

CASE REPORT

PEER REVIEWED | OPEN ACCESS

Epithelioid hemangioma of the skull: Rare case report

Ayah Rashwan, Chanpreet Riarh, Lauren Tristani, Joseph de Nanassy, Raveena Ramphal, Nagwa Wilson

ABSTRACT

Introduction: Epithelioid hemangiomas (EH) are benign vascular tumors that commonly occur in the skin, subcutis, and bone—most typically arising as nodules in the neck and head region. Pediatric cases with atypical presentations remain rare and challenging.

Case Report: The following outlines the case of a 2-year-old male who presented with a bump and swelling on the left side of his head. Imaging tests identified multiple, hyperdense, and focally ring-enhancing intra-axial lesions located mainly at the gray-white matter junction of bilateral frontal left-sided parietal, left-sided occipital, and left-sided temporal lobes. In addition, multiple lytic lesions were seen in the calvarium. Pathology confirmed EH. While one lesion required surgical resection, the remaining lesions showed spontaneous regression during follow-up.

Conclusion: This case highlights the diagnostic and management challenges associated with pediatric EH involving the head, including intracranial involvement. The patient's unique presentation, age, and lesion

involution expand the understanding of EH beyond its typical manifestation.

Keywords: Epithelioid hemangioma, Pediatric, Skull

How to cite this article

Rashwan A, Riarh C, Tristani L, de Nanassy J, Ramphal R, Wilson N. Epithelioid hemangioma of the skull: Rare case report. J Case Rep Images Pathol 2025;11(1):1–7.

Article ID: 100084Z11AR2025

doi: 10.5348/100084Z11AR2025CR

INTRODUCTION

Epithelioid hemangioma (EH), also known as histiocytoid hemangioma, and angiolymphoid hyperplasia with eosinophilia, is a rare and benign vascular lesion [1–10]. It most often occurs in the head and neck region, within the bone or soft tissue [1, 2, 5–10]. These lesions tend to be small, usually not exceeding 10 cm [10]. Epithelioid hemangioma in children lacks comprehensive knowledge regarding clinical characteristics, outcomes, and consensus on treatment [7]. Complete removal of the lesion, via surgical excision, is the commonly reported treatment, making local recurrence rare [10]. However, treatment involving incomplete resection has a reported recurrent rate of approximately 33% [10].

We present a case of EH in a 2-year-old Asian male in the left frontal bone of the skull. Diagnostic and management challenges are discussed.

CASE REPORT

A previously healthy 2-year-old male presented with pneumonia. His workup revealed a bump on the left side

Ayah Rashwan¹, Chanpreet Riarh², Lauren Tristani³, Joseph de Nanassy⁴, Raveena Ramphal⁵, Nagwa Wilson⁶

Affiliations: ¹Student, Interdisciplinary Faculty of Health Sciences, University of Ottawa, Ottawa, Ontario, Canada;

²Student, Medical Science, Western University, London, Ontario, Canada; ³Clinical Researcher, Department of Pathology, Children's Hospital of Eastern Ontario, Canada;

⁴Pathologist, Department of Pathology, Children's Hospital of Eastern Ontario, Ottawa, Ontario, Canada; ⁵Oncologist, Division of Hematology/Oncology, Children's Hospital of Eastern Ontario, University of Ottawa, Ottawa, Ontario, Canada; ⁶Radiologist, Department of Diagnostic Imaging, Children's Hospital of Eastern Ontario Research Institute, Ottawa, Ontario, Canada.

