Dubin-Johnson syndrome: an important cause of obstructive jaundice in children

Síndrome Dubin-Johnson: importante causa de icterícia colestática na infância

El síndrome de Dubin-Johnson: importante causa de ictericia colestática en la infancia

ABSTRACT

The Dubin-Johnson syndrome is clinically characterized by recurrent episodes of benign and familial obstructive jaundice. It is identified by the presence of melanin pigment in the hepatocytes. The authors report a case of Dubin-Johnson syndrome in a child with jaundice and hepatosplenomegaly, whose diagnosis was confirmed by the presence of dark brown pigment on microscopy of liver biopsy. They suggest the suspicion of this syndrome in cases of fluctuating obstructive jaundice in children.

Keywords: Cholestasis; Jaundice; Jaundice, Chronic Idiopathic.

INTRODUCTION

In 1954, the first descriptions of benign and familial jaundice with concurrent plasmatic hyperbilirubinemia at the expense of the direct fraction were described, characterized by the deposition of characteristic melanin pigment in hepatic cells. Dubin-Johnson syndrome is a rare, genetically-determined pathology of autosomal recessive inheritance, thus favored by consanguinity. Jaundice evolves in spurts, which are frequently precipitated by fatigue, string emotion, physical exercise or interspersed infections, and is often followed by discrete hepatomegaly and choluria.

In this syndrome, jaundice begins in infancy, at around two years of age, although it can also occur infrequently during the neonatal period. The onset of symptoms can be severe and accompanied by fever, similar to a viral hepatitis. There are no haematological signs attributed to hemolyses; hyperbilirubinemia occurs intermittently, with a predominance of the direct fraction, and the various liver function tests show normal values. In puberty, the symptoms include jaundice, in 100% of the cases, abdominal pain, especially in the right hypochondrium, hepatomegaly, weakness, anorexia, choluria and fecal hypocholia. Some patients have changes in the urinary excretion of coproporphyrins, such as the reduction of coproporphyrin III and a relative increase in the isomer I. These changes may be found to different degrees in family members.

The syndrome occurs due to the defective expression of the MRP2 gene, an ATP-dependent canalicular membrane transporter. The diagnosis is established by performing an oral cholecystography, a bromsulphalein test and a liver biopsy, associated with the clinical frame.

A liver biopsy is the gold standard test for diagnosing the syndrome. It shows the presence of granular brown pigment in centrilobular hepatocyte lysosomes.

In the literature, most reported cases involve young adults, and, in reports involving children, there is usually a history of cholestatic jaundice of spontaneous remission in the neonatal period without an established diagnosis.

CASE PRESENTATION

ANAMNESIS

Patient A.M.A. was female, two years and eight months of age, pardá (brown-skinned), native and resident of the municipality of Augusto Corrêa in northeastern Pará State. The patient was admitted to the pediatric ward, sick for eight months, with symptoms of abdominal distension and intermittent episodes of fever, associated with cutaneous-mucosal pallor and fluctuating jaundice. The child was born
to consanguineous parents and had a history of three episodes of malaria over the past eight months. She was treated in her hometown. There were no previous reports of jaundice.

PHYSICAL EXAMINATION

Jaundice, mucocutaneous pallor and malnutrition were present. The abdomen was distended, normotensive, painful to superficial and deep palpation in the right upper quadrant, with the liver palpable 7 cm from the right costal margin and the spleen palpable 11 cm from the left costal margin.

COMPLEMENTARY TESTS

The total bilirubin was 3.2 mg/dL, including 2.5 mg/dL direct bilirubin. Hemoglobin and protein electrophoresis values were within the normal ranges. The serology was negative for viral hepatitis, toxoplasmosis, syphilis, rubella, herpes, cytomegalovirus, HIV and visceral leishmaniasis. A plasmodium search was negative, and the myelogram presented no alterations.

The abdominal ultrasound revealed a hepatosplenomegaly with homogenous echotexture and no other abnormalities. A liver biopsy by light microscopy revealed the preserved architecture of the organ and hepatocytes with cytoplasmic deposits of fine, dark brown, granular pigment, distributed throughout the lobe. The deposits were Perls-negative and PAS-positive, consistent with Dubin-Johnson syndrome (Figure 1). No oral cholecystography exams, bromsulphalein test or mutation research by molecular biology were conducted, due to technical difficulties.

PATIENT EVOLUTION/OUTCOME

The patient had spontaneous resolution of the jaundice, with a slow regression of hepatosplenomegaly. After 25 days of hospitalization, the patient was discharged and referred to ambulatory care.

DISCUSSION

Dubin-Johnson syndrome is a rare disorder of genetic transmission, whose molecular basis is a defective gene that encodes the MRP2 organic anion transporter protein. Despite its genetic nature, there are reported cases with no family history of the disease. In the present study, the patient was born to consanguineous parents and did not report any family history of symptoms consistent with the syndrome.

The onset of symptoms at two years and eight months of age was consistent with the age group described in the literature regarding the emergence of jaundice. It is likely that *Plasmodium* infections, which cause malaria, have contributed to the manifestation of the Dubin-Johnson syndrome due to the significant hemolysis presented by this infection.

Discrete hepatomegaly is one of the symptoms of the syndrome, and there are no reports of associated splenomegaly. The marked hepatosplenomegaly found in the patient can be attributed to repeated episodes of malaria that occurred in the months before her hospitalization, since such involvement is part of the disease framework in childhood.

Figure 1 - Presence of dark brown pigment in the cytoplasm of liver cells (hematoxylin eosin, x400)
FINAL CONSIDERATIONS

Although Dubin-Johnson syndrome is a rare disease, it is extremely important to include it in the diagnostic investigation of cases of floating jaundice in childhood. The prognosis is favorable, its course is benign, and no treatment is required. Therefore, establishing the correct diagnosis is important to prevent future unnecessary procedures. The carriers of the syndrome are later during adulthood guided not to use oral contraceptives, which can compete with the hepatocyte secretion of organic anions.

Síndrome Dubin-Johnson: importante causa de ictericia colestática na infância

RESUMO
A síndrome Dubin-Johnson é caracterizada clinicamente por episódios de icterícia colestática recorrente, de caráter benigno e familiar, sendo definida pela presença de pigmento melânico nos hepatócitos. Os autores relatam um caso de síndrome Dubin-Johnson em uma criança portadora de icterícia e hepatoesplenomegalia, cujo diagnóstico foi confirmado pela presença do pigmento castanho-escuro à microscopia hepática realizada a partir de biópsia, e alertam para a necessidade de suspeição dessa síndrome em casos de icterícia colestática fluctuante na infância.

Palavras-chave: Colestase; Ictericia; Ictericia Idiopática Crônica.

El síndrome de Dubin-Johnson: importante causa de ictericia colestática en la infancia

RESUMEN
El síndrome Dubin-Johnson se caracteriza clínicamente por episodios de ictericia colestática recurrente, de carácter benigno y familiar, siendo definido por la presencia de pigmento melánico en los hepatocitos. Los autores relatan un caso de síndrome Dubin-Johnson en una niña portadora de ictericia y hepatoesplenomegalia, diagnóstico que fue confirmado por la presencia del pigmento castaño oscuro en la microscopia hepática realizada a partir de biopsia, y alertan para la necesidad de sospecha de ese síndrome en casos de ictericia colestática fluctuante en la infancia.

Palabras clave: Cholestasis; Ictericia; Ictericia Idiopática Crónica.

REFERENCES


