Congenital Epulis

Lakshmi Agarwal1 Rajendra Lalani2 Manmohan Agrawal3

1Department of Pathology, GMC, Kota, Rajasthan, India
2Department of Pathology, Dr. Rajendra Lalani Lab, Rajkot, Gujarat, India
3Department of Onco-surgery, Pushpadi Cancer Care Centre, Kota, Rajasthan, India

Address for correspondence Lakshmi Agarwal, MD, Department of Pathology, GMC, Kota, Rajasthan 324010, India (e-mail: drlaxmiagarwal@gmail.com).

Congenital epulis is a benign tumor that occurs in newborns. It was first described by Neumann in 1871; hence, it came to be known as Neumann’s tumor. It is also known as granular cell rhabdomyoma, congenital myoblastoma, or congenital granular cell tumor.1 The female infant is usually affected but has no familial tendency. It not only arises from alveolar median ridge of the maxilla and mandible but can also originate from tongue, palate, skin, etc. Therefore, the etiology and cell of origin of this tumor are still not clear.1,2 It is hypothesized that this tumor arises from several sources: undifferentiated mesenchymal cells, odontogenic epithelial cells, neuroendocrine progenitor cells, pericytes, histiocytes, fibroblasts, or myofibroblast. The steroidal hormones were believed to play a role but this was disproved due to the absence of estrogen and progesterone receptors in the tumor tissue.3

A newborn female presented with a large protruding mass in the mouth causing difficulty in breathing and breastfeeding (►Fig. 1). A probable diagnosis of congenital epulis was made based on clinical examination. Surgical excision of the mass was performed under general anesthesia (►Fig. 2). Postoperative period was uneventful. The specimen was sent for histopathological examination. On gross examination, it consists of single mass covered by gray-brown, firm mucosa with the dimensions of 2.8 × 2 × 1 cm3 (►Fig. 3). The outer surface was smooth. The cut section showed whitish and hemorrhagic areas. The nearest resected margin was found to be 0.8 cm.

Microscopy revealed large round-to-oval cells with granular eosinophilic cytoplasm and small eccentric, bland nuclei. There was presence of thin fibrovascular network separating the cells. Necrosis and mitosis were not evident. The overlying epithelium was thin and pseudoepitheliomatous hyperplasia was not evident (►Fig. 4). All the resected margins were not involved by tumor. On the basis of these findings, the final diagnosis of congenital granular cell tumor was made. Immunohistochemistry (IHC) was not performed as the microscopy findings were characteristic.

It is a rare non-neoplastic soft tissue lesion. Prenatal diagnosis can be done by ultrasonography mainly in the last week of pregnancy.4 This can help in counseling the parents about the possible complications and treatment of the tumor mass. It should be removed postnatally if the size of the mass is big as it can cause difficulty in breathing and feeding.2,4

Surgical excision is the only treatment, though spontaneous regression has also been observed in a few cases. It does not have malignant potential and recurrence rate.4 It presents as single polypoidal protruding mass covered by smooth mucosal surface and firmly attached to the gum by a broad base. The histopathology findings are characteristic. On IHC, the cells are positive for vimentin and negative for S-100 unlike in adults.1,5

It is important to diagnose it both prenatally and postnatally. This will help the oncologist involved in planning the possible intervention and thereby decreasing the morbidity and mortality. The parents can also understand the nature of the mass by consulting the oncologist.

Conflict of Interest
None declared.
References