Corresponding Author: Dr. Nagwa Wilson, 401 Smyth Rd, Ottawa, Ontario K1H 8L1, Canada; Email: NaWilson@cheo.on.ca

Received: 11 November 2024

Accepted: 04 January 2025

Published: 29 January 2025

of his head. He had no associated constitutional symptoms such as weight loss, fever, headaches, or seizures. He underwent imaging, including head and chest computed tomography (CT) and head magnetic resonance imaging (MRI). The latter revealed multiple ring-enhancing intra-axial lesions involving the gray-white matter junction of the bilateral frontal, left-sided parietal, left-sided occipital, and left-sided temporal lobes surrounded by significant vasogenic edema. The foci showed blooming artifact in the susceptibility weighted sequence (SWI) (Figure 1). The calvarium parietal bone showed the presence of multiple lytic intradiploic lesions associated with a soft tissue component (Figure 2). The skeletal survey did not reveal any lytic lesions. Abdominal ultrasound displayed an enlarged liver without focal lesions and no lymphadenopathy. The clinical considerations included inflammatory lesions, such as tuberculosis and fungal infection, and less likely multiple metastases. The patient is doing clinically well during follow-ups, and the MRI demonstrated improvements in intracranial lesions with no new parenchymal findings (Figure 1).

The lesion showed progressive growth during four months of follow-up; mandating a surgical resection of the lytic bony lesion with scalp soft component (Figure 3). Pathology revealed soft tissue and bone involvement with a vascular small- to medium-sized

proliferation lined by endothelial cells with occasional cytoplasmic vacuolization (Figure 4). The intervening stroma showed marked reactive inflammatory cells infiltrating the background, with a predominance of eosinophils (Figure 4). Brisk mitotic activity and necrosis were not identified. Immunohistochemistry showed negative CD1a. The sarcoma panel molecular testing was negative for single-nucleotide variants (SNVs) and translocations. These findings are not commonly associated with malignant lesions, thus ruling out epithelioid hemangioendothelioma and angiosarcoma. After extensive investigation and discussion, the consensus opinion is that the morphological and immunohistochemical features seen in this skull bone lesions specimen are those of an EH. The patient underwent a comprehensive surgical resection aiming to remove the lesion. Interestingly, only one lesion was surgically excised, as the remaining lesions seem to be regressing spontaneously. The patient showed no signs of recurrence during the 12-month follow-up period.

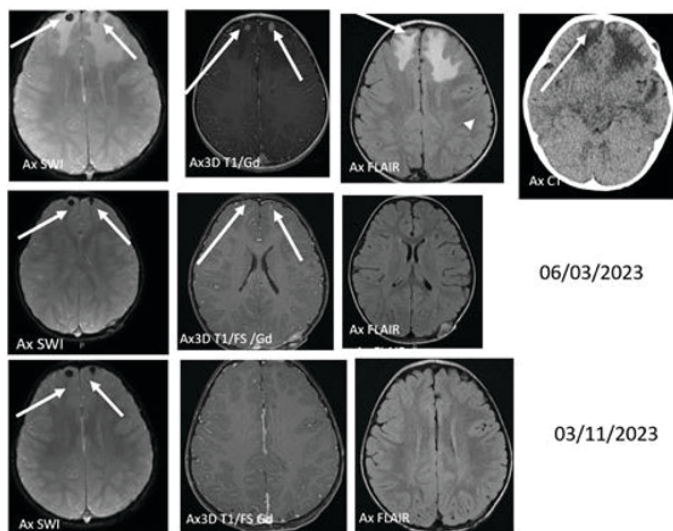


Figure 1: Intra-axial lesions with follow up AX SWI, AX 3DT1/Gd, and AX FLAIR are provided for the initial presentation and axial non-enhanced CT axial shows hyperdense foci in the bifrontal and left temporal lobes surrounded by hypodensity consistent with surrounding vasogenic edema. Follow-up after 5 months and 11 months shows interval resolution of edema on FLAIR sequence, interval decrease in the size, and enhancement of the lesions in the post-Gd sequences, and they are not visualized in the last follow-up; however, there is no change in the corresponding blooming artifact in the SWI sequence at initial presentation until last follow-up.

