Vaskularna malformacija na licu: rijedak slučaj praćen 18 godina

Vascular Malformation of the Face: a Rare Case with Follow-up of 18 Years

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Svrla rada: Vaskularne malformacije strukturne su anomalije krvnih žila. Prisutne su od rođenja i ostaju cijeli život. Mogu se klasificirati prema vrsti zahvaćenih krvnih žila (kapilarne, venske, arteriovenanske) i hemodinamskim obilježjima (1). Vinski madež je kongenitalna kapilarna malformacija koja se obično nalazi u području glave i vrata oko 0,3 posto novorođenčadi (2). Tijekom godina većina tih mrlja potiče hipertrfiju mekog tkiva, pri čemu taj rast može uzrokovati tešku deformaciju lica. Ma- terijali i metode: U ovom radu opisano je specifično za proliferativne vaskularne anomalije koja je nastala od vinskog madeža na licu. Rezultati: U prikazu se naglasak stavlja na kontinuirani i proliferativni rast lezije koja je pronađena 18 godina te na poteškoće pri utvrđivanju dijagnoze zbog nje-zine složenosti i dimenzije te okolnosti povezanih s lošim socijalno-ekonomskim statusom pacijen-ta. Zaključak: Opisivamo rijedak i neobičan slučaj divovske proliferativne vaskularne malformacije koja potječe od vinskog madeža na licu.

Uvod

Vaskularne malformacije strukturne su anomalije krvnih žila. Prema definiciji, prisutne su od rođenja i ostaju tijekom cijeloga života. Mogu se klasificirati prema vrsti zahvaćenih krvnih žila (kapilarne, venske, arteriovenanske) i hemodinamskim obilježjima (1). Vinski madež je kongenitalna kapilarna malformacija koja se obično nalazi u području glave i vrata oko 0,3 posto novorođenčadi (2). Tijekom godina većina tih mrlja potiče hipertrfiju mekoga tkiva, pri čemu taj rast može uzrokovati tešku deformaciju lica (3 - 5).

S obzirom na sve navedeno, cilj ovoga rada bio je opisati specifično za proliferativne vaskularne anomalije koja je nastala od vinskog madeža na licu. Ovaj slučaj zanimljiv je zato što je bilo moguće prati ga 18 godina, te zbog kontinuiranog i ekspanzivnog rasta vaskularne malformacije, što je rezultiralo značajnom deformacijom lica.

Introduction

Vascular malformations are structural anomalies of the blood vessels. By definition, they are present at birth and persist throughout life. These malformations can be classified according to the type of vessel involved (capillary, venous, arteriovenous) and to hemodynamic characteristics (1). Port-wine stain (PWS) is a congenital capillary malformation commonly found in the head and neck region, which is observed in approximately 0,3% of newborns (2). Over time, most of these stains result in soft tissue hypertrophy and this growth can cause severe facial deformity (3 - 5).

In view of the above considerations, the objective of this study was to describe an unusual case of a proliferative vascular anomaly arising from a PWS on the face. This case is interesting because of the possibility of follow-up for 18 years, its continuous and expansive growth, and the proliferation of the vascular malformation resulting in significant facial deformity.

Case Report

A 60-year-old male, rural worker from the Northeastern region of Brazil has been working outdoors exposed to sunshine since he was a 10-year-old boy. The patient reported having been born with a deep purple stain on the upper part
of the face, no correlated family history. As an adult, a PWS was diagnosed on the left side of the face, extending along the ophthalmic branch of the trigeminal nerve. At age 42, a tumor proliferation was evident, which involved the eyes, nose and upper lip (Figure 1). Currently, the patient is 60 years old and he presented with an exuberant tumor of vascular origin in the left upper part of the face, which was initially associated with a PWS on his hemi face. The tumor has been growing over a period of 18 years, involving an extensive area from the upper facial third to the upper lip and respecting the limits of the midline of the face and ear. The tumor reached extensive proportions, forming large, rough, red lobes of varying sizes and extending in different planes depending on the growth of these proliferations. These lobes grew as shiny, lobular, pedunculated masses measuring approximately 20 cm in diameter, joined closely together, thus reducing eye opening and deforming the nose (Figure 2). The patient felt embarrassed and suicidal and was complaining about headache,
je otkrio više nodularnih tvorbi mekoga tkiva na licu i odsutnost intrakranijalnih patoloških kalcifikacija (slika 3.). Komputerizirani tomografiji s kontrastom identificirani su oblici klastera čvrstih čvorova lociranih u koži i u prokožnom staničnom tkivu lijeve temporo-parietalne regije, bez invazije na kosti (slika 4.). Tim liječnika i stomatologa zatražio je kliničke i komplementarne prilike koje su upućivala na dijagnozu vinskog madeže, malformaciju koja je perzistirala i dosegla velike razmjere tijekom pacijentova života.

**Rasprava**

**Medunarodno društvo za proučavanje vaskularnih anomalija** takve promjene klasiﬁcira kao vaskularne tumore i vaskularne malformacije (6). Ti entiteti mogu se dijagnosticirati kliničkim instrumentima. No u neizvjesnim slučajevima kada dijagnoza nije sigurna, od pomoći mogu biti bilo obojeni Dopplerov uređaj, angiograﬁja i biopsija (7, 8). Takav pristup moga bi pridonijeti dijagnosticčkoj preciznosti, ali naš je pacijent odbijao pretrage u zatvorenim strojevima.

