Case Report

Congenital Gingival Granular Cell Tumor: A Case Report

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KEY WORDS
Congenital epulis;
Newman tumor;
Oral mucosa;

Introduction
The congenital gingival granular cell tumor (CGCT), also as known as congenital epulis, is an unusual benign oral mucosal lesion in newborns. A two-day-old female patient was admitted to the Department of Pediatric Dentistry at Gulhane Medical Academy, Ankara, Turkey with her family, and an intraoral examination showed a CGCT located in the buccal region of the maxillary right first primary molar. In this report, we present a case of CGCT in a newborn.

ABSTRACT
The congenital gingival granular cell tumor (CGCT), also as known as congenital epulis, is an unusual benign oral mucosal lesion in newborns. A two-day-old female patient was admitted to the Department of Pediatric Dentistry at Gulhane Medical Academy, Ankara, Turkey with her family, and an intraoral examination showed a CGCT located in the buccal region of the maxillary right first primary molar. In this report, we present a case of CGCT in a newborn.

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Case Report
A two-day-old female patient was admitted to the Department of Pediatric Dentistry at Gulhane Medical Academy because of swelling of the gums in her upper jaw. She had no medical problems and had been born by natural childbirth. Intraoral examination revealed a polypoid lesion (diameter 0.4 to 0.7 cm) in the area of the maxillary right first primary molar (Figure 1). The tumor had a smooth surface, was pedunculated, and was the same color as the surrounding oral mucosa. No abnormality was detected in neighboring tissue.

Figure 1: Initial appearance of the tumor before surgical excision.

Following consultation with the maxillofacial surgery department, surgical excision under general anesthesia was planned to solve the infant’s feeding problems. The operation was performed on the 12th postnatal day and a biopsy specimen was sent for histopathological examination. S100 antibody was used in the immuno-histochemical investigation. The histopathological examination of the tumor confirmed that it was a CGCT (Figure 2). Wound healing was observed at the two-week follow up, and there was no abnor-
maliy in the area of the surgery (Figure 3).

Figure 3: Healing area after two weeks.

Discussion
CGCTs, initially defined as "congenital epulis" by Ernst Christian Neumann in 1871, are rare benign mesenchymal tumors. [7] It is reported that these tumors are found three times more frequently in the maxillary anterior region than in the mandibular region. CGCTs often originate from the gingiva of the anterior maxillary alveolar ridge. [1, 8-9]

CGCTs are seen most frequently in newborn females (F/M 8:1) as a result of the stimulation of the intrauterine endogenous hormone. [5, 8] The tumor is often single but it has been reported to be multiple in 5 to 16% of cases. Multiple tumors appear to occur mainly in the maxilla or mandible and have rarely been reported in the tongue. The diameter of the tumor can range from 0.1 to 4 cm. The diameter of the largest tumor reported was 7.5 cm. [9-10]

GCTCs, in the clinical differential diagnosis, include teratomas, congenital dermoid cysts, congenital fibrosarcomas, hemangiomas, lymphangiomas, leiomyomas, rabdomyomas, heterotopic gastrointestinal cysts, congenital cystic choristomas, and congenital lipomas. CGCTs appear as normal-colored mucosa, while hemangiomas and lymphangiomas have a red or dark blue area that is spongy on palpation. Findings from palpation for fluctuation and Doppler ultrasound may be used for definitive diagnosis of CGCTs and cystic lesions. Radiological examination for GCTCs includes a margin of soft tissue that is not related to hard tissue. This is done mainly to differentiate GCGTs from solid tissue tumors, especially to eliminate the suspicion of malignancy. [11]

In terms of histology, GCGTs look like adult granular cell tumors. The tumor cells are polygonal with large granular cytoplasm and small eccentric nuclei. There is no mitosis and necrosis, but there is a network of capillaries among the tumor cells. Pseudoepitheliomatous hyperplasia is not found in CGCTs overlying the multilayered squamous epithelium. It is thought that this tumor has undifferentiated mesenchymal cells, fibroblasts, myofibroblasts, histiocytes, Schwann cells or odontogenic epithelial progenitor cells. [12]

When the tumor increases in size, it can cause feeding and respiration problems. Therefore, surgical excision is required without delay. No recurrences have been observed in cases where a CGCT has been completely removed. [5, 13] In our case, the tumor was removed completely.

Conclusion
CGCTs are rare benign tumors, and surgical resection is required to prevent feeding and respiration problems. Histopathologic examination is a critical component in the diagnostic process.
Conflict of Interest
None to declare.

References