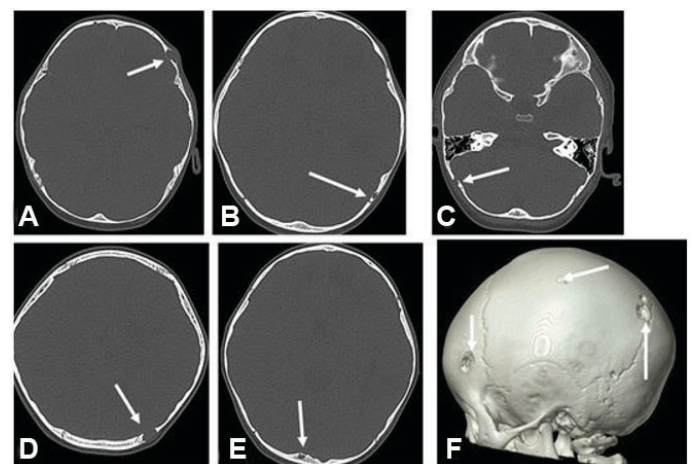


Figure 2: Axial CT in bone window. (A–E) demonstrate the presence of multiple lytic lesions (white arrows); (F) 3D reformat image shows multiple calvaria lesions.

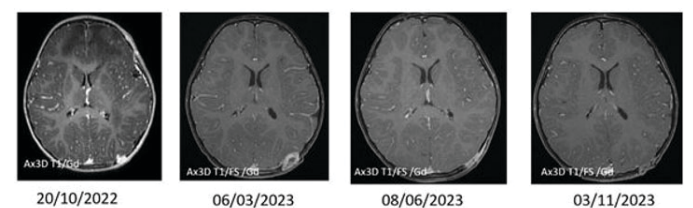


Figure 3: Calvaria lesions follow-up: Initial presentation with multiple calvaria lesions. The provided MRI sequence is 3D/T1/Gd, showing multiple heterogeneously enhancing lesions scattered, one of which is seen in the left posterior parietal bone. It showed interval increase in size in the follow-up performed after five months on 06/03/2023 which was later resected.

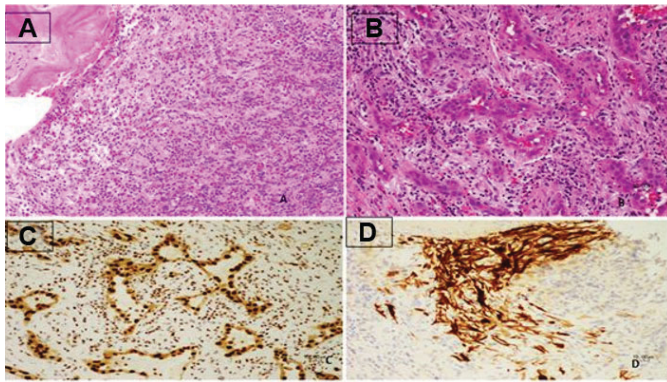


Figure 4: (A) Proliferation of spindled and polygonal myofibroblasts with numerous lymphocytes and eosinophils in the background. Piece of (skull) bone in left upper corner, H&E $\times 200$. (B) Lesion cells strongly positive by immunohistochemistry, Caldesmon $\times 200$. (C) Proliferation of capillary vessels with plump endothelial cells in an inflammatory background, H&E $\times 200$. (D) Strong nuclear staining in prominent epithelioid endothelial cells by immunohistochemistry, ERG $\times 200$.

DISCUSSION

In pediatric patients, EH occurs more frequently in males with a median age of 14.4 years [7]. Among adult populations; however, the literature reports a higher prevalence in women with a median age ranging from 20–50 years old and a mean onset of 30–33 years old [8–18]. Our current case involves a 2-year-old Vietnamese patient

diagnosed with EH. Epithelioid hemangioma is most prevalent among Asian and Caucasian populations, a trend that aligns with our case, and is considered rare in elderly and non-Asian pediatric populations [10, 15, 17, 18].

Epithelioid hemangioma is commonly seen in the head and neck region in both pediatric and adult populations [1, 2, 5–10, 12–22], aligning with our case presentation in the left frontal bone of the skull. However, the location of EH lesions can vary throughout the body [5, 7, 8, 10] (Tables 1 and 2).