**Vinske mrlje** najprije se pojavljuju kao ružičaste makule koje se povećavaju kako dijete raste dok ne dosegnu stupanj hipertrofije mekoga tkiva koja može uzrokovati nodule, pa čak i tumore (4, 5). Unatoč nedostatku istraživanja koja po- vezuju izloženost suncu i proliferaciju vaskularnih malformacija, činjenica da je pacijent bio poljski radnik izložen cijelog života suncčevoj svjetlosti, povećava mogućnost da upravo to može biti etiološki čimbenik koji je potaknuo znatni rast kongenitalne tvorbe (2). U medicinskoj literaturi nisu dostupni nikakvi podaci o ovoj vrsti lezija koje nastaju iz vinskih madeže i razvijaju se u takvim razmjerima.

Na temelju tih nalaza i uz pomoć multidisciplinarog tima koji je bio uključen u slučaj, može se zaključiti da je pacijent imao kongenitalnu krvazožilnu malformaciju koja je dosegla značajne razmjere tijekom života. Taj je poremećaj verovatno bio arterijskog podrijetla, jer su venske malformacije asimptomatske. Pacijent je cijeli život imao osetljivost na zahtjev područja. No zbog nemogućnosti pod- vrgavanja angiograﬁji, lezija se najbliže može dijagnosticirati kao arterijska malformacija, jer su one venske obično tamno ružičaste, a arterijske plave i pacijenti rijetko imaju simptome (7). Proliferativne malformacije kod našeg pacijenta bile su pretežno crvene.

Prognoza i liječenje vaskularnih malformacija koje nastaju iz vinskih madeže kontroverzni su. Cerri i suradnici u svojem radu iz 2014. smatraju da je terapija sigurna i odsutna, uz minimalno povećani rizik od krvenja, jer se malformacija uglavnom nalazi u površinskim tkivima (3). No naš je pacijent imao velike lezije (> 20 cm), pa bi tehnike liječenja narekle u literaturi svakako bile izazov s povećanim rizikom od smrti.

dizziness and a hot, cold and pulsatile feeling in the affected region. A cranial X-ray revealed multiple nodular soft tissue images on the face and the absence of intracranial pathological calcifications (Figure 3). Contrast-enhanced computed tomography identified soft tissue densities formed by clusters of solid nodules located in the skin and subcutaneous cellular tissue of the left/temporoparietal region without bone invasion. (Figure 4). A team of physicians and dentists requested clinical and complementary exams, which suggested the diagnosis of a PWS, a malformation that persisted and reached large proportions during the life of the patient.

**Discussion**

The International Society for the Study of Vascular Anomalies classified vascular anomalies into vascular tumors and vascular malformations (6). These entities can be diagnosed by clinical means. However, in uncertain cases, color Doppler, angiography and biopsy can be of help when the diagnosis is not certain (7, 8). This approach could contribute to diagnostic precision, but our patient was extremely reluctant to undergo examinations in closed devices.

Port-wine stains first appear as a pink macule that progresses in size as the child grows, reaching a stage of soft tissue hypertrophy that can cause nodules and even tumors (4, 5). Despite the scarcity of studies correlating sun exposure with the proliferation of vascular malformation, the fact that the patient was a rural worker exposed to sunlight throughout his life, raises the possibility that the sun exposure could be an etiological factor that promoted considerable growth of the congenital PWS reported by the patient (2). No reports of this type of lesion, which arise from a PWS and develop to such a great extent, are available in the medical literature.

Based on these findings and with the help of the multidisciplinary team involved in the case, it can be concluded that the patient has a congenital vascular malformation that reached considerable proportions during his life. This malformation was probably of arterial nature since venous malformations are asymptomatic. The patient reported a constant pulsatile feeling on the affected region throughout his life. However, due to the patient’s unwillingness to take angiography, it is best to diagnose the lesion as an arterial malformation, since patients with venous malformations are usually deep blue in color and they seldom report symptoms (7). The proliferative malformations in our patient were essentially red.

The prognosis and treatment of vascular malformations arising from PWS is controversial. In 2014, the treatment was considered safe and effective by Cerri et al, with only a minimally increased risk of bleeding, since the malformation is mainly located in superficial tissues (3). However, our patient had extensive lesions (> 20 cm), therefore, the treatment techniques described in the literature would certainly be a great challenge for our healthcare team, with an increased risk of death.
Abstract
Objective: Vascular malformations are structural anomalies in the blood vessels. They are present at birth and persist throughout life. These malformations can be classified according to the type of vessel involved. A port-wine stain is a vascular malformation characterized by an increased number of ectatic vessels in the dermal vascular plexus, which can be found in any part of the body, including the head and neck region. Over time, most of these stains result in soft tissue hypertrophy and this growth can cause severe facial deformity. Materials and Methods: This study describes a rare case of a giant proliferative vascular anomaly arising from a port-wine stain on the face. Results: The report highlights the continuous and proliferative growth of the malformation observed after follow-up of 18 years, as well as the difficulty in establishing the diagnosis due to the complexity and dimension of the lesion and the conditions related to the patient’s low socioeconomic status. Conclusions: We have described a rare and unusual case report of a giant proliferative vascular malformation arising from a Port-wine stain on the face.

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