thosis, extended rete ridges, and various degrees of hyperkeratosis in the epidermis. The origins of the mucin and the mechanism of its development are not known, but previous studies suggested that mucin formation might increase as a result of fibroblast upregulation. Because the present case showed a decrease in fat composition and an increase in fibrosis in the NLCS lesion over time, it is considered to be an interesting case supporting this correlation. Apart from fibroblast upregulation, external stimulation by constant friction, and coincidental co-occurrence of mucinous nevus and NLCS should be regarded as possible hypotheses, and more cases and study will be needed to better understand the relation between two diseases and mechanism of mucin deposition.

The clinical features of the present case were similar to those of soft fibromas or neurofibromas, so histopathological examination and special staining were required for differential diagnosis. In conclusion, we report a rare case of acquired mucinous nevus in NLCS. Our case provides additional support for the hypothesis that mucin deposition may develop in association with fibroblast.

CONFLICTS OF INTEREST

The authors have nothing to disclose.

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A Familial Case of Aplasia Cutis Congenita in Two Korean Siblings: A Review of Genetic Aspects

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Dear Editor:

Aplasia cutis congenita (ACC) is a rare congenital malformation, characterized by a localized defect of epidermis, dermis, and sometimes subcutaneous tissues. Although it has been suggested that ACC results from disrupted development or degeneration of skin in utero, the pathogenesis remains unclear. There are five familial cases of ACC reported in Korea (Table 1). Three cases were type

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Table 1. Summary of reported cases of familial ACC in Korea

| Author          | Sex | Location   | Family relationship   | Hereditary nature | Frieden’s classification | Associated abnormalities |
|-----------------|-----|------------|-----------------------|-------------------|--------------------------|-------------------------|
| Kim et al.      | M   | Right leg  | Mother (right leg)    | AD or AR          | Type VI                  | Epidermolysis bullosa   |
| Kim et al.      | M   | Scalp      | Brother (scalp)       | AR or mosaicism   | Type II                  | Adams-Oliver syndrome   |
| Lim et al.      | F   | Left leg   | Sister (both legs)    | AR or mosaicism   | Type VII                 | None                    |
| Kim et al.      | F   | Right foot | 2nd-degree relative   | AR or mosaicism   | Type VII                 | None                    |
| Jin et al.      | F   | Right leg  | Sister (left leg)     | AR or mosaicism   | Type VII                 | None                    |
| Present case    | M   | Scalp      | Brother (scalp)       | AR or mosaicism   | Type I                   | None                    |

ACC: aplasia cutis congenita, M: male, F: female, AD: autosomal dominant, AR: autosomal recessive.

Fig. 1. (A) Clinical manifestation of irregular crusted skin defects at one month after birth. (B) Clinical features with non-hairy large shiny scar at 18 months after birth. (C) His 8-year-old brother also presented with scalp aplasia cutis congenita (ACC). (D) Histopathological findings revealed dermal fibrosis with absence of adnexal structure, consistent with ACC (H&E, ×200).

VII ACC localized to legs1–3. The other cases were combined with syndromic conditions such as epidermolysis bullosa and Adams-Oliver syndrome4,5. Herein, we report 18-month-old male and his 8-year-old brother with ACC on their scalps, which represent type I ACC in siblings without associated congenital anomalies (Fig. 1A–C).

A 13-day-old infant was transferred to us for evaluation of a large ulcerated-crusted scalp defect, which was present since birth (Fig. 1A). The infant was born at term (38 weeks, birth weight 2,580 g) by caesarean section. The newborn presented with a well-demarcated irregular ulcerated scalp defect covering an area 10×10 cm. Outside brain computed tomography and neurosonography showed no skull defects or abnormalities. There was no chromosomal abnormality such as trisomy 13 or 4p-syndrome. Clinical diagnosis was ACC, and he was treated conservatively by simple dressing with antibiotic ointment. After 18 months, he returned to discuss advanced treatment (Fig. 1B). We referred him to plastic surgeon for staged repair. A histological study confirmed the diagnosis of ACC, and there were no remnant viable hair follicles on the biopsy specimen (Fig. 1D).

ACC is a rare congenital disorder with an underlying mechanism that remains unclear. It is characterized by localized or widespread areas of affected skin most commonly on the scalp, and usually present at birth3. ACC may be associated with defects of the underlying skull, especially when the skin defect is larger than 10 cm2. Our patient had no underlying skull abnormalities, although he had a relatively large scalp lesion. According to Frieden’s classification of ACC, type I (ACC of the scalp without multiple abnormalities) is most common with autosomal dominant or sporadic inheritance1. In our case, the parents had no evidence of ACC, indicating the inheritance pattern could be autosomal recessive or represent genetic mosaicism, both of which differ from the typical hereditary pattern of type I ACC.

In conclusion, we report a rare case of familial scalp ACC in Korea which follows atypical inheritance patterns. Our case could provide additional support for the diverse hereditary characteristics of type I ACC.

CONFLICTS OF INTEREST

The authors have nothing to disclose.

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