Additionally, our patient displayed symptoms of lesion growth and pain, mirroring symptoms reported in other pediatric EH cases [2, 7, 11]. However, the commonly reported symptoms among other cases include pruritus and pain [19]. Imaging revealed multiple ring-enhancing intra-axial lesions located predominantly in the gray-white matter junction of bilateral frontal left-sided parietal, left-sided occipital, left-sided temporal lobes surrounded by significant vasogenic edema. Also, the calvarium showed multiple lytic diploic lesions predominantly on the left side of the calvarium and with soft tissue components. A skeletal survey revealed no other lytic lesions. Abdominal ultrasound showed an enlarged liver with no focal lesion, normal spleen, and no lymphadenopathy. Interestingly within the adult literature, only one case has been reported in a 66-year-old woman with primary EH affecting the central nervous system (CNS) and manifesting in the CNS [22].

Table 1: Pediatric cases of EH

Case/ [Reference]	Age at diagnosis (years)/ gender	Presenting symptoms	Size of lesion	Location side + body part	Diagnostics	Treatment	Follow- up (mo/ yr)
1/[1]	9y/F	Dysphagia	4.8 $\times$ 3.7 $\times$ 3.1 cm	Right side + neck	CT, angiography, ultrasound	Embolization, local excision of mass with nerve monitoring, Extubating post- procedure	6 mo, 1 yr
2/[2]	14y/M	Enlarging painless scrotal mass	3.0 $\times$ 2.5 cm	Scrotal wall	Ultrasonography, ultrasound, scrotal exploration, complete biopsy	Primary scrotal reconstruction, surgical excision	–
3/[4]	15y/M	Severe swelling, tenderness, redness, and warmth	3 $\times$ 212 cm (2nd toe and metatarsal bone); 3 $\times$ 4 $\times$ 8 cm (left inguinofemoral region)	Left foot + 2nd toe. Left + inguinofemoral region	Histological examination, MRI, core needle biopsy, immunohistochemistry	Ray resection, surgery	3 yr, 4 yr
4/[6]	15y/M	Painless penile glans mass	2.5 cm glandular lesion	Dorsal + penile glans	Ultrasound, cystoscopy	Surgical excision, urethroplasty	Initially lost to FU; re- evaluated 18 mo post- resection due to recurrence 2 mo

Table 1: (Continued)

Case/ [Reference]	Age at diagnosis (years)/ gender	Presenting symptoms	Size of lesion	Location side + body part	Diagnostics	Treatment	Follow- up (mo/ yr)
5/[7]	Median age 14.4 y (n = 11)/n = 9M, n = 2F	Localized tenderness, localized pain, CN deficits	1.5–6.4 cm	Face, leg, skull, foot, spine, pelvis	Biopsy	Sirolimus, resection, biopsy, embolization, propranolol, curettage, zoledronate, interferon, cryoablation, partial corpectomy; resection	0.3–7.2 mo
6/[8]	11y/M	Migraines, headaches, syncope, seizures	–	Right-side + orbital mass	CT and MRI, biopsy	Oral propranolol	–
7/[9]	4y/M	Painless penile skin lesions lasting 1 yr	–	Right edge + glans and right distal penile shaft	Immunohistochemistry	Propanol (unsuccessful), excision—skin graft	–
8/[23]	1d/M	Normal pulse, heart rate, and grade 2–3/6 ejection systolic murmur	3.2 cm diameter	Right + atrial tumor	Echocardiography, CT, chest X-ray, electrocardiogram	Tracheal intubation and mechanical ventilation (due to respiratory failure), the patient underwent cardiac surgery on the 9th day of life	–
9/[10]	16y/M	Painless and enlarging scapula mass	15 × 15 cm	Left + scapula	Ultrasound, MRI, and core needle biopsy	Preoperative embolization, surgical excision	6 mo
10/[11]	17y/M	Pain	17 mm × 4 mm × 15 mm	right + wrist	MRI, radiograph	Intralesional curettage surgery	5 mo

Abbreviations: CN, cranial nerve; F, female; M, male.

