CEREBROFACIAL ARTERIOVENOUS METAMERIC SYNDROME (CAMS) 2 RARE ENTITY: HOW TO DIAGNOSE

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ABSTRACT

Cerebrofacial arteriovenous metamic syndrome (CAMS) is a recent classification of vascular malformations that encompasses a spectrum of phenotypic expression involving arteriovenous malformations (AVMs) of the cerebral, orbital and facial region. Recognizing the CAMS is important for diagnosing other AVMs along the same metamic level. We present a case with intraparenchymal AVM and maxillofacial AVM without optic nerve involvement, characteristics of CAMS 2. The diagnosis of intracranial and facial AVM was made by magnetic resonance imaging of the brain and confirmed with digital subtraction angiography.

Key words: Cerebrofacial Arteriovenous Metameric Syndromes (CAMS), natural history, angioarchitecture

Introduction

A rare case of Cerebrofacial Arteriovenous Metameric Syndrome (CAMS) in a 28 years old man is described with arteriovenous malformation involving simultaneously brain and the face. This case represents CAMS 2 according to the classification described by Lasjaunias et Al. The metamic distribution of the cerebrofacial arteriovenous syndromes is well illustrated in this case. The extracranial and intracranial involvement is described and appearance on imaging is detailed. In our opinion this case demonstrates a spectrum of disease expression of cerebrofacial vascular structures related by the same neural crest origin, named cerebrofacial arteriovenous metamic syndrome (CAMS) by Bhattacharrya et Al.1 We describe CAMS in terms of natural history, clinical presentation, angioarchitecture, intracranial AVM and proper investigation to diagnose this suspicious case.

Case Report

A 28 year old man presented at the age of 28 years with facial asymmetry. This started at infancy as a small mark on right face and developed into a very large maxillo-facial venous malformation. He underwent near total surgical debulking in 2008 (at 21 years of age). The swelling regrows after 3 months interval. Now for the past one year he had occasional episodes of bleeding from skin locally along with epistaxis and swelling becomes painful as well. A month ago patient was referred to our department for diagnosis and treatment purposes. During history taking he disclosed trivial headaches. As a part of investigation magnetic resonant imaging of brain done which revealed intracranial AVM which was missed. He works as a mechanic. There is no past history of seizures or any family history of venous malformation.

Clinical examination revealed a physically well developed 28 year old right handed man. He had an asymmetry of face due to swollen right cheek. He is neurologically intact. (Fig-1a-1b)

Imaging MRI examination done recently demonstrated a right sided facial venous malformation involving the soft tissues of the cheek and lower eyelid and extending into deeper structures of infratemporal fossa on right side. (Fig.2a-2b-2c)
MRI of the brain revealed sulcal AVM right post temporal lobe along the marginal/triangular gyrus (Fig. 3a-3b).
Diagnostic angiography showed the high flow arteriovenous malformation locating right cheek, fed by the right internal maxillary artery, draining through facial vein. Selective angiography of right internal carotid artery showed right posterior temporal lobe sulcal arteriovenous malformation, fed by the inferior trunk of middle cerebral artery, draining through superficial venous system. (Fig. 5a-5b-5c)

CT angiography showed right cheek arteriovenous malformation and intracranial AVM which was missed on imaging. (Fig. 4)
Discussion

An association between arteriovenous malformation of face, retina and brain was first recognized by Bonnet, Dechaume and Blanc in 1937 and Wyburn-Mason in 1943. Furthermore, in 1995 Couly demonstrated that endothelial cells of cephalic region have a regionalized origin providing blood vessels to specific regions of face and brain. The neural crest and mesodermal cells arising from a certain transverse level occupy the same facial territories. Since the endothelium and media of blood vessels are derived from mesoderm and neural crest respectively, Lesjau-nias group used this approach to regroup the neural tube and neural crest derivatives to associate between certain maxillofacial and central nervous system malformations. Moreover, the case report of Gupta et al describing two patients there were unilateral cerebral and orbital AVMs with facial involvement (a venous malformation in one case and a maxillofacial AVM in the other). One of these patients had optic atrophy but in neither did fundoscopy identify a retinal AVM. Lesjau-nias group suggested the name Cerbrofacial Arterial/venous Metameric syndrome (CAMS or CVMS) for conditions with arterial or venous malformations simultaneously involving the brain and the face. According to new classification three subgroups are identified within CAMS or CVMS. CAMS 1 or the “median prosencephalic group” involves the hypothalamus and the hypophysis as the intracranial component and nose as the facial component. CAMS 2 or the “lateral prosencephalic group” involves the occipital lobe, thalamus, optic tract and retina intracranially and the maxilla as its facial component. CAMS 3 or the “lateral rhombencephalic group” involves the cerebellum and pons intracranially and the maxillary as the facial component. Using this classification our case represents CAMS 2. The intracranial findings in our case, although predominantly ipsilateral to the facial involvement but does not involve retina. This could be explained by Couly’s comment that the regionalization of the cephalic neural crest and mesoderm is not strict. According to him this may allow some fluidity in the final expression. Furthermore, this is supported by similar bilaterality reported by Boukobza et al in one of their unilateral facial malformation patients.

Conclusion

The diagnosis of cerebrofacial arteriovenous metameric syndrome (CAMS 2) encompasses a spectrum of phenotypic expression. A portion of these patients manifest the complete expression of the disease with arteriovenous malformation of brain and additional high flow arteriovenous malformation of the maxillofacial region. These presents the additional risks of life threatening hemorrhage. We suggest any patient who present with facial arteriovenous malformation or venous malformation must be investigated further for intracerebral arteriovenous malformations.

References


