THE IMPORTANCE OF THE DENTIST FOR THE STRUGER WEBER SYNDROME

MONFERDINIV*, Ferreira L*, Dos Santos N*, Leinmuller MMC***, Giro EMA***, Hebling J***, Souza YTCS***, Oliveira CF***
Laboratorio De Patologia, Faculdade De Odontologia Da Universidade de Ribeirão Preto (UNAERP)

Struger Weber syndrome is characterized by an abnormality in the development of the primary capillaries in the leptomeninges, eyes and face. This syndrome is a rare congenital malformation, not hereditary, with pathophysiological manifestations. It is characterized by a corticocerebral angiomatosis, cerebral calcifications, epilepsy, oculars disorders, mental retardation and easily nerve, following the trigeminal nerve. This study aims to report the case of a patient with the syndrome Struger Weber (MHB), 16 year-old male who sought Service to Patients with Special Needs (Serviço de Atendimento ao Paciente com necessidades especiais) - UNESP for dental treatment. On examination, it can be observed hemangiomatous gums lesions, palate and buccal mucosa located in the left epsilateral maxilla region. Flameo nervus (or stains coming from Porto) can be seen in his left hemiface, affecting the region of the forehead, nose and eye, extending to the upper lip, generating mental retardation and epilepsy. The pre-and-post-surgical become more complicated and hampered in this type of syndrome, whereas for surgery there is a high risk of severe bleeding in the regions affected by the PCH, so it is extremely important to emphasize the need for preventive procedures on this type of patient.