Table 2: Adult cases of EH

Case/ [Reference]	Age at diagnosis (years)/ gender	Presenting symptoms	Size of lesion	Location side + body part	Diagnostics	Treatment	Follow-up (wk/mo/ yr)
1/[12]	43y/M	Gait disturbance, back pain, lucent mass	–	Vertebral body T3	CT, MRI, biopsy, histopathology	Laminectomy, embolization	12 mo
2/[13]	30y/F	Swelling, pain	1 × 2 cm (lingual alveolar mucosa) and 4–5 mm (labial surface)	Lingual alveolar mucosa and labial surface	Radiography, blood investigation, immunohistochemistry, histopathology	Excisional biopsy	1 mo

Table 2: (Continued)

Case/ [Reference]	Age at diagnosis (years)/ gender	Presenting symptoms	Size of lesion	Location side + body part	Diagnostics	Treatment	Follow-up (wk/mo/ yr)
3/[14]	35/F	Breast lump	2.5 × 2.2 cm	Left + breast	Biopsy, histopathology	Excision	2 yr
4/[19]	53/M	Right epistaxis (recurring)	–	Right + middle turbinate	Cerebral magnetic resonance imaging and CT, histopathology, immunohistochemistry	Endoscopy resection	12 mo
5/[15]	47/M	Painless, slowly growing mass	2.6 cm long	Left + orbit	Hertel exophthalmometry, chest X-ray, CT, MRI, histopathology	Surgical excision, propranolol	16 mo
6/[20]	50/M	Pain	6 × 8 mm	Mid shaft of the penis dorsally	Local excision, immunohistochemistry, histopathology		3 mo
7/[16]	29/F	Multiple small red/pink nodules, severe itching	~1–2 cm ² dimension	Left + external auditory canal skin	Biopsy, histopathology	Steroid cream, excision	6 mo, 9 mo
8/[21]	52/M	Nodular lesion	3 cm	Left + buccal mucosa	Histopathology, immunohistochemistry	Excisional biopsy	18 mo
9/[22]	66/F	Headaches, vision impairment	24 mm	Primary EH of the CNS	MRI, CT, histopathology, immunohistochemistry	Surgical excision	–
10/[17]	51/M	Solitary mass	8 × 6 cm	Right + cheek	Ultrasound, CT, histopathology	Full excision	2 wk, 18 mo
11/[18]	25/F	Painless submucosal nodule	5 mm × 5 mm	Left angle of mouth + lower lip	Complete blood investigation, histopathology	Excisional biopsy	–

Regarding the pediatric population, there is no current treatment consensus for EH due to its rarity [7, 10]. Pediatric cases of EH [1–5, 10], including the presented case, typically present as well-circumscribed lesions, allowing for feasible complete surgical excision, a pattern similarly observed in most adult lesions [13, 14, 17, 19]. In our case, the lesion was dissected freely from the bone—falling in line with the most common method of treatment for EH reported in both pediatric and adult populations [1, 2, 4, 6–8, 10, 14–22]. Surgical excision typically results in low morbidity and has been deemed the most successful treatment [1, 2, 4, 6, 8, 10]. Similarly, the literature reports complete excision with follow-up as the most effective treatment of EH in adults [14–22].

The rarity of EH within the pediatric population has contributed to the difficulty in the characterization of diagnoses, as highlighted in previous literature [3, 4, 7, 10]. Similarly, the adult literature also indicates a lack of therapeutic protocols for EH in the brain due to its infrequent occurrence [22]. Despite these challenges, our case aligns with the treatment and outcome consensus observed in pediatric cases of EH. Successful outcomes and no recurrence are evident post-treatment and follow-up, mirroring findings in the literature [6]. While some cases, both in pediatric and adult populations, have

reported medical treatments involving propranolol, sirolimus, and steroids, the necessity for long-term follow-up is emphasized to ensure successful outcomes in patients undergoing such interventions [7, 8, 14–16].

Our patient experienced numerous back-to-back viral infections following surgical treatment. However, he is doing well with no recurrence of EH during the 12-month follow-up.

