CASE REPORT

Infantile GM1-Gangliosidosis Revealed by Slate-Grey Mongolian Spots

Ghizlane Draïss¹, Adil Fouad¹*, Nourddine Rada¹, Ouafa Hocar², Naima Fdil³ and Mohamed Bouskraoui¹

¹Department of Pediatrics, Mohamed VI University Hospital, Marrakesh, Morocco
²Department of Dermatology, Mohamed VI University Hospital, Marrakesh, Morocco
³Department of Biochemistry, Medical School of Marrakesh, Cadi Ayyad University, Marrakesh, Morocco

Abstract:

Introduction: GM1-gangliosidosis is an inherited metabolic disease caused by mutations in the GLB1 gene resulting in deficiency of β-galactosidase. Three forms have been identified: Infantile, juvenile, and adult. The infantile type progresses rapidly and aggressively and a delayed diagnosis hampers the prevention of many neurological deficits. This delay in diagnosis may be due to the variability of clinical expression of the disorder.

Hypothesis: Extensive Mongolian or slate-grey spots deserve special attention as possible indications of associated inborn errors of metabolism, especially GM1-gangliosidosis and mucopolysaccharidosis. Only symptomatic treatments are available for GM1-gangliosidosis; research is underway.

Observation: In this article, we report a case of infantile GM1-gangliosidosis revealed by slate-grey Mongolian spots, a rare condition in Morocco, and a review of the literature.

Conclusion: The finding of persistent and extensive slate-grey mongolian spots in infant could lead to early detection of GM1-gangliosidosis before irreversible organ damage occurs.

Keywords: GM1-gangliosidosis, Metabolic disease, Mongolian spots, Slate-grey spots, GLB1 gene, Mucopolysaccharidosis.

Article History

Received: August 2, 2018
Revised: November 19, 2018
Accepted: December 20, 2018

1. INTRODUCTION

Gangliosidoses are a group of autosomal recessive inherited metabolic disorders, in which accumulation of gangliosides in the CNS leads to severe and progressive neurological impairment. GM1-gangliosidosis is caused by mutations in the GLB1 gene, resulting in the deficiency of β-galactosidase and subsequent accumulation of GM1-ganglioside that accumulates in the brain as well as in the liver, spleen and kidney. Clinically, GM1-gangliosidosis can be classified into three types, type I (infantile), type II (juvenile) and type III (adult). Type I shows a rapid clinical course at the onset of the infantile period and is characterized clinically by neurological symptoms, bone abnormalities and hepatosplenomegaly [1, 2].

* Address correspondence to this author at the Department of Pediatrics, Mohamed VI University Hospital, PB 2360, Ibn Sina Avenue, Marrakesh, Morocco; Tel: +212524300700; E-mail: adilfouadi@hotmail.fr

Mongolian or slategrey spots are grey-blue, congenital pigment spots that suggest an accumulation of dopapositive melanocytes. They are usually round or ovoid, single or multiple and occur in the sacrococcygeal region. Classically, benign and limited, they appear at birth or in the first months of life and disappear spontaneously before puberty. However, when extensive, they deserve special attention. An association between Mongolian spots and inborn errors of metabolism, especially GM1-gangliosidosis and mucopolysaccharidoses, has been described [3]. We present a case of infantile GM1-gangliosidosis revealed by slate-grey spots, a rarely described disease in Morocco, and a review of the literature.

2. CASE REPORT

The patient, a 7-month-old male infant, the only living son of parents of a second degree consanguineous marriage, was born at full-term and adapted well to extrauterine life. His
psychomotor development was normal until the age of 4 months (gesticulated normally, smiled, followed light). An older brother had died at the age of 6 months due to respiratory distress with a clinical picture of psychomotor retardation, macrocephaly, abdominal distension, and blue-grey pigmented cutaneous spots. The present patient was hospitalized (1st hospitalization) for respiratory distress with the appearance of cutaneous spots and psychomotor regression since the age of 4 months. He had coarse facial features, macrocephaly (cranial circumference + 3DS), abolished tendon reflexes, significant psychomotor retardation (axial hypotonia, no voluntary prehension, no smile, sitting position even with unearned support), slate-grey spots scattered on the trunk and limbs, a negative Darier sign, and some atrophic lesions on the trunk (Fig. 1). The ophthalmological examination suggested bilateral blindness (did not follow light), a pale papilla, but no cherry-red macular spot. He had no skeletal deformities. Brain imaging showed matter myelination abnormalities, enlargement of frontotemporal subarachnoid space and hypoplasia of the corpus callosum (Fig. 2). Abdominal ultrasonography revealed homogeneous hepatosplenomegaly. He had no atypical or vacuolated lymphocytes on the blood smear, liver and thyroid tests were normal, the amino acid chromatography was normal and the urinary excretion of glycosaminoglycans was negative. The presence of oligosaccharides in the urine, and serum beta-galactosidase deficiency confirmed the diagnosis of GM1-gangliosidosis. The progression was marked by death at the age of 9 months after severe neurological deterioration. The infant died before genetic testing could be performed and genetic counselling was planned for the parents.

