Thrombocytopenia Absent Radius Syndrome: Rare Cause of Constitutional Thrombocytopenia

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Abstract: The Thrombocytopenia absent radius syndrome is a very rare congenital malformation of genetic transmission which seems autosomal recessive, characterized by a thrombocytopenia associated with absent rays. The presence of thumb is pathognomonic.

Through our work, we illustrate the case of a 6-month-old infant suffering from syndromic thrombocytopenia. His symptomatology dates back a month before his admission by the appearance of bruises on the face. The clinical examination on admission objectified the presence of conjunctival hemorrhage and petechiae of the forehead, adysmorphia and radial club hands. The malformation assessment objectified minimal hydrocephalus and inter-auricular communication. The blood count showed severe thrombocytopenia responsible for externalized hemorrhage requiring transfusion of the red blood cell and platelet. Genetic confirmation was not done since the mother left against medical advice.

1. INTRODUCTION

Thrombocytopenia in children is defined by a platelet count below 50 g / L. Normal values before the age of 15 are, for 95% of children, between 165 g / L and 473 g / L with a median value of 299 g / L [1]. The discovery of isolated thrombocytopenia most often evokes the diagnosis of immunological thrombocytopenic purpura in children. However, in some cases, other diagnoses such as bone marrow hypoplasia, myelodysplasia, or genetic thrombocytopenia should be considered.

We illustrate through our work, the case of a 6-month-old infant suffering from syndromic thrombocytopenia.

2. CASE REPORT

This is a 6 month old female infant from non-consanguineous parents with no specific history. His symptomatology dates back a month before his admission by the installation of bruises on the face. She was referred to our training for suspected leukemia since the presence of a leukocytosis at 73030 E / mm3 and thrombocytopenia at 14000 E / mm3.

Clinical examination on admission showed a conscious, non-pyretic infant with conjunctival hemorrhage and petechiae of the forehead. An estimated weight of 5kg (-2DS), a size of 50cm (-4DS) and a normal head circumference at 42cm. we also noticed facial dysmorphia (picture 1) with plagiocephaly, hypertelorism, low implanted ears, flattening of the base of the nose, radial club hands without associated anomalies of the fingers (picture 2).

Figure1: Facial dysmorphia

The cardiac auscultation showed the presence of a cardiac murmur in connection with a minimal inter-auricular communication estimated at 6.6mm. The radiography of the two hands (picture 3) showed bilateral radial agenesis. The rest of the malformation report showed minimal
hydrocephalus and the absence of an abnormality of the urinary tree and other skeletal anomalies.

**Figure 2: Radial club hand**

**Figure 3: Bilateral radial agenesis**

The biological assessment showed a hemoglobin at 6g / dl, VGM at 76fl, TCMH at 22pg, hyperleukocytosis at 73030 E / mm3 (lymphocytes at 32940 E / mm3, neutrophiles at 17240 E / mm3, monocytes at 20010 E / mm3) and a thrombocytopenia at 14000E / mm3. The myelogram showed the presence of 3% of blasts, of a discreet multiline dysmyelopoiesis with monocytosis. The case has been discussed by hematologists. It is a pre leukemic condition, which required a myelogram check in a month, but the mother left against medical advice.

Therapeutically, the infant was put on a blood and platelet pellet transfusion in the presence of hemorrhagic signs.

Regarding facial dysmorphism, bilateral radial agenesis and thrombocytopenia, we thought of thrombocytopenia of constitutional origin, notably the thrombocytopenia-radial agenesis syndrome. The genetic study was planned but the mother decided to go out against medical advice before genetic confirmation.

**3. DISCUSSION**

Thrombocytopenia absent radius syndrome was first described in 1951[2] and defined as a syndrome in 1969 [3] by Judith Hall. It is a very rare congenital malformation (<< 1 / 100,000 live births) with a sex ratio of 1. Its genetic transmission seems autosomal recessive, while an autosomal dominant transmission has also been proposed [4, 5].

Clinically, it is characterized by thrombocytopenia associated with absent rays; clinical sign allowing the diagnosis to be made at birth [3]. The presence of the thumb is pathognomonic. Other abnormalities of the skeletal and cardiac systems (23%) were present, most often a tetralogy of Fallot and septal atrial defects. Genitourinary anomalies including the duplex ureter and horseshoe kidney [3, 6-9] and brain abnormalities “hypoplasia of the cerebellar vermis and corpus callosum” have also been reported [10]. In case of our patient, we objectified as anomalies, bilateral radial agenesis, facial dysmorphism, inter-auricular communication and minimal hydrocephalus.

The number of platelets in infants is low, ranging from 10 X 10⁹ / L to 100X 10⁹ / L. The bone marrow reveals few megakaryocytes. In the case of our patient, we found severe thrombocytopenia and multiline dysmyelopoiesis with monocytosis, which may be related to a pre leukemic state. In the literature, 4 cases of leukemia have been reported in patients suffering from this syndrome [6-8].

The Thrombocytopenia disappears after childhood, although platelet counts may remain below average until adulthood [11]. The etiology and mode of transmission remain unclear. In all patients, a deletion of chromosome 1q21.1 has been found [12]. However, to this date, the cause and effect link between this deletion and the Thrombocytopenia absent radius syndrome remains unclear. In case of our patient, the karyotype and the genetic study were not done given the discharge of the patient against medical advice.

**4. CONCLUSION**

The Thrombocytopenia absent radius syndrome is a rare anomaly, which must be evoked in front of any dysmorphism associated with a thrombocytopenia. Mortality during the first year of life is linked to massive haemorrhage, particularly intracranial [5]. Most patients with this syndrome who survived childhood have a normal lifespan [5,9].
REFERENCES