CONCLUSION

We report a rare case of EH in a 2-year-old male. Notably, our patient's very young age, multifocal lesion distribution, and unexpected outcome with lesion involution during follow-up distinguish this case. Additionally, the novelty of intracranial involvement expands the understanding of EH beyond its typical presentation limited to the skull and scalp.

REFERENCES

1. Abraham MJ, Gregory ND, Chennupati SK. Epithelioid hemangioma of the internal carotid artery: A case report supporting the reactive pathogenesis

- hypothesis of this vascular tumor. *Int J Pediatr Otorhinolaryngol* 2014;78(7):1186–9.
2. Avallone AN, Avallone MA, Share S, Rubin BP. Epithelioid hemangioma—A rare scrotal tumor of childhood. *Urology* 2012;80(3):707–9.
3. Chakravarthi SS, Veeraraghavan S, Fukui M, Jennings J, Rovin RA, Kassam AB. Diagnostic dilemma: An unusual case of primary intracranial, intraparenchymal epithelioid hemangioma. *J Neurol Stroke* 2018;8(5):263–5.
4. Floris G, Deraedt K, Samson I, Brys P, Sciort R. Epithelioid hemangioma of bone: A potentially metastasizing tumor? *Int J Surg Pathol* 2006;14(1):9–15.
5. Huang SC, Zhang L, Sung YS, et al. Frequent FOS gene rearrangements in epithelioid hemangioma: A molecular study of 58 cases with morphologic reappraisal. *Am J Surg Pathol* 2015;39(10):1313–21.
6. Kohli P, Wang Y, Baker Z, et al. Epithelioid hemangioma of the glans penis in an adolescent male: A case report. *Urology* 2022;170:189–92.
7. Liu KX, Duggan EM, Al-Ibraheemi A, Shaikh R, Adams DM. Characterization of long-term outcomes for pediatric patients with epithelioid hemangioma. *Pediatr Blood Cancer* 2019;66(1):e27451.
8. Moss HB, Sines DT, Blatt J, Dutton JJ, Proia AD. Epithelioid hemangioma responsive to oral propranolol. *Ophthalmic Plast Reconstr Surg* 2012;28(4):e88–90.
9. Murshed K, Farghaly H. Multifocal epithelioid hemangioma of the penis in a 4-year-old child: A case report. *Am J Dermatopathol* 2020;42(5):372–4.
10. Wiggins CJ, Dibbs RP, Bartlett EL, Ashton DJ, Maricevich RS. Atypical presentation and management of an epithelioid hemangioma: A case report and review of the literature. *Ann Pediatr Surg* 2020;16(1):53.
11. Bregman JA, Jordanov MI. Epithelioid hemangioma occurring in the radial styloid of a 17-year-old boy—An unusual presentation of an uncommon neoplasm. *Clin Imaging* 2014;38(6):899–902.
12. Okada E, Matsumoto M, Nishida M, et al. Epithelioid hemangioma of the thoracic spine: A case report and review of the literature. *J Spinal Cord Med* 2019;42(6):800–5.
13. Nangia R, Puri A, Gupta R, Bansal S, Negi A, Chauhan I. Epithelioid hemangioma of lingual alveolar mucosa: An immunohistochemical case report. *Case Rep Med* 2014;2014:436240.
14. Nair M, Aron M, Sharma MC. Angiolymphoid hyperplasia with eosinophilia (epithelioid hemangioma) of the breast: Report of a case. *Surg Today* 2000;30(8):747–9.
15. Lee SM, Khwarg SI, Ahn JH. Epithelioid hemangioma of the orbit treated with oral propranolol in adult. *Can J Ophthalmol* 2018;53(6):e244–6.
16. Hussain JA, Al Zamel H, Nawaz B. Angiolymphoid hyperplasia with eosinophilia (epithelioid hemangioma) of the external auditory canal, an unusual presentation in an adult female: A case report. *J Surg Case Rep* 2018;2018(7):rjy145.
17. Al-Muharrari MA, Faqi MK, Uddin F, Ladak K, Darwish A. Angiolymphoid hyperplasia with eosinophilia (epithelioid hemangioma) of the face: An unusual presentation. *Int J Surg Case Rep* 2011;2(8):258–60.
18. Aggarwal A, Keluskar V. Epithelioid hemangioma (angiolymphoid hyperplasia with eosinophilia) in the oral mucosa. *Indian J Dent Res* 2012;23(2):271–4.
19. Makhoul M, Hourseau M, Nassereddine H, Hamad WA, Khoeir N. Epithelioid hemangioma of the nasal cavity with atypical features: A case report and review of the literature. *SN Compr Clin Med* 2023;5(1):235.
20. Ismail M, Damato S, Freeman A, Nigam R. Epithelioid hemangioma of the penis: Case report and review of literature. *J Med Case Rep* 2011;5:260.
21. Henriques ÁCG, Vidal MTA, Gurgel CA, Rocha SLD, Júnior BC, de Azevedo RA, dos Santos JN. Epithelioid hemangioma in the oral mucosa – A case report. *Srp Arh Celok Lek* 2016;144(9–10):535–40.
22. Covelli C, Parente P, Icolaro N, Dimitri LMC, Vigna B, Popolizio T, Graziano P. Primary epithelioid hemangioma of the central nervous system: A case report and review of the literature. *J Neuropathol Exp Neurol* 2021;80(7):717–9.
23. Malakan Rad E, Radmehr H, Vasei M, Rahimi Rastgoo B. Giant congenital right atrial epithelioid-capillary hemangioma with prolonged QT interval: Case report and practical surgical treatment strategy for primary cardiac tumors in children based on 25-year review of 299 cases. *Echocardiography* 2018;35(9):1471–81.

