Case Report

Congenital dermoid fistulas of the anterior chest region (CDFACR): usefulness of sonography for complete resection

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A B S T R A C T

Congenital dermoid fistulas of the anterior chest regions (CDFACRs) consist of a skin orifice at the anterior border of the sternocleidomastoid muscle with fistulas extending caudally in the subcutaneous tissue near the sternoclavicular joint. We report 2 pediatric CDFACR cases with pathognomonic sonography findings. By using sonography, we could diagnose the fistulas as CDFACRs by focusing on their location and direction and could reveal the distal side for complete resection. We suggest that sonography, which does not involve radiation or require sedation, is a better choice for the initial examination of CDFACRs than computed tomography or MRI.

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Introduction

Congenital dermoid fistulas of the anterior chest regions (CDFACRs) consist of a skin orifice at the anterior border of the sternocleidomastoid muscle with fistulas extending caudally in the subcutaneous tissue near the sternoclavicular joint [1–7]. Histologically, CDFACRs have a keratin-containing tubular structure lined with keratinizing stratified squamous epithelium [1–7] and are perhaps connected to the sternoclavicular joint via fibrous tissue [1]. They have also been termed congenital fistulas of the sternoclavicular joint area [2], congenital cutaneous fistulas at the sternoclavicular joint [6], and congenital sternoclavicular dermoid sinuses [7].

The prevalence of CDFACR cases is unclear, and only a few case reports have described their physical diagnosis [1–8] or

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presented computed tomography (CT) images [3,8], MRIs [1], or fistulagrams [7]. Because CDFACRs are congenital and occur in infants and children, radiation exposure should be minimal. Complete resection is required to prevent recurrence [2,7], and preoperative radiologic images of the caudal part of the fistula are important. To the best of our knowledge, sonography findings for CDFACRs have not been reported thus far. Herein, we describe 2 pediatric CDFACR cases and the pathognomonic sonography findings that confirmed their diagnosis.

Case report

The clinical details of 2 CDFACR cases are presented below.

Case 1

An 11-month-old girl, who had a cutaneous pit at the slightly superior to the left sternoclavicular joint at birth, was referred to our institution for a detailed examination. Physical examination detected a dermal pit 1 mm in diameter (Fig. 1A). Sonography without sedation revealed a fistula extending caudally from the pit toward the left sternoclavicular joint (Figs 1B and C). Surgical and histological examination showed that the fistula was lined with squamous or ciliated epithelium with a blind end in the subcutaneous tissue. These findings confirm a diagnosis of a CDFACR with no evidence of malignancy.

Case 2

A 4-year-old girl was referred to our hospital for a detailed examination owing to slight leakage of an effusion from a cutaneous pit that was detected at the slightly superior to the left sternoclavicular pit joint when she was an infant. Physical examination detected a pit 1 mm in diameter. Sonography without sedation showed a fistula extending caudally from the dermal pit toward the left sternoclavicular joint (Figs 2A and B). MRI performed without sedation revealed a well-circumscribed mass in the subcutaneous tissue in the sternoclavicular joint area (Figs 2C–F). Surgical and histological examination showed that the fistula was lined with squamous or ciliated epithelium (Fig. 2G) with a blind end in the subcutaneous tissue. The skin orifice was situated at the anterior border of the sternocleidomastoid muscle and in line with the clavicular bone. These findings confirm a diagnosis of a CDFACR with no evidence of malignancy.

Discussion

Although CDFACRs are thought to be congenital, their etiology is unclear. There are 2 hypotheses. First, as suggested by Araki et al [2], CDFACRs result from disorderly fusions between the sternum and clavicle. The sternum arises from 2 bands of somatopleural mesenchyme near the median plane of the thorax about 6 weeks after fertilization, when the blastemal clavicles are appearing [9,10]. The proximal portion of the clavicles fuses to the upper part of the sternum, specifically, the interclavicular blastema [10]. In some cases, the distal region of the CDFACR extends and, via fibrous tissue, connects to the sternoclavicular joint [2]. In the present case, sonography showed that the caudal region of the CDFACR was also connected to the sternoclavicular joint. Second, as proposed by Ohno et al [6], CDFACRs originate from remnants of the fourth branchial cleft. Moreover, Liston [11] and Moore [9] suggest that the dermal pit of the fistula in the fourth branchial cleft is located near the anterior border of the sternoclavicular joint. Ohno et al [6] reported CDFACR tended to located left side as location of the pyriform sinus fistula. Our sonography findings, which showed the CDFARC on the left side of the chest and extending into the sternoclavicular joint, did not allow us to distinguish between the 2 hypotheses.

Differential diagnoses of a DFACR include a congenital dermoid cyst [12] and a congenital skin fistula with sternal cleft [13] as well as either a lateral cervical sinus/fistula or a pyriform sinus fistula [6]. However, using sonography, we can
distinguish CDFACRs from these other fistula types. First, CDFACRs usually occur on the left side, rather than in the median portion, of the chest [2,6,7]. Congenital skin fistulas with sternal cleft occur in the midline of the anterior chest wall [13], and congenital dermoid cysts occur at the mid-ventral or mid-dorsal embryogenic fusion site [12]. However, similar to CDFACRs, dermoid cysts are thought to result from fusion of the 2 bands of the somatopleural mesenchyme of the sternum [13]. Second, the direction in which a fistula extends is useful for differentiating a CDFACR from a branchial sinus/fistula, as CDFACRs extend caudally. A lateral cervical sinus, also known as a second pharyngeal or branchial cleft fistula/sinus, arises from the remnants of the second branchial cleft, and the sinus or fistula is directed toward a region between the internal and external carotid arteries [14]. A pyriform sinus fistula originates from the remnants of the fourth branchial cleft, with the fistula extending toward the pyriform sinus [15].

Previous studies have reported CT and MRI findings for CDFACRs. CT and MRI allow visualization of the anterior sternal region including sternal anomalies [8,13] and well-circumscribed masses near the sternoclavicular joint [1], respectively. However, for visualization of cutaneous regions, sonography provides higher resolution images compared with CT or MRI. Unlike CT or MRI, sonographic examination of congenital anomalies does not involve radiation or require sedation, which carry small risks to infants and children. Furthermore, sonography provides the caudal side of CDFACRs, and it is useful to perform complete resection and prevent recurrence. For these reasons, sonography may be a better choice for the initial examinations evaluating CDFACRs than CT or MRI.

Conclusion

Sonography allowed visualization of CDFACRs extending to the sternoclavicular joint on the left side of the sternum, thus facilitating their definitive diagnosis. Detection the caudal side of CDFACRs is useful to perform complete resection. Because it does not involve radiation or require sedation, sonography is a viable option for identifying CDFACRs in infants and children.

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