Introduction

Sturge-Weber syndrome (SWS), also known as encephalotrigeminal angiomatosis, is a rare, neurocutaneous, non-inherited neurovascular disorder characterized by abnormalities in the vasculature associated with angiomas involving the face (PWB), choroid, cerebral vasculature alterations (leptomeningeal vascular malformation), and ocular disorders [1,2]. Primarily affects the brain, skin, and eyes and causes seizures, developmental delay, intellectual disability, learning disabilities, and glaucoma. The incidence is not well known and estimated to be 1 in 20,000 – 50,000 live births, this disease affects males and females equally and there is no race predilection [1,3]. The most common somatic mutation (p.Arg183Gln, p.R183Q) in the guanine nucleotide-binding protein G(q) subunit alpha (GNAQ) gene [4,5] although GNA11 and GNB2 have also been implicated.

These patients have stroke-like episodes, due to impaired cerebral perfusion with increased risk of stroke, commonly appear as transient episodes, lasting more than 24 hours of hemiparesis or visual field defects, and are often difficult to distinguish from ongoing seizure activity or postictal Todd’s paresis, for which an important part is a treatment with aspirin, in addition to ASM [6-8]. 75% of patients present seizures during the first year of life, mainly focal motor seizures, with or without consciousness impairment.
more often diffuse in nature but can be circumscribed and is usually ipsilateral to the facial PWB. It is caused by aberrant and enlarged vessels and vascular channels that cause variable clinical features including reduced visual acuity, refractive errors, scotoma, or retinal detachment. The ophthalmological examination is also recommended every 3 months during the first year and yearly afterward to monitor for glaucoma and choroidal hemangiomas [10].

Case report

This is a 33-year-old patient with a diagnosis of PWB at birth (Figure 1), in addition, seizure onset at age of 4 months of age with 3 seizure types: 1) Focal onset seizure with impaired awareness, left hemibody clonic to bilateral tonic-clonic, 2) Focal onset seizure with impaired awareness, myoclonic motor, and 3) Focal onset seizure with retained awareness, the tonic motor with a Jacksonian gait, characterized by onset in the left thoracic limb that spreads to the left hemibody. With a past medical history of delay in the milestones, accompanied by cognitive impairment. SWS was diagnosed based on clinical characteristics despite no genetic study being performed during the diagnostic protocol. The patient was seizure free for one year during childhood, adolescence, and adulthood, but experienced breakthrough seizures during their last admission. Her seizure rate average was usually 5 per month, with catamenial exacerbation and multiple admissions due to intractable epilepsy and once status epilepticus (SE). Previously under treatment with gabapentin 300-300-600 mg and phenytoin 150 mg TID. Previously under treatment with gabapentin 300-300-600 mg and phenytoin 150 mg TID. Her last admission was due to breakthrough seizures secondary to presenting fever and cough with expectoration, for which community-acquired pneumonia was diagnosed, culminating in a CSE. The symptoms did not respond after the administration of benzodiazepines and ASM, for which she required sedation, advanced airway management, and antibiotic treatment, remaining for 9 days in the intensive care unit. Subsequently, as part of the approach, a thyroid profile was performed, which reported thyroid-stimulating hormone 1.77 mUI/ml, T3 0.95 ng/ml, T4 12.56 um/dl, an electroencephalogram under sedation effect, which was reported with epileptiform activity over the right frontotemporal region and continue generalized slowing (Figure 2), and non-contrast brain computed tomography (CT) showing brain parenchyma with decreased volume in the right frontal and parietal region, with hyperdense linear lesions due to vascular malformation, right cerebellar atrophy, and gyri form calcifications are classically described as “tram-track sign” (Figure 3) (Singh & Keenaghan, 2023). Brain magnetic resonance imaging (MRI) was performed, where right cerebral atrophy, secondary frontal hyperostosis, and abnormal capillary vessel on the right cerebral convexity were observed (Figure 4).

