Liver Transplantation in a Myopathic Patient with Glycogen Storage Disease Type IIIa and Decompensated Cirrhosis

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ABSTRACT

Glycogen storage disease (GSD) type IIIa (Forbes-Cori disease) can be associated with severe liver disease. A patient with GSD type IIIa may therefore be a potential candidate for liver transplantation. Progressive myopathy makes uncertain the outcome of the patient and the transplant. Herein, we report on the good results of liver transplantation up to 28 months after the transplantation in a 40-year-old man with liver cirrhosis and significant muscle weakness due to GSD type IIIa.

KEYWORDS: Cirrhosis, Glycogen storage disease types III; Liver transplantation; Liver diseases; Liver cirrhosis

INTRODUCTION

Glycogen storage disease type III (GSD III) is a rare autosomal recessive hepatic disease characterized by glycogen accumulation due to defect in the gene AGL with deficiency of glycogen debranching enzyme activity that converts glycogen to glucose [1]. Glycogen accumulation occurs primarily in the liver and in skeletal and cardiac muscles with systemic consequences [2]. GSD III accounts for approximately 24% of GSDs with an estimated incidence of 1 case per 83,000 live births in Europe and 1 in 100,000 in North America [2, 3]. Four subtypes of GSD III, based on differences in tissue expression of the deficient enzyme, are recognized [4]. Relatively, 85% of GSD III is GSD IIIa subtype (Forbes-Cori disease) that involves both liver and muscle; 15% are of type IIIb, which have only liver involvement [5]. Two other subtypes, GSD IIIc and GSD IIIId, are extremely rare [5]. Faroe Islands have the highest prevalence of GSD IIIa [6]. This relatively mild form of GSD presents in childhood and has a good prognosis [7]. Typical features include hepatomegaly that gets smaller with time, hypoglycemia, poor growth with eventual catch-up, skeletal myopathy that gradually worsens with time, and cardiomyopathy [7]. Hepatomegaly, hypoglycemia, hyperlipidemia, and growth retardation usually improve with age and disappear after puberty [4]. Muscle weakness in GSD IIIa is minimal during childhood; however, it can become predominant in adulthood and show signs of neuromuscular involvement with slowly progressive weakness and distal muscle wasting [6]. Long-term complications in adults mainly related to the muscles, heart and liver involvement. Hypertrophic cardiomyopathy usually develops during childhood in the majority of patients with GSD IIIa, but the seriousness of signs and symptoms varies from asymptomatic in the majority to severe heart disease and rarely heart failure [7]. Liver cirrhosis is a rare complication of GSD III [8]. There is no specific treatment for GSDs, but diet therapy with high-protein, frequent meals with corn starch supplements or nocturnal gastric drip feeding, constitutes effective therapy and im-
proves hypoglycemia, liver size, and overall
growth and development [2]. The detection
of the molecular defect has facilitated the di-
agnosis and has offered the opportunity for
prenatal diagnosis in these patients [9]. Cur-
currently, there is no effective treatment for the
progressive myopathy or cardiomyopathy [4].
Diabetes mellitus (DM) in GSD III is difficult
to treat because patients are prone to hypogly-
cemia [5]. Herein, we report on a case of GSD
IIIa presenting with great skeletal myopathy
and end-stage cirrhosis without hypoglyce-
mia and cardiomyopathy that underwent liver
transplantation (LT).

