Isolated and Syndromic Congenital Sternal Cleft

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Introduction
Chest wall malformations represent a large phenotypic spectrum. Among them, congenital sternal clefts are rare abnormalities, and have been classified as complete (the rarer form) or incomplete. The incidence is of 1:100,000 cases per live birth, and represents about less than 1% of all chest wall deformities [1].

In embryonic life, the sternum is derived from the lateral plate mesoderm of either side of midline, which becomes fused during the 10th weeks. The sternum development appears during the 6th week of embryology as a pair of condensed and parallel mesenchymal bands. Cells migrate from two lateral plates on either side of the anterior chest wall to fuse in the midline in a craniocaudal fusion. The original sternum is cartilaginous then multiple ossification centers develop in a craniocaudal sequence (Figure 1). This anomaly results from a failure fusion of the sternum. Very rarely, the sternal bars fail to join in the midline, which results in a complete or incomplete sternal cleft. The clinical severity of sternal cleft ranges from the benign sternal cleft to the potentially lethal ectopia cordis. Generally sternal cleft is observed at birth. The prognostic depends mostly on associated abnormalities as congenital cardiac diseases. It
leads to severe cardiovascular disturbance during anesthesia particularly when pulmonary hypertension is associated [2]. Sternal cleft may be associated also with defects in the anterior chest wall such as ectopia cordis, or Cantrell’s pentalogy, with a combination of defects involving the abdominal wall, sternum, diaphragm, pericardium and heart, possibly diagnosed prenatally with three-dimensional ultrasonography.

Surgery is indicated to protect the heart and major vessels, of injuries by external trauma. It improves respiratory dynamics and esthetics. The challenge to the anesthetist is the risk of paradoxical respiration especially if the defect is incomplete.

We report our experience in the treatment of sternal abnormalities and we discuss the results and evolutions of these patients.

Materials and Methods

From 1994 to 2015, we reviewed five cases of neonatal sternal defects. The majority of patients were girls (4:1). Four cases were discovered at birth and one prenatally and confirmed by MRI. The mean weight at birth was 3.2 kg (2.2-4.1). Sternal abnormalities types were: - Upper sternal cleft in two cases; one case isolated abnormality, and the other associated with PHACES syndrome (posterior fossa brain malformations, facial hemangiomas, arterial anomalies, cardiac defects, eyes abnormalities, sternal cleft, and supra umbilical raphe) - Lower sternal cleft in two cases; one associated to PHACES syndrome (Figure 1), intestinal atresia and gyral pattern disorders. The other abnormality associated as part of a field defect known as Cantrell’s Pentalogy (ectopia cordis, deficiency of the anterior diaphragm, supra umbilical abdominal wall defect, defect in lower sternum, and congenital intra-cardiac abnormalities) - One total sternal cleft associated with Cantrell’s pentalogy. Characteristics of patients are reported in the Table 1.

All patients were studied preoperatively by chest X-ray, computerized tomography, cardiologic evaluation, genetic evaluation. When the sternal abnormality was associated to hemangiomas, cerebral MRI and ophthalmic evaluation were done (Figure 2).

In three cases, surgical correction by primary closure was done (two upper and one complete). Theses patients presented retraction at the defect with inspiration and protrusion with exhalation or Valsalva maneuver.

During surgery, a cerebral and splanchnic monitoring was evaluated with the use of near infrared spectroscopy (NIRS). Postoperatively, all infants were admitted to the neonatal intensive care unit.

Surgical approach

Under endo-tracheal general anesthesia, the patient was placed in decubitus position, the skin incision was vertical on the midline (Figure 3), and the pectoral muscles were dissected to expose the sternal bars and then, the medial edges of the two sternal halves were freed from the underlying pleura and pericardium. The inferior aspect of partial sternal cleft was incised, when U shaped, to make closure easier. The sternal bars were approximated on the midline by non-absorbable sutures (Figure 4).

If complete sternal cleft was too wide or there was a mediastinal compression after sternal bar juxtaposition, we closed the defect by placing non-absorbable prostheses in multiple layers and then, the defect and the primary repair was delayed using the technique described previously.