Author Contributions

Ayah Rashwan – Conception of the work, Design of the work, Acquisition of data, Analysis of data, Interpretation of data, Drafting the work, Revising the work critically for important intellectual content, Final approval of the version to be published, Agree to be accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved

Chanpreet Riarh – Interpretation of data, Revising the work critically for important intellectual content, Final approval of the version to be published, Agree to be accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved

Lauren Tristani – Conception of the work, Design of the work, Acquisition of data, Analysis of data, Interpretation of data, Revising the work critically for important intellectual content, Final approval of the version to be published, Agree to be accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved

Joseph de Nanassy – Interpretation of data, Revising the work critically for important intellectual content, Final approval of the version to be published, Agree to be accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part

of the work are appropriately investigated and resolved

Raveena Ramphal – Interpretation of data, Revising the work critically for important intellectual content, Final approval of the version to be published, Agree to be accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved

Nagwa Wilson – Acquisition of data, Interpretation of data, Revising the work critically for important intellectual content, Final approval of the version to be published, Agree to be accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved

Guarantor of Submission

The corresponding author is the guarantor of submission.

Source of Support

None.

Consent Statement

Written informed consent was obtained from the patient for publication of this article.

Conflict of Interest

Authors declare no conflict of interest.

Data Availability

All relevant data are within the paper and its Supporting Information files.

Copyright

© 2025 Ayah Rashwan et al. This article is distributed under the terms of Creative Commons Attribution License which permits unrestricted use, distribution and reproduction in any medium provided the original author(s) and original publisher are properly credited. Please see the copyright policy on the journal website for more information.

