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In this issue

Case Report

Open Access Case Report PTZAID:AHCRR-7-140

Recurrent primary cutaneous anaplastic large cell lymphoma in young female patient from Ethiopia: A Case Report

Published On: August 30, 2022 | Pages: 019 - 022

Author(s): Temesgen Assefa Ayele*, Abel Tenaw Tessema and Fisihatsion Tadesse

Primary Cutaneous Anaplastic Large Cell Lymphoma (PC-ALCL) is a CD30+ lymphoproliferative disease of the skin characterized by single or focal nodules or plaques that ulcerate over time. Diagnosis of PC-ALCL relies heavily on clinicopathological correlations because of the potential morphological, clinical, and molecular overlap with other cutaneous CD30+ LPDs. Histop ...

Abstract View Full Article View DOI: 10.17352/ahcrr.000040

Open Access Case Report PTZAID:AHCRR-7-139

Synchronous seminoma and Langerhans cell histiocytosis with a multilocular thymic cyst: A rare case report

Published On: August 23, 2022 | Pages: 015 - 018

Author(s): Ji Hye Kim and Hee Jeong Cha*

A 28-year-old man with an anterior mediastinal mass underwent total thymectomy. The surgical specimen showed multilocular cysts and nodules. On microscopic examination, the cysts were lined by squamoid epithelium, consistent with thymic cysts and tumors showing two different histologic features were identified. One tumor component was composed of sheets of monotonous, ...

Abstract View Full Article View DOI: 10.17352/ahcrr.000039

Open Access Case Report PTZAID:AHCRR-7-138

Hereditary elliptocytosis discovered during work-up for infective endocarditis: About a case

Published On: May 05, 2022 | Pages: 013 - 014

Author(s): Mahjouba Baiya*, Imane Elkhannouri, Ibtissam Mhirig and Sanae Sayagh

Hereditary elliptocytosis is a group of red blood cell membrane disorders that are characterized by elliptical-shaped erythrocytes and shortened red blood cell survival [1]. It is due to protein abnormalities involving the horizontal skeletal network of the red cell membrane, including the spectrin dimer-dimer interaction or the spectrin-actin-protein 4.1 junction com ...

Abstract View Full Article View DOI: 10.17352/ahcrr.000038

Open Access Case Report PTZAID:AHCRR-7-137

Coronavirus Disease 2019 (COVID-19)- Associated central retinal vein occlusion: A case report and literature review

Published On: March 03, 2022 | Pages: 009 - 012

Author(s): Mohammad Hassan Hodroj, Ziad F Bashshur, Charbel Wahab, Kawthar Jarrah and Ali Taher*

Coronavirus Disease 2019 (COVID-19) is caused by the Severe Acute Respiratory Syndrome Coronavirus 2(SARS-CoV-

2) that is associated with several inflammatory and vascular endothelial complications. Ocular vascular occlusive events were reported in COVID-19 patients including both Central Retinal Vein Occlusion (CRVO) and Central Retinal Artery Occlusion (CRAO). We rep ...

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Open Access Case Report PTZAID:AHCRR-7-136

Hemophagocytic lymphohistiocytosis secondary to epstein-barr virus reactivation in a patient with COVID-19

Published On: February 16, 2022 | Pages: 006 - 008

Author(s): Nurfiza Ladak*, Kenneth Csehak, Justin Chan and Farnoush Moen

Hemophagocytic lymphohistiocytosis (HLH) in coronavirus disease 2019 (COVID-19) is a recognized complication of severe illness. However, this phenomenon has been reported most often in the setting of acute infection. Here we present a case of a patient with a history of severe acute respiratory syndrome Coronavirus 2 (SARS-CoV-2) infection that subsequently developed ...

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Combined treatment with vemurafenib and cobimetinib in Langerhans cell histiocytosis and Erdheim-Chester disease overlap syndrome: A case Report

Published On: January 20, 2022 | Pages: 001 - 005

Author(s): Hidalgo-Soto Marta*, Poza-Santaella María, Pita-Suárez Daniel, Calbacho-Robles María, Pina-Sánchez José, González-Medina José and Baumann Tycho Stephan

Histiocytoses are clonal disorders diseases derived from the monocyte-macrophage lineage. The Erdheim-Chester Disease (ECD) and Langerhans Cell Histiocytosis (LCH) may occur in association with overlapping clinical, histopathological and molecular features, harboring somatic MAP2K1 mutations in more than 50% of patients. BRAF and MEK inhibitors have shown to be effica ...

Abstract View Full Article View DOI: 10.17352/ahcrr.000035