Results

Surgical correction was performed in 3 of them. During primary closure, none signs of compression of the mediastinum appeared. None intra-operative complications occurred. Near infrared spectroscopy indices stayed normal and regular all along the surgeries. Mean operative time was 120 min (100-150). In cases of upper sternal cleft, primary closure was performed at 5 and 10 week-old respectively. In complete sternal cleft, prosthetic closure was done at 14 days-old, and the primary repair was done seven months later. In cases of lower sternal cleft non-surgical
correction was necessary due to asymptomatic patients and minor defect.

The mean follow-up period was 9.5 years (3.6-21 years). No complications were reported. All patients achieved good functional and cosmetic results after correction.

**Discussion**

Sternal cleft malformation is caused by a fusion failure of the sternal elements. The aetiology is unknown, thought to arise from a failure of ventral fusion of the sternal bars during the 8th week of gestation. There is a female predominance. Alcohol intake and methylcobalamine or riboflavine deficiency is associated with sternal cleft, and studies in mice indicate abnormalities in HOX b gene expression as a possible factor [3].

Sternal diagnosis is prenatally difficult with ultrasonography. Twomey considers this abnormality as a thinned and sunken midline anterior chest wall with intact skin coverage but no identifiable cartilaginous sternum [4].

Sternal clefts are classified as being total or partial. The partial malformation can be superiorly or inferiorly located, with the upper sternum being the most common and almost always isolated abnormality [5]. Occasionally it can be associated with PHACES syndrome [6]. The rare inferior variety is often associated with a thoraco-abdominal *ectopia cordis*, being part of a Cantrell’s Pentalogy [7].

Superior clefts have orthotopic normal heart, normal skin coverage, and intact pericardium. With isolated cleft, the patient is usually asymptomatic, except during cries or coughs (upper and midline thoracic depression during inspiration and a prominence during expiration) [1]. When a sternal cleft is diagnosed, associated anomalies must be searched by careful physical examination, chest X-ray and echocardiography, because the prognostic changes if the cleft had associated abnormalities. Genetic evaluation can be useful [6].

When haemangioma was associated, other criteria’s of PHACES syndrome must be searched and then, a cerebral RMI and an ophthalmologic evaluation are completed. A consensus statement on diagnostic criteria was published in 2009 [6].

Regarding Cantrell’s Pentalogy, the prognosis depends mainly on the cardiac abnormalities [7].

Even for asymptomatic patients, sternal surgery is necessary to offer protection to the heart, to prevent recurrent respiratory infections due to paradoxical respiratory movements, to maintain the growth potential of the chest wall and to offer a good cosmetic outcome. Chest computerized tomography or MRI is preoperatively required to confirm the presence of a sternal cleft and establish the type (presence or not of bar inferiorly) and then to plan the surgery [8].

Surgical correction is indicated to protect the heart and major vessels, to improve respiratory dynamics and aesthetic profile. Only in very minor defects, particularly in lower sternal cleft, conservative treatment can be propose for asymptomatic patients. In our series, in two cases of lower sternal cleft, non-surgical correction was done with uneventful course.

The patient age is an important factor for primary repair, preconized if possible in the neonatal age to achieve primary closure. In some cases the reduction in thoracic volume can cause cardiovascular impairment. It is validated that after 3 months of age, the chest wall becomes relatively rigid and more complicated techniques may be required such as use of prostheses, partial or total thymectomy, sliding chondrotomies, and clavicle dislocation. These techniques can be an alternative to primary closure if the latter is challenging or impossible due to a stiff thorax [1].

For patients 1 and 2, sternal surgery was accomplished respectively at 10 weeks and 40 days, by primary closure alone. For patient 4, primary closure was impossible at 14 days because of a great distance between the sternal halves. A prosthetic material was necessary, and direct approximation of the sternal halves was performed at 7 months.

