Yolk sac tumor of ear: a case report and literature review of the last 30 years

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Abstract: Extra-gonadal pure yolk sac tumor of the ear is very rare. We report a case of a yolk sac tumor of the ear and review the English literature in PubMed. The initial complaint was a mass protruding out of the external auditory canal. Computed tomography (CT) showed a large irregularly enhancing isodense mass lesion measuring 42*16 mm in the right external auditory canal, the right mastoid process, and extending to the right back parapharyngeal space with unclear border. Laboratory studies revealed that serum alpha fetoprotein (AFP) was significantly elevated at 664.60 ng/ml (range, 0 to 25 ng/ml), and neuron-specific enolase (NSE) was 28.3 ng/ml (range, 0 to 16.3 ng/ml). After finishing 6 cycles of chemotherapy, the patient underwent a total resection of yolk sac tumor of the ear. In addition, we review the English literature of the yolk sac tumor of the ear.

Keywords: Yolk sac tumor, ear, external auditory canal

Yolk sac tumors, also known as endodermal sinus tumors are one of the common malignant germ cell tumors which account for 3% of all childhood malignant tumors. Yolk sac tumors often arise in gonadal sites such as ovary and testis. Yolk sac tumors of the extra-gonadal region are rare. It has been reported that extra-gonadal occurrence of Yolk sac tumors are involved in the spinal cord [1], urachal [2], abdominal wall [3], mediastinum [4], vagina [5], gluteus [6], the brain [7, 8], the liver [9], head and neck region [10], etc. Yolk sac tumor of the ear is rare [11-13]. To our knowledge, there are only 9 cases reported about the yolk sac tumor of the ear. We aimed to report another case of primary yolk sac tumor of external auditory canal and review the previous cases in the literature.

Case report

A 18-month-year old male child presented to the Department of Otorhinolaryngology, The First Affiliated Hospital of Sun Yat-sen University. His parents found a mass in his right side of the external auditory canal approximately measuring 5*3 mm.

One month later, the mass growth to 30*20 mm with a little bleeding. The parents took the patient to the Department of Otorhinolaryngology, Sun Yat-sen Memorial Hospital of Sun Yat-sen University. Endoscopic examination revealed a dark red mass blocked and protruding out of the right external auditory canal. Laboratory studies revealed that serum alpha fetoprotein (AFP) was significantly elevated at 664.60 ng/ml (range, 0 to 25 ng/ml), and neuron-specific enolase (NSE) was 28.3 ng/ml (range, 0 to 16.3 ng/ml). Other tumor biomarkers including b-HCG, carcinoembryonic antigen (CA125, CA15-3, CA19-9, CA211, CA72-4), and other serum examination were normal. Temporal bone computed tomography scan showed a large irregularly enhancing isodense mass lesion measuring 42*16 mm in the right external auditory canal, the right mastoid process, and extending to the right back parapharyngeal space with unclear border. There were several enlarged lymph nodes in the II level of the bilateral neck region. Biopsy of the mass was then performed.

Microscopically, the tumor was located underneath the epidermis with partially ulceration.
The loose meshwork of interconnected spaces and small cysts were lined by round or ovarian tumor cells with primitive appearance. The papillary fibrovascular core lined by a single layer of tumor cells, which pathognomonic Schiller-Duval bodies were poorly formed. Hyaline globules could be found in the stroma (Figure 1). Immunohistochemistry staining showed that tumor cells were positive for cytokeratin, SALL4, glypican-3, focal positive for EMA, vimentin, CD10, and CD34, and negative for AFP, HCG, PLAP, p63, SMA, CD30, CD31, CD56, S-100 protein, synaptophysin, chromogranin A, RCC, androgen receptor, and GCDFP-15. According to the histopathological findings and elevated serum AFP level, the pathologist diagnosed as a yolk sac tumor. However, the parents refused the chemotherapy due to the economic condition.

Four months later, with the rapid growth, the tumor reached about 87*66 mm (Figure 2). The patient was admitted to Sun Yat-sen University Cancer Center and Guangzhou Red Cross Hospital for the chemotherapy. Temporal bone computed tomography scan showed a large irregularly enhancing isodense mass lesion measuring 87*66 mm in the right exter-
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Figure 3. Temporal bone computed tomography scan showed a large irregularly enhancing isodense mass lesion measuring 87 × 66 mm in the right external auditory canal, the right mastoid process, and extending to the right back parapharyngeal space with unclear border. (Illustration: Diao Zehong, 14 months, M.0877639; Head examination; Sep/05/2015; Patient’s position: HFS; Examination #: 201511050252; Series: 7.5 mm Stnd (A-C), 7.5 mm Stnd (D-F); GE MEDICAL SYSTEMS Optima; 100 KV, 9 mAs (A), 55 mAs (B-F), Sweep range: 320.00 mm; 100% pixel; Original resolution).

Figure 4. Temporal bone computed tomography scan showed no mass in both the right middle ear and external auditory canal.

The child accepted BEP regimen chemotherapy for 6 cycles, consisting of bleomycin (6.4 mg; first day), VP-16 (42 mg; five days), DDP (8.4 mg; five days). After finishing the 6 cycles of chemotherapy, the tumor dramatically reduced to a small soft mass in the external auditory canal.

The patient presented to our hospital for advanced therapy. Temporal bone computed tomography scan revealed there still remained a soft mass in the right external auditory canal. The patient underwent a total resection:

1. The shape of the mass was irregular and the texture of the mass was harder than the temporal bone.
2. The tumor with a wide base of anterior, superior and posterior bone wall of external acoustic meatus. Otherwise, there was not an obvious boundary between them. The peripheral blood supply was not abundant.
Margins of resection, lymph node, the mass were deemed free of tumor.

