Case Report

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Sternal reconstruction by extracellular matrix: a rare case of phaces syndrome

DOI 10.1515/med-2016-0037
received January 27, 2016; accepted March 7, 2015

Abstract: Congenital defects of the sternum are rare and due to a failure of midline development and fusion of the sternal bones. Surgical correction of a sternal cleft should be preferred during infancy for functional reasons. Chest wall reconstruction represented a complex problem in the last decades. We report our successful outcome of sternal reconstruction in a rare case of PHACES syndrome, in which the patient was submitted to reconstruction of the sternum and complete closure of the thoracic defect by the employ of an extracellular matrix XCM Biologic tissue matrix.

We promote the use of extracellular matrix in surgical reconstruction of chest defects for its maneuverability, plasticity, tolerability and the possibility of growing with the children’s chest getting a good compliance and optimal cosmetic results.

Keywords: Phaces syndrome, Sternal reconstruction, Midline development defects.

1 Introduction

Congenital defects of the sternum are rare and due to a failure of midline development and fusion of the sternal bones. The sternal cleft is more common in females and in association with other anomalies like posterior fossa malformations, hemangiomas, arterial anomalies, cardiac defects, eye anomalies and supraumbilical midline raphe better known as PHACES syndrome (OMIM 606519) [1]. Surgical correction of a sternal cleft should be preferred during infancy for functional (respiratory impairment and potentially dangerous injuries to the mediastinal organs) and mechanical (more elasticity and thoracic compliance) reasons [2]. Chest wall reconstruction represented a complex problem in the last decades due to the difficulty to choose the best surgical approach for closing the defect without compromising the stability and the following evolution of the thoracic wall. We report our successful outcome of sternal reconstruction by bioprosthetic mesh (extracellular matrix) in a rare case of PHACES syndrome.

2 Case report

A 4 year-old girl was referred to our hospital with a congenital defect of the anterior chest wall. Her antenatal history was unremarkable and family history revealed parents as cousins of first grade, no traumas or surgical operations were referred. At delivery, the birth weight was 2600 g and no episode of respiratory distress was reported. At clinical examination she presented a wide gap in the upper part of anterior wall chest with intermittent herniation of cardiac structures during crying, while its pulsations were easily seen at rest, and also it was evident a paradoxical excursion of the thorax during respiratory acts (Figure 1). Cutaneous examination revealed an erythematous plaque as left sided facial hemangiomas extending to neck (Figure 2 a-b) and a cutaneous raphe from sternal defect to umbilicus (Figure 1). Ophthalmologic, routine blood
samples, genetic analysis, thyroid profile and cerebral cranial magnetic resonance were normal. Cardiovascular evaluation revealed a small atrial septal defect hemodynamically stable.

Chest-X ray and computed tomographic scan (CT) with three-dimensional reconstruction showed the presence of a congenital diastasis of sternal manubrium (37 mm) with agenesis of the corpus sternum, with V-shaped cartilages (24 mm cranio-caudal diameter) in whose concavity the right heart ventricle was made superficial (Figure 3 a-b). After an accurate surgical planning, the patient was submitted to reconstruction of the sternum and complete closure of the thoracic defect by the employ of an extracellular matrix XCM Biologic tissue matrix®. The skin overlying the sternal defect was incised along the midline from the ideal manubriosternal joint to the ideal xiphisternum separating the subcutaneous tissues from the underlying fused pericardium. Then the pericardium was mobilized all along the edges and sutured in midline without tension. The second layer was provided by an extracellular matrix previously modeled and then anchored in double layers as a sandwich, at the medial margins of the bilateral ribs with multiple intermittent not absorbable suture (Figure 4). The pectoral muscles were reapproximated to the midline and drainage was positioned before closure the skin that was removed on the third day. The post-operative recovery was regular even if there was an episode of bradycardia during extubation treated with administration of atropine and epinephrine with rapid cardio-respiratory stabilization. The patient was discharged on the 8th postoperative day with a good cosmetic and functional

Figure 1: Sternal cleft in the upper part of anterior wall chest associated with supra umbilicus raphe.

