Case Report / Приказ болесника

Milan Filipović¹,², Tatjana Filipović¹, Nebojša Videnović¹,², Zlatan Elek¹,³, Aleksandar Božović¹,³,*

A very rare cutaneous epithelioid hemangioendothelioma in the right auricle on a male adult

Веома редак кожи епителоидни хемангіоендоцителом десне ушне школке код одраслог човека

¹University of Priština – Kosovska Mitrovica, Faculty of Medicine, Kosovska Mitrovica, Serbia;
²Priština Clinical Hospital Centar – Gračanica, Gračanica, Serbia;
³Kosovska Mitrovica Clinical Hospital Centar, Kosovska Mitrovica, Serbia

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*Correspondence to:
Aleksandar BOŽOVIĆ
University of Priština – Kosovska Mitrovica, Faculty of Medicine, Department of Surgery, Anri Dinana b.b., 38220 Kosovska Mitrovica, Serbia
E-mail: aleksandar.bozovic@med.pr.ac.rs
A very rare cutaneous epithelioid hemangioendothelioma in the right auricle on a male adult

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Main treatment for local changes is surgical removal. Chemot

SUMMARY
Introduction Epithelioid hemangioendothelioma is an extremely rare tumor of vascular origin, which mainly affects soft tissues, and in rare cases, also affects the skin.
Case outline A 71-year-old man came for an examination due to a tumor localized on the antihelix of the right ear. A radical excision of the tumor and reconstruction of the defect with a preauricular transposition tunnelized Banner flap was performed.
Conclusion Epithelial hemangioendothelioma of the skin is an extremely rare, vascular tumor. It is characterized by local aggression, the appearance of local recurrences and metastatic hematogenous and lymphogenetic spread. Standard dermoscopic examination of this tumor is not important. The appearance of the tumor itself has no clinical specificity, and most often resembles nodular basal cell carcinoma. An early visit to the doctor and removing the change early are the key to success in treating this tumor. So far, no case has been reported that is localized on the antihelix of the ear. Radical surgical excision is the therapy of choice.
Keywords: epithelioid hemangioendothelioma; vascular neoplasm; auricle

SAŽETAK
Увод Епителоидни хемангиоендотелиом је изузетно редак тумор васкуларног порекла, који углавном захвата мека ткива, а у ретким случајевима, захвата и кожу.
Приказ болесника Приказани је мућкац старости 71 године, који долази на преглед због туморске промене локализоване на антхеликсу десне ушне школке. Болеснику је урађена радикална екцизција тумора и реконструкција транспозиционим тунелизованим Банеровим флагом.
Закључак Епителоидни хемангиоендотелиом коже представља изузетно редак, тумор васкуларног порекла. Карактеристична је локалном агресивношћу, појавом локалних рецидива и метастатским хематогеним и лимфогеним ширањем. Стандардни дермоскопски преглед код овог тумора није од велике значаје. Изглед саме туморске промене клинички нема специфичности, а најчешће подсећа на нодуларни базалноклетачки карцином. Рани одлазак код лекара и рано уклањање промене су кључ успеха у лечењу овог тумора. До сада није објављен ниједан случај који је локализован на антхеликсу ушне школке. Радикална хируршка екцизција је терапија избора.
Кључне речи: епителоидни хемангиоендотелиом; васкуларна неоплазма; ушна школка

INTRODUCTION
Epithelioid hemangioendothelioma (EHE) is a vascular tumor of endothelial cell origin. EHE is extremely rare, with an incidence of 1: 1,000,000 [1], and the literature is limited to case reports and several retrospective studies. It most commonly occurs in the lungs, liver, and bones, but can also occur in the skin [2]. EHE was first described by Weiss and Enzinger in 1982. as a soft tissue tumor [3]. Clinically, the neoplasm usually presents as a slightly raised, erythematous, sometimes painful dermal nodule.

According to the World Health Organization classification from 2020. (WHO 2020), EHE is a tumor that is locally aggressive with metastatic potential [4]. The rate of local recurrence of epithelioid hemangioendothelioma is 10 to 15%, and the level of lymphatic and systemic metastases is between 20 and 30% [5]. Pathohistological analysis is the gold standard for definitive diagnosis, where tumor cells are CD31, CD34, FLI-1 and ERG positive [1]. The main treatment for local changes is surgical removal. Chemotherapy and radiotherapy have not
been shown to be effective due to slow tumor growth [6]. The prognosis of surgically treated local changes is good, with a five-year survival rate of 75.3% [7].

**CASE REPORT**

This case demonstrates a rare neoplasm which unusually presented as a cutaneous lesion. The location of the lesion on the ear is a challenging area for the surgeon to reconstruct. Local flaps, preauricular transposition tunnelized flap in this case, provide adequate reconstruction and are cosmetically acceptable. EHE requires aggressive management, given its potential to metastasize to lymph nodes, and it requires continued surveillance after treatment. It is important to consider alternative histological diagnoses in patients presenting with cutaneous lesions and how this affects management and prognosis.

A seventy-one-year-old patient was admitted to the surgery department of the Clinical Hospital Centar Priština - Gračanica, due to a tumor localized on the antihelix of the right ear. On physical examination, in the area of the antihelix of the right ear, there is a nodule measuring 0.5 × 0.5 cm, above the level of the skin, with clearly defined edges, firm consistency, immobile in relation to the surface, Figure 1.

