Non-syndromic megalencephaly and epilepsy: Our findings

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Abstract

Megalencephaly (MEG) and macrocephaly are defined as a head circumference measurements two standard deviations above the age-related mean. A distinction between megalencephaly and macrocephaly has been proposed despite the fact that these terms encompass individuals with a large head and, in some cases, with similar neurologic manifestations including intellelctive disability of various degree, epileptic seizures, and motor impairment. Nevertheless they differ widely in causal events, cerebral structural anomalies, approach in the work-up, treatment and prognosis for which a clinical distinction is justified. From July 2013 through July 2019, 10 children with non-syndromic MEG were selected (7 male, 3 female), and followed up at the Pediatric Unity of the University Hospital Policlinic-Vittorio Emanuele, Catania, Italy for pediatric and neuropediatric disorders; 5 of them (4 male, 1 female) have an abnormally large head, mild/moderate developmental delay, and seizures.

Introduction

Megalencephaly (MEG) and macrocephaly are defined as a head circumference measurements two standard deviations above the age-related mean [1,2]. A distinction between megalencephaly and macrocephaly has been proposed despite the fact that these terms encompass individuals with a large head and, in some cases, with similar neurologic manifestations including intellectual disability of various degree, epileptic seizures, and motor impairment. Nevertheless they differ widely in causal events, cerebral structural anomalies, approach in the work-up, treatment and prognosis for which a clinical distinction is justified [2,3].

MEG is a sign of several unusual conditions and can be differentiated into three groups: idiopathic, metabolic, or anatomic. Anatomic MEG manifests with neurologic signs and may be isolated (“non-syndromic”) or part of complex syndromes, such as Megalencephaly-Capillary Malformation-Polymicrogyria (MCAP) and macrocephaly-cutis marmorata telangiectasica (M-CMTC) [4-6].

Macrocephaly is referred to individuals in whom the brain enlargement is secondary to events inside the brain such as intracranial masses, abnormal ventricular dilatation, hydrocephalus ex vacuo, and increase of bone skeletal structures. On the other hand, megalencephaly refers to anomalous structural cerebral events such as ineffective molecular control of neuronal growth during the various stage of the brain development or to inborn errors of metabolism [2-4]. There are examples where the adverse events causing megalencephaly and macrocephaly may co-exist but semantic distinction among these conditions is suitable. In a previous article, our group seek to underline the clinical aspect of megalencephaly, emphasizing the main disorders that manifest with this anomaly in an attempt to properly categorize these disorders within the megalencephaly group [7].

We reviewed our patient with non-syndromic MEG, harboring minor dysmorphism, mild/moderate developmental delay and epileptic seizures as associated signs of the disorder: we enrolled 10 patients of MEG: five of them (unrelated patients) were affected by MEG and epilepsy.

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Editorial

Among all the forms of MEG, the idiopathic form is the most frequent and is not associated with neurologiical manifestations, intellectual disabilities or other symptoms involving other parts of the body. In this disorder, the patient’s head circumference increases gradually until an age of 18 months and then becomes more stable during the course of development. The children’s parental OFC measurements were within normal ranges. This situation is opposite to what is typically reported in the literature for benign MEG (i.e., a large parental head is frequently reported) [5].

Brain MRI malformative anomalies were uncommonly reported in children with MEG. This situation confirms the hypothesis of Berg and Dobyns [4], who maintain that genetic anomalies affecting brain development in the first steps of neuronal growth may be the cause of cerebral involvement; structural cerebral anomalies may not always be noticeable in brain imaging techniques.

This cohort of children exhibits a set of clinical signs consisting of an abnormally large head circumference (above the 97th percentile),
a mild-to-moderate intellectual disability, epileptic seizures in 5 cases and, in some cases, minor dysmorphism especially facial. In the absence of signs suggestive of known syndromes, a diagnosis of anatomic, non-syndromic MEG was made. Nine of the ten patients did not show structural abnormality of the brain, in the group with epilepsy four of the five patients failed to exhibit structural anomalies in their brain MRI. Only in one case we find a cerebral cortex dysplasia and microgyria. Among the children with epilepsy the EEG was abnormal and indicative of epilepsy diagnosis in all our patients; dysmorphisms were noted in three children; these conditions were not severe and mainly affected the face.

A differential diagnosis between idiopathic MEG and anatomic, non-syndromic MEG is not simple prior to the onset of clinical signs. Developmental delays, although usually mild and moderate, and dysmorphisms, if not severe, may be useful for a diagnosis.

We maintain that an abnormally large head deserves particular attention because this condition may hide relevant neurologic symptoms in the absence of evident brain anomalies. A large head in children with mild-to-moderate developmental delays and minor dysmorphisms may be a clue for subsequent epileptic seizures and a diagnosis of anatomic MEG.

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