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In this issue

Research Article

Open Access Research Article PTZAID:GJRD-7-134

Communicative and pragmatic skills: down syndrome vs williams syndrome

Published On: October 18, 2022 | Pages: 011 - 016

Author(s): Esther Moraleda-Sepúlveda*, Patricia López-Resa, Noelia Pulido-García, Paula López-Peces, Noelia Santos-Muriel and Gabriela Gutierrez-Arce

Down Syndrome (DS, hereinafter) is a chromosomal alteration produced by changes in the DNA sequence of chromosome 21, in addition to being the main cause of intellectual disability in the world [1]. According to the World Health Organization [2], DS has a worldwide prevalence of 1 in every 1,000 live births, but these figures vary due to different factors such as pren ...

Abstract View Full Article View DOI: 10.17352/2640-7876.000034

Review Article

Open Access Review Article PTZAID:GJRD-7-132

Erythromelalgia: Definition, clinical contexts, differential diagnosis, and therapy. Single case and literature update

Published On: April 12, 2022 | Pages: 001 - 007

Author(s): Giulio Perrotta* and Emanuele Guerrieri

Clinical publications on this morbid condition report as many as 16 different Nav1 7 channel substitutions. They also report more than 40 pharmacological treatments, including agents with sodium channel blocking but nonspecific activity (mexiletine, lidocaine, carbamazepine), anti-neuropathic drugs (gabapentinoids and antidepressants), antidepressant drugs with effica ...

Abstract View Full Article View DOI: 10.17352/2640-7876.000032

Case Report

Open Access Case Report PTZAID:GJRD-7-133

Shy-Dragger Syndrome – An underdiagnosed sad reality?

Published On: August 29, 2022 | Pages: 008 - 010

Author(s): Vasco Pinto Neves*, Maria Pacheco, Maria Inês Silva Gonçalve, Joao Rodrigues, Inês Salvado, Jessica Fidalgo and Pedro Daniel Carneiro Ventura

Multiple System Atrophy (MSA) regards a group of neurodegenerative diseases sharing the same physiopathology. It is a rare group of diseases and often represents a diagnostic challenge for clinicians. Mild symptoms are present at the onset of the disease and are often neglected by patients. The case report describes a 62-year-old female with multiple episodes of synco ...

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