3. DISCUSSION

GM1-gangliosidosis is a neurodegenerative condition for which three main clinical forms have been identified: Type I (severe), type II, and type III. The incidence of GM1-gangliosidosis is considered to be between 1/100,000 and 1/200,000 live births. The infantile type is characterized by psychomotor regression by the age of 6 months, visceromegaly, cherry-red macular spots with facial and skeletal abnormalities. In some patients with the infantile form, cardiomyopathy may sometimes be present. It is thought that impaired elastogenesis and cardiomyopathy in GM1-gangliosidosis are caused by defects in the elastinbinding domain of elastinbinding protein [4, 5]. Juvenile GM1-gangliosidosis usually starts between 7 months and 3 years of age with slowly progressing neurological signs including early
motor problems, strabismus, muscle weakness, seizures, lethargy, and susceptibility to infections. The adult form, the attenuated phenotype, with the onset between 3 and 30 years, is characterized by cerebellar dysfunction, dystonia, slurred speech, short stature and mild vertebral deformities [6, 7].

Among different ethnic groups, over 90% of Native Americans and people of African descent, approximately 80% of Asians, about 70% of Hispanics, and fewer than 10% of Caucasians have Mongolian spots [8]. Typical Mongolian spots are limited benign blue or slate grey macular lesions of varying size, most commonly located in the lumbosacral region. Mongolian spots usually disappear gradually with age, but may occasionally persist. Those associated with inborn errors of metabolism are extensive and show no signs of resolution. Extensive Mongolian spots are most frequently associated with GM1-gangliosidosis, Hurler-Scheie syndrome, Niemann-Pick disease, Hunter syndrome and α-mannosidosis. The findings of generalized Mongolian spots in an infant may represent underlying storage disorders thereby allowing identification of families at risk. They may also take on a much darker nuance in colour [9, 10]. Mongolian spots result from entrapment of melanocytes in the dermis because of arrested transdermal migration from the neural crest into the epidermis. The accumulated metabolites in GM1-gangliosidosis are thought to bind tightly to tyrosine kinase protein, resulting in an increased nerve growth factor. This factor then binds to chemotactic melanocyte receptors, leading to abnormalities in melanocyte migration and producing the large dermal melanocytosis seen [11]. Demonstration of decreased beta-galactosidase activity in cultured skin fibroblasts, urine, white blood cells or amniotic fluid cells establishes the diagnosis [12]. Our patient had facial dysmorphism, hepatosplenomegaly, and slate grey dermal spots suggesting a storage disease. Results of a skin biopsy were inconclusive but a urinary metabolite investigation revealed oligosaccharides suggestive of GM1-gangliosidosis, subsequently confirmed by enzymatic assay.

There are as yet no approved treatments for GM1-gangliosidosis. Research is underway in animal models evaluating gene therapy technologies and intravenous enzyme replacement therapies [13]. At present, only symptomatic treatments are available. Substrate reduction therapy mediated by N-butyldeoxyxynojirimycin (NB-DNJ, Miglustat) has been used with encouraging results for the treatment of patients with mild to moderate type 1 Gaucher disease, Niemann-Pick disease type C and Sandhoff type chronic GM2 gangliosidosis [7].

Substrate reduction therapy using miglustat has been tried in the infantile gangliosidosis patient population (both GM1 and GM2-gangliosidoses). It has been generally well tolerated and has been shown to be safe in this population. It has not been observed to result in marked improvement in symptom management or disease progression. Safety of low-carbohydrate diets in an infantile patient group, however, may be of concern in terms of maintaining adequate nutrition [13]. Intracranial delivery of recombinant Adeno-Associated Virus (AAV) vectors has been shown to be highly effective in animal models. The choice of injection sites and promoters driving transgene expression are important parameters to translate into clinical trials a safe and effective AAV-based therapy for GM1-gangliosidosis [14]. However, some researchers find that cyclodextrins are useful for the treatment of GM1-gangliosidosis [15]. On the other hand, other researchers find that N-Functionalized amino (hydroxymethyl) cyclo-pentanetriols are potent inhibitors of β-D-galactosidases and, for the first time, could be shown to act as pharmacological chaperones for GM1-gangliosidosis-associated lysosomal acid β-galactosidase thus representing a new structural type of pharmacological chaperones for this lysosomal storage disease [16]. In the absence of a specific treatment, especially in Morocco, our patient was initially given symptomatic treatment: Treatment of pulmonary and urinary infections, motor and respiratory physiotherapy. The infant died as a result of severe neurological deterioration before a genetic study could be performed.

CONCLUSION

The broad variability of clinical symptoms in patients with GM1-gangliosidosis disease may cause delays in diagnosis.

In the absence of screening programmes, the finding of persistent, extensive slate-grey Mongolian spots in newborns or infants could lead to early detection and treatment of GM1-gangliosidosis before irreversible organ damage occurs.

ETHICS APPROVAL AND CONSENT TO PARTICIPATE

Not applicable.

HUMAN AND ANIMAL RIGHTS

No animals/humans were used for studies that are the basis of this review.

CONSENT FOR PUBLICATION

A written informed consent was obtained from the parents when they were enrolled.

CONFLICT OF INTEREST

The authors declare no conflict of interest, financial or otherwise.

ACKNOWLEDGEMENTS

The authors thank the patient’s parents, Dr. Klevor Raymond, and the staff of the pediatric department for their help and support.

REFERENCES


