Mauriac Syndrome: A Rare Complication of Poorly Controlled Diabetes Mellitus

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Abstract

Mauriac syndrome is one of the complications of poorly controlled diabetes mellitus type 1. Common presenting features include short stature, growth retardation, moon facies, protuberant abdomen, and proximal muscle wasting. We hereby report a case of 15 year old female who presented with clinical features of Mauriac syndrome.

Keywords: Mauriac syndrome; Diabetes mellitus; Complication

Introduction

Poorly controlled diabetes is a frequent problem in developing countries leading to many complications related to underinsulisation and Mauriac syndrome is one of them. Mauriac syndrome is a complication in diabetes mellitus type 1, which includes short stature, glycogen laden enlarged liver, limited joint mobility, tight waxy skin, growth maturation delay, moon facies, protuberant abdomen, and proximal muscle wasting and also it is frequently associated with retinopathy and nephropathy [1, 2]. We hereby report a case of mauriac syndrome as very few cases have been reported previously in India [3].

Case Report

A 15-year-old female, referred to us for evaluation of short stature and delayed puberty. Patient also had decreased vision and tingling sensation in the upper and lower limbs since one month. She was diagnosed to have type 1 diabetes mellitus (DM) 10 year back, and was on subcutaneous premixed insulin. However, compliance as well as sugar monitoring was very poor. Therefore, she had been repeatedly hospitalized with frequent manipulations in the insulin dosing and schedule. On examination, patient was short statured (102 cm, less then 3rd percentile), her weight for age was less then 3rd percentile, BMI was 14 (between 5 - 10th percentile), had protruded abdomen with hepatosplenomegaly, muscle wasting and delayed puberty (SMR stage 1). She was hypertensive, had ketonuria (+++) and glycosuria. Investigations showed normal hemogram, normal liver and renal function test. Ultrasound abdomen revealed hepatosplenomegaly with altered liver echotexture and fatty infiltration of the liver. Her premeal blood sugar was 420 mg%, HBA1C was 13.1% (normal < 6%), microalbuminuria > 160 μg/mL (normal < 18 μg/mL) with higher albumin/creatinine ratio. Patient also found to have motor axonal neuropathy on nerve conduction velocity, microaneurysm on fundus examination and osteopenia on skeletogram. Blood gas analysis suggested no acid base imbalance, and no electrolyte abnormality. Thyroid function tests were within normal range. Based on history, examination and investigation findings, final diagnosis of Mauriac syndrome was made. Patient was shifted to strict dietary management and high dose premixed insulin, high dose calcium, vitamin D3 and multivitamin. After 2 months of therapy, patient improved symptomatically, premeal sugar came down to 100 - 150 mg% and there was reduction in hepatomegaly.

Discussion

Mauriac syndrome was first described by Mauriac in 1930 in children with type 1 DM presenting with clinical features of growth failure, maturation delay, hepatomegaly and abdominal distension [4]. Most of the cases occur in adolescence with equal sex ratio. On the basis of the presence or absence of obesity, 2 different forms of Mauriac syndrome have been described. Pathogenesis of the growth retardation is not clear but thought to be multifactorial. Inadequate glucose to the
tissues, decreased insulin like growth factor 1 and growth hormone level, and hypercortisolism may contribute to delayed growth and puberty. Hepatomegaly is thought to be due to glycogen deposition in the liver [5, 6]. Growth failure, delayed puberty and hepatomegaly in Mauriac syndrome improve with glycemic control [3]. This is a case of adolescent female with poorly controlled DM who found to have short stature. With the additional findings of gross hepatomegaly, delayed puberty and a Cushingoid habitus, she was diagnosed with Mauriac syndrome. In this case, poor compliance because of poor literacy was the main cause behind the poorly controlled diabetes.

Disclaimer

None.

Competing Interest

None.

References