Medical treatment is adjusted with levetiracetam 1500 mg BID, valproate acid 600-400-600 mg, topiramate 50 mg BID, clonazepam 250 mg every 24 hours, and aspirin 100 mg every 24 hours, with clinical improvement. Currently, the patient has an improvement in the seizure rate, without presenting SE, currently with 3 seizures per month. She is following up with ophthalmology, physical rehabilitation, and neurology.
Discussion

This case presentation highlights essential considerations for health professionals regarding the study of patients with this disease. SWS is a rare congenital vascular disorder characterized by a cutaneous capillary malformation, [11] that does not present mendelian inheritance. F. Parkes Weber whose 1922 paper led to the naming of the syndrome as Sturge-Weber. Historically SWS was categorized as one of the phakomatoses [12]. Characteristically, these patients present a capillary malformation, called PWB, a leptomeningeal malformation that typically enhances contrast on MRI [13]. Steve Roach in 1992 proposed a classification for the spectrum of "encephalotrigeminal angiomatosis". The Roach Scale was used for SWS classification. Type I patients have both facial and leptomeningeal angiomats. Type II patients have isolated facial angiomata (no CNS involvement). Type III patients have isolated leptomeningeal angiomatas. Type IV patients have facial angiomata with CNS involvement [14]. Nonetheless, this outdated classification incorrectly implies that all patients with facial capillary malformation (without brain or eye involvement) have type II SWS. At present, we now know that the syndromic association to SWS and the risk of both brain and eye involvement can be more accurately stratified based on the extension and location of the facial capillary malformation [10]. SWS can also be accompanied by ocular vascular malformation, which can cause glaucoma, which has been reported as the most frequent ocular manifestation, with an incidence of 30% - 70%. [15] Pressure is treated with eyedrops, such as timolol and latanoprost, which decrease fluid production in the eye. Frequently, medical management fails, or the glaucoma is fulminant (particularly in infants) and surgery is required with shunt placement or other means of relieving ophthalmic pressure [3].

Patients with this diagnosis may present episodes of stroke-like events characterized by focal seizures, hemiparesis, speech, and cognitive disorders. Recurrent thrombosis has also been postulated as a potential cause. Based on these suppositions, preventative treatment using low-dose aspirin at 3 to 5 mg/kg/day has been proposed and used by many to limit or prevent stroke-like episodes and improve neurodevelopmental outcomes. [3,6,10]. This dual positive effect may reflect the prevention of thrombosis in the setting of venous stasis, thereby improving blood flow to the brain, and reducing the risk of seizures [10]. EEG can be normal or show asymmetrical background activity amplitude or frequency as well as interictal activities or seizures [1]. EEG is an important test to perform for SWS patients with a febrile illness and altered mental status (unexplained confusion or unresponsiveness), as they are at risk [6].

Some differential diagnoses are Blue rubber bleb nevus syndrome, Klippel-Trenaunay-Weber syndrome, PHACES (posterior fossa abnormalities, hemangiomas, arterial anomalies, cardiac, eye, and sternal anomalies), and Wyburg-Mason syndrome [1].

In the case of our patient, who has been diagnosed with SWS since birth, presenting the characteristic facial angioma in the right frontotemporal V1 trigeminal region, Type I according to the Roach Scale [14], who started with seizures at 4 months of age, which have been refractory according to the ILAE criteria [16]. The patient has a history of breakthrough seizures, generally triggered by upper and lower airway infections. Among the main causes of SE are the reduction of ASM at 34%, metabolic causes at 15%, and systemic infections at 7% [17]. It has been observed that up to 57% of patients with SWS present SE during their evolution [18]. Among the challenges to be dealt with in this patient is proper management and avoiding triggering events that may exacerbate seizures.

Conclusion

This case illustrates the natural history of SWS, ranging from facial angioma to the onset of seizures before the first year of age, together with cognitive alterations and difficult-to-control seizures. The drugs most used in this disease are ASM and aspirin, which are postulated to reduce stroke events. Currently, the use of cannabidiol has also been reported for patients with refractory seizures, with which there is a decrease in seizures, improvement in quality of life, and subjective improvement in language, motor, and cognitive skills. The use of mTOR inhibitors has also been mentioned, hypothesizing a decrease in the risk of cerebrovascular events, normalizing the function of the affected vessels, with reports of significant improvement in anger control, depression, and quality of life. This disease presents with a wide spectrum of multisystemic manifestations, so a multidisciplinary approach is required to achieve the expected quality of life, despite the challenges that this disease provides us.
Ethics statement

Written informed consent was obtained from the individual(s) for the publication of any potentially identifiable images or data included in this article.

References