Imerslund-Grasbeck Syndrome: Association with Diabetes Mellitus

S Madhavan, M Vijaykumar*, Sarala Rajajee† and BR Nammalwar*

From the Departments of *Pediatrics, Pediatric Nephrology, and †Pediatric Hematology, †Kanchi Kamakoti CHILDs Trust Hospital, Chennai, India.

CASE REPORT

A 14-year-old male adolescent born of 2nd degree consanguineous marriage presented with asymptomatic proteinuria and severe anemia. He had leucopenia, anisopoikilocytosis, megaloblastic erythropoiesis, megakaryocytes with low serum B$_{12}$ level. His younger sibling was similarly affected. This combination suggested Imerslund-Grasbeck syndrome. The hemoglobin levels improved with injection of vitamin B$_{12}$ but proteinuria persisted. During follow-up, he developed ketoacidosis due to insulin-dependent diabetes mellitus. This rare combination has not been reported in the Indian literature.

Key words: Imerslund-Grasbeck syndrome, Diabetes mellitus, Vitamin B$_{12}$

Correspondence to:
Dr M Vijaykumar,
Flat No 4, Muktha Vandan,
New No 7, Old No 4,
Ramanathan Street, Kilpauk,
Chennai, India.
E-mail: doctormvk@gmail.com

Imerslund-Grasbeck syndrome (IGS) is a rare autosomal recessive disease characterized by vitamin B$_{12}$ (cobalamin) deficiency due to selective malabsorption of this vitamin resulting in megaloblastic anemia appearing in later childhood. The anemia responds well to parenteral vitamin B$_{12}$ therapy. This syndrome is frequently accompanied by proteinuria and sometimes neurological symptoms(1). Associated genitourinary tract abnormalities(2), dolichocephaly(3), beta-thalassemia trait(4) and diabetes mellitus(5) have been reported. We present a case of IGS with diabetes mellitus.

CASE REPORT

A 11-year-old boy, first born to second degree consanguineous parents was referred for persistent asymptomatic proteinuria since one year. There was no hematuria, oliguria, skin rash, joint pain or swelling. At the age of 2 years, during a diarrheal illness, he had periorbital puffiness, severe anemia, hyper-pigmentation of the hands and 2+ proteinuria. With vitamin B$_{12}$ injection and oral medications, he had improved.

On examination, his growth and development were normal and his weight was 36 kg. He did not have anemia or pedal edema. He had hyperpigmentation of the skin over the dorsum of the fingers and toes. Blood pressure was normal. Cardiovascular and respiratory system were clinically normal and he had no ascites or organomegaly. Investigations showed proteinuria of 2+ by dipstick. Blood counts and peripheral smear was normal, hemoglobin was 11.5 g/dL. Renal function tests were normal. Serum proteins and cholesterol was normal. 24-hours urine protein excretion was 294 mg/day. Ultrasonogram of the abdomen was normal. Serum ANA was negative. Renal biopsy was reported as minimal lesion nephropathy by light microscopy and immunofluorescence showed no immune deposits. Enalapril and losartan potassium were given for renoprotective and antiproteinuric effects.

He remained asymptomatic for 3 years but had persistent non-nephrotic proteinuria with normal hematological values. His weight increased to 44 kg. At 14 years, he was hospitalized for fever with vomiting. On this occasion, he had severe anemia,
facial puffiness and loss of weight to 37 kg (25th percentile). His height was 158 cm (50th percentile). There was no neurological deficit, optic atrophy, or hearing defect. Blood pressure was normal. Investigations revealed severe anemia with leucopenia and thrombocytopenia; hemoglobin was 4.9 g/dL. There were macrocytes, hypersegmented neutrophils and anisopoikilocytosis in the peripheral smear. Bone marrow revealed megaloblastic erythropoiesis with occasional megakaryocytes. Serum LDH levels were elevated. In view of megaloblastic anemia, persistent proteinuria and absence of chronic diarrhea, parasitic infestations or obvious nutritional deficiency, the possibility of IGS was considered and serum vitamin B12 levels were estimated. This was low (vitamin B12 97 [211-911 pg/mL]). Injection of vitamin B12 was given with advice for follow-up B12 injections. At follow-up, his general condition was good with an increase in weight to 39 kg. Hemoglobin increased to 11.9 g/dL. Three months later, he was admitted for severe diabetic ketoacidosis (fasting urine sugar 4+, fasting plasma glucose 421 mg/dL, post prandial plasma glucose 590 mg/dL). With human insulin and four monthly injections of Vitamin B12 he remained asymptomatic with a weight gain to 42 kg.

His ten years old younger sibling, who was asymptomatic but for hyperpigmentation of the dorsum of the fingers, was administered 2 monthly injection of B12 by his pediatrician in view of the elder sibling’s diagnosis. The hemogram done subsequently was normal with non-nephrotic proteinuria. His vitamin B12 level was normal (444 [211-911 pg/mL]) and glucose tolerance test was normal. We plan to stop vitamin B12 injections for him and recheck his serum B12 values after 6 months.

DISCUSSION

In the absence of history of exposure to nitrous oxide, vegetarian diet, parasitic infestation or chronic gastroenterstinal malabsorption syndromes, the cause of macrocytic and megaloblastic anemia is likely to be due to an inherited disorder of folate or vitamin B12. In clinical practice, diagnosis can be nearly reached by observing the response to treatment with oral dose of folate or injection vitamin B12. The response in this child favours the diagnosis of a B12 deficiency state. The presence of proteinuria favors the diagnosis of IGS, which is present in about 70% of the patients.

Imerslund-Grasbeck syndrome was first described in Finland and Norway where the prevalence is about 1:200,000. The clinical features include megaloblastic anemia, failure to thrive, recurrent infections, neurological manifestations and asymptomatic proteinuria which is neither glomerular nor tubular with no signs of kidney disease. These symptoms may manifest at any time between 1 to 15 years of age with the mean age being 3.5 years.

The selective malabsorption of vitamin B12 and proteinuria involves a mutation in one of two genes, cubulin (CUBN) on chromosome 10 or amnionless (AMN) on chromosome 14. Both proteins are components of the ileal enterocyte receptor for the vitamin B12 intrinsic factor complex and the receptor mediating the tubular reabsorption of protein from urine. Life long treatment with injection B12 leads to complete recovery. Proteinuria persists with normal renal function. Schroder, et al. (5), reported two siblings with IGS and in one of them type 1 diabetes mellitus (DM) preceded the diagnosis of IGS by 7 years(5). Our child developed DM with ketoacidosis later on. This presentation is because of the rare association and unexplainable common etiological factor.

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