Access full text article on
other devices



Access PDF of article on
other devices





INTERNATIONAL JOURNAL OF
CASE REPORTS AND IMAGES



VIDEO JOURNAL OF
CLINICAL RESEARCH



VIDEO JOURNAL OF
BIOMEDICAL SCIENCE



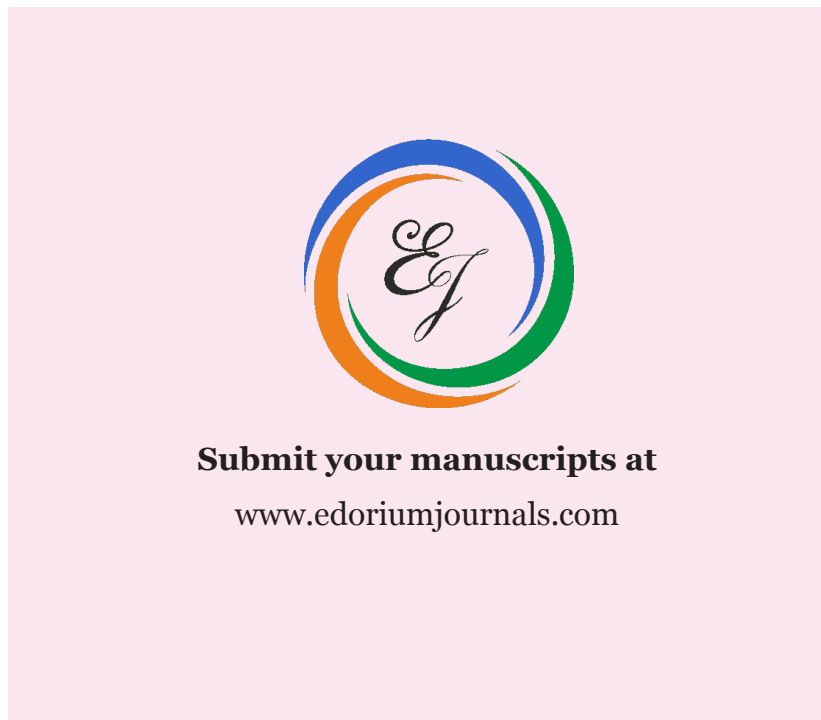
INTERNATIONAL JOURNAL OF
HEPATOBIILIARY AND
PANCREATIC DISEASES



INTERNATIONAL JOURNAL OF
BLOOD TRANSFUSION AND
IMMUNOHEMATOLOGY



EDORIUM JOURNAL OF
OPHTHALMOLOGY



EDORIUM JOURNAL OF
MEDICINE



EDORIUM JOURNAL OF
CARDIOTHORACIC AND
VASCULAR SURGERY



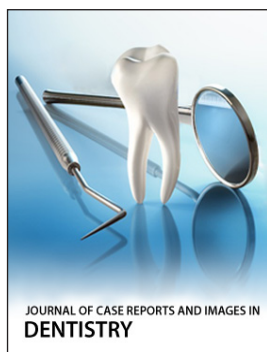
JOURNAL OF CASE REPORTS
AND IMAGES IN ORTHOPEDICS
AND RHEUMATOLOGY



EDORIUM JOURNAL OF
PSYCHOLOGY



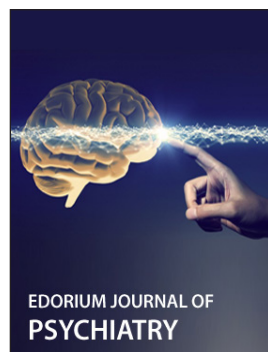
EDORIUM JOURNAL OF
CELL BIOLOGY



JOURNAL OF CASE REPORTS AND IMAGES IN
DENTISTRY



EDORIUM JOURNAL OF
CANCER



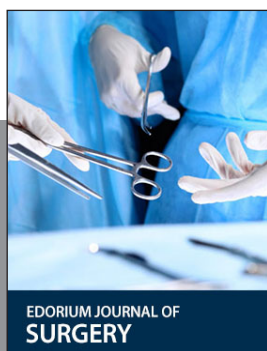
EDORIUM JOURNAL OF
PSYCHIATRY



JOURNAL OF CASE REPORTS AND
IMAGES IN INFECTIOUS DISEASES



EDORIUM JOURNAL OF
ANATOMY AND EMBRYOLOGY



EDORIUM JOURNAL OF
SURGERY



JOURNAL OF CASE REPORTS
AND IMAGES IN PATHOLOGY



EDORIUM JOURNAL OF
ANESTHESIA