Six months after the surgery, temporal bone computed tomography scan showed no mass in both the right middle ear and external auditory canal (Figure 4). The patient was still alive during the 13 months follow up period.

Discussion

Yolk sac tumors of the ear are rare and predominantly occur in early-childhood and pediatric patients. English literature from PubMed was surveyed to found the cases with yolk sac tumor of the ear. There were only 10 cases including our current case have been reported [11-19] (Table 1).

Yolk sac tumors often arise in gonadal sites such as ovary and tetis. Yolk sac tumors of the extra-gonadal region are reported in 20% of the cases. The exact etiology of the extra-gonadal germ cell tumors is still not well understood. Filho provided 4 hypothesis: (1) Detachment of germ cell during migration from the vitelline sac to gonadal crest during embryogenesis and their subsequent malignant transformation. (2) Germ cells present in extragonadal tissues. (3) Pluripotent cells that escape from the influence of primary development. (4) The tumors have a genetic origin arising through somatic cells or germ cell mitoses [20, 21].

Yolk sac tumors, also known as endodermal sinus tumors, account for 3% of all childhood malignant tumors. Only 5% of germ cell tumors are found in the head and neck region (except for the central nervous system). There are a few cases reported in the head and neck including ear [11], nasal cavity and sinus [20, 21], nasopharynx [22], orbit [23], parotid gland [24], and submandibular gland [25]. According to previous studies, yolk sac tumor of the head and neck show a highly aggressive behavior which might be attributed to the difficulty of tumor approach and excision. The poor prognosis is related to incomplete resection and anterior skull base involvement. Therefore, what should we do to monitor the recurrence and metastasis status of extra-gonadal and gonadal yolk sac tumors?

The diagnosis of yolk sac tumors mainly depends on the histological findings and serum biomarkers. Both the yolk sac tumor derived from gonad and extra-gonadal regions have similar characteristics, such as specific growth patterns and AFP immunoreactivity. Yolk sac tumor cells display vacuolated cytoplasm, a single large nucleolus, and some small eosinophilic hyaline globules. The tumor cells com-

Table 1. Report of cases of yolk sac tumor of the ear

<table>
<thead>
<tr>
<th>No.</th>
<th>Sex</th>
<th>Age (months)</th>
<th>Position</th>
<th>Serum α-feto protein level</th>
<th>Treatment (months)</th>
<th>Follow up</th>
<th>References</th>
</tr>
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<tbody>
<tr>
<td>1</td>
<td>Female</td>
<td>11</td>
<td>Left external auditory canal</td>
<td>Elevated</td>
<td>Partial resection Chemotherapy 15 m, alive</td>
<td>[14]</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>Female</td>
<td>21</td>
<td>Left temporal bone</td>
<td>Elevated</td>
<td>Surgical resection Chemotherapy 16 m, alive</td>
<td>[15]</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>Female</td>
<td>8</td>
<td>Left external auditory canal</td>
<td>Elevated</td>
<td>Surgical resection Chemotherapy 13 m, alive</td>
<td>[16]</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>Female</td>
<td>26</td>
<td>Left external auditory canal</td>
<td>Elevated</td>
<td>Surgical removal              3 m, dead</td>
<td>[17]</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>Female</td>
<td>18</td>
<td>Left temporal bone</td>
<td>Elevated</td>
<td>Surgical resection Chemotherapy 36 m, alive</td>
<td>[18]</td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>Male</td>
<td>60</td>
<td>Left external auditory canal</td>
<td>Elevated</td>
<td>Surgical resection Chemotherapy -</td>
<td>[16]</td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>Female</td>
<td>24</td>
<td>Left postauricular region</td>
<td>Elevated</td>
<td>Surgical resection Chemotherapy 9 m, alive</td>
<td>[13]</td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>Male</td>
<td>30</td>
<td>Left temporal bone</td>
<td>Elevated</td>
<td>Surgical resection Chemotherapy 9 m, alive</td>
<td>[12]</td>
<td></td>
</tr>
<tr>
<td>9</td>
<td>Male</td>
<td>9</td>
<td>Right external auditory canal</td>
<td>Elevated</td>
<td>Partial resection          3 m, alive</td>
<td>[11]</td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>Male</td>
<td>18</td>
<td>Right external auditory canal</td>
<td>Elevated</td>
<td>Surgical resection Chemotherapy 13 m, alive</td>
<td>Our case</td>
<td></td>
</tr>
</tbody>
</table>
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monly form a meshwork of microcystic, labyrinthine loose, eosinophilic, reticular stroma [26]. In addition, almost all the patients with yolk sac tumors are reported with serum elevated AFP level. There are different patterns under the microscopy, such as pseudopapillary, polycystic, glandular, hepatoid, and solid patterns [27, 28]. In our case, we observed polycystic patterns and Schiller-Duval body which has a characteristic pseudopapillary pattern.

Conclusion

Current therapies for yolk sac tumors are chemotherapy and surgical resection. Extra-gonadal yolk sac tumors have a strong relationship to poor prognosis, owing to the tumor position and total resection of the tumor. However, all the reported cases in the literature were still alive during the follow-up period. In our case, the child received chemotherapy and subsequent surgical resection, and was alive at the time of this writing. Therefore, we can closely test the serum level of AFP to monitor the status of yolk sac tumors.

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Disclosure of conflict of interest

None.

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References


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