The CTS may be associated with other CHD in 24–80% of the cases [21]. Partial or total anomalous pulmonary venous connection is the most commonly associated cardiac pathology, which has been reported in nearly one-third of the patients with CTS [21]. We encountered PAPVC and ASD as the leading associated CHDs in our patient population.

Surgical correction is indicated in patients with obstructive symptoms regardless of age [21]. The urgency of operation is primarily determined by the severity of the presenting symptoms, but CTS can remain asymptomatic until late adulthood. Therefore, it is reasonable to observe patients with an incidental diagnosis, provided the opening in the diaphragm is non-restrictive and regular follow-up is possible [21]. The specific surgical approach is chosen primarily based on the presence of associated lesions and the size of the atria. In our surgical experience, we predominantly preferred a right atrial approach for the resection of the membrane as well as the correction of the associated intracardiac pathologies. In elder children with isolated CTS and an enlarged LA, a left atriotomy may be preferred. A combined biatrial approach could be necessary in order to facilitate the surgical exposure in selected patients.

In cases of CTS, the timing of surgical intervention primarily depends on the severity of stenosis at the left atrial venous return and the coexistence of other CHD. We operated two critically ill patients during the newborn period. In cases where the membrane at the LA does not lead to severe stenosis, the coexisting CHDs predominate the timing of surgery. The increased utilization of the transthoracic echocardiography in the infancy led us to encounter low age and body weights at the time of diagnosis and surgical repair. Eleven of our cases were operated at the infancy period. The follow-up echocardiography measurements revealed encouraging results with a mean left atrial pressure below 5 mmHg.

Although percutaneous balloon dilatation of the obstructing membrane in simple CTS may be performed in selected cases at older ages [22], most of the patients have co-existing CHDs, which will eventually need to be treated surgically.

**CONCLUSION**

CTS is a rare cardiac anomaly that can be detected in any age. Correct preoperative diagnosis of the membrane separating the LA and the associated cardiac anomalies is important in the management of these patients. Early and long-term results may be variable according to the associated CHD; otherwise, the mid-term results are excellent.

**Conflict of interest:** none declared.

**REFERENCES**


**DISCUSSION**

CTS is a rare morphological cardiac abnormality. The embryological basis of CTS remains controversial along with three possible explanations. The first one is the idea of malincorporation that suggests that the common pulmonary vein fails to incorporate into the LA, which results in the development of an obstructing fibromuscular membrane at the junction of these structures during the fifth week of gestation [14, 15]. Although this theory is widely accepted, it fails to explain the presence of fossa ovalis and atrial muscular fibres in the proximal chamber [16]. The malseptation theory proposes that the abnormal membrane that is located at the LA is a result of septum primum overgrowth. The theory of entrapment considers that the left horn of the sinus venosus entraps the common pulmonary vein, which prevents its incorporation into the LA [17]. As a variation of the entrapment theory, the impingement of the PLSVC on the LA is suggested to result in the production of the obstructing membrane in the LA [13, 18].

Two basic classifications are favoured in CTS. Loffler [19] first classified CTS into three groups in 1949, depending on the size and number of fenestrations at the membrane: (i) no communication between proximal and distal chambers, (ii) communication through small perforations at the membrane and (iii) incomplete membranous subdivision of the LA. Lam et al. [20] suggested another classification in 1962. Type A (also named as the classical form) consists of a proximal chamber that receives all the pulmonary veins and a distal chamber that receives the LAA and the mitral valve. There is no ASD and the proximal and distal chambers connect through the small holes on the membrane in the classical form. The subtype A1 defines an ASD between the RA and the proximal chamber, whereas subtype A2 defines an ASD between the RA and the distal chamber. Type B defines hearts in which the pulmonary veins drain into the coronary sinus, which is also accepted as a variant of total anomalous pulmonary connection. Type C defines hearts in which the proximal chamber receives no pulmonary veins (the rarest variant) [20]. In our patient population, all the patients were classified as Type A, with an exception of 1 case classified as Type B.