Many authors agree to say that primary repair is best accomplished in the neonatal period, benefiting from of the pliability of the chest wall and less compression of underlying structures. If signs of mediastinum compression appear during surgery, primary repair should be terminated for a more elaborate approach (sliding or rotating chondrotomies, pectoralis major myoplasty, clavicular fracture, or even autologous tissue flaps, myocutaneous flaps, prosthetic materials) [5,8,9].
Table 1. Data base of patients (ICA: internal carotid artery; ACA: Anterior Cerebral Artery; PCA: Posterior Cerebral Artery).

<table>
<thead>
<tr>
<th>PATIENTS</th>
<th>Year of birth</th>
<th>Year of birth</th>
<th>Sex</th>
<th>Gestational weeks (WG)</th>
<th>Birth weight (g)</th>
<th>Associated Syndrome</th>
<th>Type of cleft</th>
<th>Phenotype</th>
<th>Associated anomalies</th>
<th>Sternal surgery</th>
<th>Follow-up</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2007</td>
<td>2012</td>
<td>M</td>
<td>38</td>
<td>2200</td>
<td>Superior</td>
<td>Superior</td>
<td>Large left and right frontotemporal segmental facial hemangiomas, abdominal and thoracic hemangiomas, hypoplasia of the right ICA, supraventricular raphe, optic nerve hypoplasia</td>
<td>Jejunal atresia, gyration disorders</td>
<td>Stomal closure alone</td>
<td>3 years and 10 months: Good evolution with propranolol, normal neurodevelopment, ingestive angiomatosis, treatment with enterocolitis, jejunal atresia, digestive haemorrhage, digestive angiomas.</td>
</tr>
<tr>
<td>2</td>
<td>2012</td>
<td>2012</td>
<td>F</td>
<td>38</td>
<td>3300</td>
<td>Superior</td>
<td>Inferior</td>
<td>Large left and right frontotemporal segmental facial hemangiomas, abdominal and thoracic hemangiomas, hypoplasia of the right ICA, supraventricular raphe, optic nerve hypoplasia</td>
<td>Jejunal atresia, gyration disorders</td>
<td>Stomal closure alone</td>
<td>3 years and 6 months: Ulcero-necrotizing enterocolitis, jejunal atresia, digestive haemorrhage, digestive angiomas, treatment with propranolol, seizures controlled by antiepileptic, normal neurologic examination.</td>
</tr>
<tr>
<td>3</td>
<td>2004</td>
<td>2004</td>
<td>F</td>
<td>38</td>
<td>2810</td>
<td>Superior</td>
<td>Inferior</td>
<td>Large left and right frontotemporal segmental facial hemangiomas, abdominal and thoracic hemangiomas, hypoplasia of the right ICA, supraventricular raphe, optic nerve hypoplasia</td>
<td>Jejunal atresia, gyration disorders</td>
<td>Stomal closure alone</td>
<td>10 weeks: Primary closure alone</td>
</tr>
<tr>
<td>4</td>
<td>1994</td>
<td>2012</td>
<td>F</td>
<td>39</td>
<td>3450</td>
<td>PHACES</td>
<td>Inferior</td>
<td>Small omphalocele, anterior diaphragmatic defect</td>
<td>Complete common mesentry</td>
<td>Prosthetic closure at 14 days, primary closure at 7 months</td>
<td>40 days: Primary closure alone</td>
</tr>
<tr>
<td>5</td>
<td>2004</td>
<td>2004</td>
<td>F</td>
<td>38</td>
<td>4184</td>
<td>Kanavel's Reform</td>
<td>Inferior</td>
<td>Small omphalocele, anterior diaphragmatic defect</td>
<td>Complete common mesentry</td>
<td>Prosthetic closure at 14 days, primary closure at 7 months</td>
<td>40 days: Primary closure alone</td>
</tr>
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In conclusion, congenital sternal clefts are uncommon pathologies and require a multidisciplinary approach to optimize the patient’s outcome. They can be isolated or part of a syndrome, justifying complementary explorations. The surgical approach is not required for all patients. Surgical treatment depends on the type of sternal malformation. Best results are obtained during the neonatal period, when the chest wall is maximally flexible to make the repair easier. Further, if necessary, a prosthetic closure can resolve the malformation in better conditions.
References


