Henna-Induced Haemolysis in an Un-diagnosed G6PD Deficient Arabian Baby-Case Report

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Keywords: Henna; Lawsone; Haemolysis; G6PD deficiency

Introduction

Worldwide Glucose-6-Phosphate Dehydrogenase (G6PD) deficiency is the most prevalent enzymopathy that affects over 400 million individuals [1]. G6PD deficiency, prevalence varies by geographical location. It is higher in some parts of the world and low in others and differs among different ethnic groups like in Sub-Saharan Africans (7.5%), Middle East (6.0%) and Asians (4.7%) are affected more than the Americans (3.4%), Europeans (3.9%) and the Pacific Islanders (2.9%) [2].

Most people with G6PD deficiency, however, do not exhibit symptoms unless exposed to stressors like, infections, oxidative medicine and or fava beans. The main clinical manifestations of G6PDd are acute haemolytic anaemia, chronic non-spherocytic haemolytic anaemia, neonatal jaundice and favism [1,3].

Henna also called “Mehendi” is a traditional cosmetic agent, used worldwide especially in Asia, Middle East and Africa, not only for cosmetic purposes to stain the hairs, skin and nails in ceremonial and social events throughout the Asia, Middle East and Africa that contain an active dye ingredient lawsone. Like other ortho-substituted 1,4-naphthaquinone, lawsone induce oxidative injury to red blood cells, that could be fatal in G6PD deficient infants and rarely in children and adults. We report a previously undiagnosed G6PD deficient Arabian baby with life threatening haemolysis after topical application of henna.

Case Report

A 4-month old Arabian baby girl, previously undiagnosed with G6PD deficiency, who developed life threatening hemolysis after topical application of henna to her body.

Physical examination revealed an icteric baby with an axillary temperature of 36.5°C, pulse 110/min, respiratory rate of 28/min, blood pressure of 119/55 mmHg and oxygen saturation of 98% in room air. Other clinical findings were unremarkable except a short systolic murmur on left sternal border.

Laboratory work-up showed haemoglobin 6.4 g/dL, hematocrit 22.4%, white blood cell count of 23.50 x 103/ µL, red blood cell count 2.45 x 106/µL and platelet count 343 x 103/µL, MCV 82fL and RDW 13%.

Peripheral blood smear showed 27.1% neutrophils, 66.2% lymphocytes, anisocytosis (+), poikilocytosis (+), spherocytosis (-), Heinz body (+) and reticulocyte count 5.5%. Biochemical analysis yielded total serum bilirubin 13 mg/dL, indirect bilirubin 2.9 mg/dL, blood asparate aminotransferase 131 iu/L, serum urea 12 mg/dL, creatinine 0.36 mg/dL, and other biochemical parameters were within normal limits. Serology tests for HAV, HBV, and HCV were negative. Blood test for malaria, direct coombs test and blood culture was negative. Haemoglobin electrophoresis was within normal range. Urine analysis showed yellow colored urine, specific gravity: 1015, bilirubin (Nil), hemoglobinuria was (Nil). Quantitative G6PD enzyme deficiency of the infant was established and it was low, 104 mU/10⁹ RBCs (Ref. Range 166–445).

A presumptive diagnosis of henna induced acute haemolysis was made, as other causes of haemolysis like infection and exposure to chemical agents were ruled out, which then came out as G6PD deficient by quantitative analysis.

The patient neither developed renal impairment nor Disseminated Intravascular Coagulation (DIC) and had uneventful recovery after blood transfusion and supportive treatment including adequate hydration and the patient symptoms and laboratory data was recovered. The infant progress was uneventful and discharged on 7th day of admission, with follow-up a week later in good health while the parents were counselled regarding the condition with absolute avoidance of henna and given a list of drugs and food to be avoided.
Discussion

Henna obtained from the powdered leaves of a shrub that are used not only in religio-cultural events, widely in Asia, Middle East and Africa but also for therapeutic purposes to treat various skin infections. Lawsonia that constitute about 1% of the powdered form of henna leaves is a potent oxidant of G6PD deficient red cells like 1,4 naphthoquinone-a naphthalene metabolite [7,8].

Despite the potential risk of henna induced haemolysis, there are few reported cases of life threatening hemolysis in G6PD-deficient patients due to topical application of henna. The clinical symptoms severity depends on the degree of enzyme deficiency although other factors such as oxidative agent dose, exposure time, patient age, haemoglobin level and concurrent infection do contribute [1].

In the study of Kandel et al. [9] 15 G6PD deficient new borns were reported that were admitted with haemolysis after 24-72 hours of topical henna application. Raupp et al. [10] reported 4 cases of acute hemolysis in G6PD deficient children post-henna application.

In another study Syeyedzadeh et al. [11] reported a 42-days old G6PD-deficient infant who developed acute hemolysis after topical application of henna to treat his napkin dermatitis. Kheri A et al. [12] reported life-threatening haemolysis in a 6-year old boy with undiagnosed G6PD deficiency after topical application of henna, while Soker et al. [7] reported an eleven years old G6PD deficient boy, with henna induced haemolysis to treat his psoriatic skin lesions.

In a recent systemic review, Lee et al. [13] reported 20 cases of hemolysis in G6PD deficient patients due to henna application. Taken together, the available data indicates that henna could increase the risk of hemolysis in G6PD deficient individuals. Our case is the first reported case of henna induced severe haemolysis in a previously undiagnosed G6PD deficient patient, in our paediatric unit of tertiary care hospital despite the high prevalence of G6PD deficiency and popularity of henna in the Arab culture.

The report of this case highlights the risk of life threatening risk of henna to cause acute haemolysis in G6PD deficient patients in specific and undiagnosed cases in general. Thus, henna application should be discouraged generally in unscreened infants and specifically individuals of any age group with a family history or known cases of G6PD deficiency.

Consent

Written informed consent was obtained from the patient’s guardian for publication of this case.

Conflict of interest

The authors declare no conflict of interest.

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