CASE REPORT

A 40-year-old man presented with gastroin-
testinal bleeding due to esophageal varices
and liver cirrhosis. Review of his records re-
vealed that he had been diagnosed with GSD
at the age of two years following a liver biopsy,
when he had a distended abdomen due to hepa-
tomegaly. He had problem with weakness from
early childhood. In his childhood, he had fre-
quent nose bleeding and easy fatigability. His
growth was very slow. He was on no specific
therapy and never suffered from symptomatic
hypoglycemia. He went into a late puberty at
about 17–18 years, but grew very rapidly and
became much taller than his parents. During
that period, his muscle weakness worsened
and he had been in a wheelchair since he be-
came 24 years old. His weakness in his arms
also dated back to his puberty. At the time of
presentation, he was unable to brush his hair
and had to lean forward to eat. He had not had
any heart problem. He had not had any sleep
problems or chest infections. He suffered from
low back pain after sleeping. His diet was high
in protein with lots of meat, nuts, eggs, and
yoghurt. He had never tried uncooked corn
starch. GSD IIIa, for the first time, was con-
firmed at the age of 28 in the National Hos-
pital for Neurology and Neurosurgery, Queen
Square in London, on April 7, 2004. His par-
ents were unrelated, alive and doing well.
There was no family history of other muscle
disorders. He had no other medical problems.
He had normal schooling and studied engi-
neering at university but had to stop in his
3rd year because of the weakness. He was on
no regular medication. He was unable to sit-
ing in wheelchair and needed assistance to
be transferred. On physical exam, he appeared
dysmorphic, thin, disable but intelligent. His
vital signs, chest and heart were normal with-
out any evidence of cardiomyopathy. Abdomi-
nal examination revealed a firm and enlarged
liver, splenomegaly, and ascites. Marked prox-
imal and distal weakness, not involving face,
was detected. The patient was unable to lift
his arms or legs against the gravity. No deep
tendon reflexes were presented. Muscular
enzymes were elevated; muscle biopsy per-
formed at age of 30 years. Because of massive
upper gastrointestinal bleeding due to grade 4
esophageal varices, ascites, low platelet count,
and fatigue, despite severe muscle weakness
with weight, height and BMI of 56 kg, 180
cm, and 17.2 kg/m², LT was performed. On
28th month after transplantation, the patient
had normal liver function and metabolic con-
trol without any restrictions and some weight
gain up to 60 kg. However, he still had muscle
weakness with elevated muscle enzymes. He
was treated with prednisone 5 mg, mycophe-
nolate 1500 mg, and tacrolimus 4 mg for pro-
phylaxis of graft rejection.

DISCUSSION

This is the first report of patient with GSD
IIIa who underwent LT in the middle-age, be-
cause of cirrhosis and variceal hemorrhage as
an index complication despite loss of muscle
mass and strength. The first report on LT in
GSD III performed in 1989 [10]. GSD IIIa in-
volves muscle and muscle weakness is mostly
minimal at childhood, however, it progresses
slowly and becomes prominent during the
third and fourth decades of life [11, 12]. Proxi-
mal muscles are primarily involved, but distal
muscle wasting including the calves, hands,
and peroneal muscles are also involved [11, 12].
Liver fibrosis is also a common feature of
GSD III, and micronodular cirrhosis in some
cases, has been found but appears not to pro-
gress in most instances. Hepatic adenoma or
hepatocellular carcinoma has been reported
in some cases [13]. LT is indicated when end-stage cirrhosis or hepatic malignancy is detected, but it has also been used with success in patients who are recalcitrant to medical therapy [14]. Ultimately, GSD IIIa is a multisystem disorder, and long-term success of LT, particularly with regard to myopathy or cardiomyopathy, is not known [15]. LT may adversely affect the symptoms of myopathy and cardiomyopathy [16].

Matern and colleagues demonstrated post-transplantation correction of metabolic abnormalities [5]. LT is a life-saving measure for patients with chronic end-stage liver disease and supposed to correct the primary hepatic enzyme defects and the deleterious complications of GSD with acceptable outcome and good long-term results [14, 17]. The extrahepatic manifestations of GSD often complicate post-transplantation period [18]. LT is contraindicated for patients who are unlikely to survive the procedure or receive long-term benefits. Patients are considered individually and their candidacy is assessed by a formal multidisciplinary evaluation process [19]. LT does not cure the heart and muscle problems; transplantation has been associated with worsening myopathy and cardiomyopathy [20]. No systematic study has analyzed the risk factors for neuromuscular complications in LT. High doses of corticosteroids and the use of non-depolarizing neuromuscular blocking agents are reported to favor quadriplegia after LT [21]. Use of prednisone with dose of <10 mg/day is rarely associated with glucocorticoid-induced myopathy [22]. Therefore, discontinuation of prednisone probably has little effect on muscle weakness.

CONFLICTS OF INTEREST: None declared.

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