Figure 2 a-b: Left sided facial hemangiomas extending to neck.

Figure 3 a-b: Three-dimensional reconstruction of CT images showed the presence of a congenital diastasis of sternal manubrium (37 mm) with agenesis of the corpus sternum, with V-shaped cartilages (24 mm cranio-caudal diameter).
result (Figure 5). At the follow up after 1 year the patient had a normal looking chest wall with absence of abnormally thoracic excursion and a good integration of the extracellular matrix with surrounding tissues as shown to chest X-ray and CT results (Figure 6 a-b).

**Ethical approval:** The research related to human use has been complied with all the relevant national regulations, institutional policies and in accordance the tenets of the Helsinki Declaration, and has been approved by the authors’ institutional review board or equivalent committee.

**Informed consent:** Informed consent has been obtained from all individuals included in this study.

### 3 Discussion

Sternal cleft is a rare congenital anomaly caused by failed midline development and fusion of the mesodermal lateral plates at about 8 weeks of intrauterine life [3]. It is commonly classified in two types: complete, when the sternum is split from the manubrium to the xiphoid, and incomplete (inferior and superior), when a chondral bar bridges the midline [4]. In about 10% of patients with sternal cleft is reported an associated supra umbilical raphe and cardiac anomalies in about 30% [5]. As in our case, the sternal defect can also be a part of a complex clinical condition such as PHACES syndrome. This is a neurocutaneous syndrome of which most common features include posterior fossa malformations, hemangiomas, arterial anomalies, cardiac defects, eye anomalies and sternal clefting or superumbilical raphe [6]. The first case of syndrome was described by Pascual-Castroviejo in 1978 [7], while Frieden et al. used the acronym PHACE in 1996.
later modified in PHACES by Boulinguez et al. [8]. The aetiology is unknown and there is predominance in females [9], rising the hypothesis of dominant X-linked condition. Diagnostic criteria are the presence of hemangioma and the coexistence of at least one among associated anomalies; 70 % of affected children have only one of extracutaneous manifestation [1]. Only 44 previously reported cases of sternal cleft associated with hemangiomas and supra umbilical raphe has been described in literature [10]. The possible cardiac and cerebrovascular anomalies associated must be carefully searched and excluded before any surgical procedures for avoiding major complications. In literature a surgical correction of sternal defect is highly recommendable and should performed as soon as possible during infancy for an higher thoracic compliance at this age [11]. The choice of best surgical approach is controversial and many techniques have been described in literature for reconstructing sternum including primary approximation, sliding or rotating chondrotomies and muscle flaps preferably performed when the flexibility of the chest wall is maximal and compression of the underlying structures is minimal [12]. In some cases the cleft is wide and the sternal remnants hypoplastic, so a primary repair could not be possible. A valid alternative is the employment of prosthetic implantable materials, but there is not consensus about the type to use. Le Roux and Shama reported the ideal characteristics of prosthetic material: rigidity to abolish paradoxical movement with a minimal compression of thoracic organs, inertness to allow in-growth of fibrous tissue and decrease the likelihood of infection, malleability and radiolucency [13]. In our case, we have chosen to use XCM Biologic tissue matrix®, a sterile non-cross-linked 3-D matrix derived from porcine dermis which is composed of cells and extracellular matrix (combination of proteins, proteoglycans, glycosaminoglycans and other biological materials) submitting to procedures of sterilization, decellularization and inactivation of viruses. The result is a strong biologic implant providing a structure that can be infiltrated by body’s cells for a better integration with surrounding tissues and gives a good resistance and durability [14]. The advantage of decellularized dermis is that is gradually revascularized and remodeled into autologous tissue while maintaining its structural integrity. Even if more research is needed to elucidate the indications, contraindications and outcomes of these materials we promote the use of extracellular matrix in surgical reconstruction of chest defects for its maneuverability, plasticity, tolerability and the possibility of growing with the children’s chest getting a good compliance and optimal cosmetic results.

Conflict of interest statement: Authors state no conflict of interest.

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