The anamnestic patient states that he noticed the change four months before and that it increased rapidly. Upon admission to our institution, the patient is in good general condition, with orderly laboratory findings, conscious, oriented, normocardic, eupnoic. Under conditions of local anesthesia was performed a radical excision of the tumor. The reconstruction of the resulting defect was performed using a preauricular transposition flap, (Banner flap) tunnelized through a cartilaginous fistula created at the crus helicis. The flap was deepithelialized at its base to allow the closure in a single step. The donor site was primarily closed. Figure 2.

In the laboratory for pathohistology, pathohistological and immunohistochemical analysis of the tumor change was performed.

Histopathological finding, Figure 3.:

- Separated from the epidermis by a narrow free zone, limited, non-encapsulated tumor nodule built of epithelioid cells, more abundant eosinophilic cytoplasm, round-oval nucleus with noticeable nucleolus and rare mitoses (2–3 mitoses/mm²).

- Scattered multinuclear cells and small foci of lymphocytic infiltrates

- Focal, more pronounced superficial, hyalinized stroma between tumor cells
Immunohistochemical findings:

- PRAME: + (paler coloring)
- CD34: -
- BAP1: +
- ERG: +, Figure 4.
- CD68: - (rare scattered single cells +)
- FLI-1: +
- CD10: diffuse +
- Ki-67: about 20%
- CD31: -

  Maximum tumor thickness: 4.3 mm
  Minimum distance from the resection line in depth: 0.2 mm.

After the early postoperative period, which passed without complications, the patient came for regular check-ups at 1, 3, 6 and 12 months postoperatively, Figure 5.

At each examination, clinical, laboratory and ultrasound evaluation was performed in order to confirm the occurrence of possible metastatic changes. After a one-year follow-up, which is without clinical manifestations of disease recurrence, we will continue the follow-up in the next four-year period.

**DISCUSSION**

Cutaneous epithelioid hemangioendothelioma is most common in adults, both men and women, but there are several cases where cutaneous EHE has been described in children [8]. According to the World Health Organization, EHE is a malignant tumor, with a metastasis incidence of 20 to 30% and a mortality of 10 to 20% [4]. Deyrup et al., in 2008, analyzed a series of 49 EHE subjects and concluded that large tumors (> 3 cm in diameter) with high mitotic activity (more than 3 mitoses) have a more aggressive clinical course [9]. In our case, the mitotic activity of the tumor is low, as well as the diameter of the tumor itself, which indicates that it is a tumor with a low risk of local recurrence and systemic metastasis.
According to the literature, the clinical findings of EHE vary from dermal or subcutaneous nodules [10], through non-healing ulcers to small multiple red papules [11].

Pathohistologically, tumor cells with eosinophilic cytoplasm, organized in hyaline or mucoid stroma, are present in EHE [12]. A large number of endothelial proteins may be useful in the diagnosis of EHE. FLI-1 protein shows greater sensitivity and specificity in the diagnosis of EHE than CD-31 and CD-34. CD-34 is positive in more than 90% of vascular tumors, so this marker has low specificity because it occurs in a large number of soft tissue tumors [13]. Based on research in 2000, Miettinen et al showed that the transcription factor ERG is a tumor marker that occurs in 42 of 43 cases of EHE [14], Figure 6. Based on the analysis of Uta F. et al., in 2014, all 39 subjects with EHE tested on ERG were positive, as were all five tested on FLI-1 [15].

EHE is an extremely rare tumor of vascular origin. It is characterized by local aggression, local recurrences and metastatic, hematogenous and lymphogenic spread. Standard dermoscopic examination of this tumor is not important. The appearance of the tumor itself has no clinical specificity, and most often resembles nodular basal cell carcinoma. Going to the doctor early and removing the change early are the key to success in treating this tumor. Radical surgical excision is the therapy of choice. There are various surgical procedures described for the reconstruction of the anterior auricle, including local flaps, skin grafts and even healing by secondary intention. The authors consider the tunnelized preauricular transposition flap a good option especially when compared to interpolated flaps, since it allows a reconstruction in a single surgical step with decreased morbidity and favorable cosmetic results.

Ethical standards: Written consent for publication of this article has been obtained by the patient.

Conflict of interest: None declared.
REFERENCES


Figure 1. Appearance of tumor preoperatively
Figure 2. Appearance at the end of the operation
Figure 3. Histopathological findings at different magnifications show: a – 10 × 0.25; b – 10 × 0.35; c – 10 × 0.40 [non-encapsulated tumor nodule built of epithelioid cells, more abundant eosinophilic cytoplasm, round-oval nucleus with noticeable nucleolus and rare mitoses (2–3 mitoses/mm²), scattered multinuclear cells and small foci of lymphocytic infiltrates, and focal, more pronounced superficial, hyalinized stroma between tumor cells]; and d – 4 × 0.25 (the ratio of tumor tissue to healthy tissue)
Figure 4. Positive nuclear ERG expression in tumor cells (10 × 0.25)
Figure 5. Postoperative appearance after six months