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Short Communication

Early care of N-acetyl glutamate synthase (NAGS) deficiency in three infants from an inbred family

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ABSTRACT

N-acetyl glutamate synthase (NAGS) deficiency is the rarest urea cycle defect presenting as neonatal onset life-threatening hyperammonemia. We report here a family history of severe NAGS deficiency: after the index-case with severe hyperammonemia, one patient benefited from antenatal diagnosis, and from primary care at birth, another one was diagnosed at 2-days and immediately treated with carbaglumic-acid. Finally, we report excellent tolerance to long-term carbaglumic-acid treatment, with no side effects, and healthy neurological and psychomotor development.

1. Introduction

N-acetyl glutamate synthase (NAGS) deficiency is an autosomal recessive disease that usually presents as neonatal onset life-threatening hyperammonemia. NAGS catalyzes the formation of N-acetyl glutamate by the combination of glutamate and N-acetyl CoA. This molecule activates, in an allosteric manner, carbamoylphosphate synthetase I (CPSI), which is the first and rate-limiting enzyme of the urea cycle. This enzyme combines ammonia with bicarbonate to produce carbamoyl-phosphate. High plasma ammonia, glutamine, and glutamate, and barely detectable citrulline concentrations characterize NAGS deficiency [1] [2] [3] [4].

Severe NAGS deficiencies entailing the complete lack of the enzyme lead to severe neonatal hyperammonemia; late-onset presentations related to a partial lack of the enzyme have been reported in infancy, childhood, and adulthood [7]. All patients present a physiological adaptation to extra-uterine life, and neurological deterioration appears within a few hours to a few months. NAGS deficiency is the only urea cycle disorder that is curable. This disease is currently treatable using carbaglumic acid (Carbaglu*), an analog of N-acetyl glutamate, and a low protein diet when initiating the treatment [5]. Management of hyperammonemia is challenging in the acute phase.

We report here a family with severe NAGS deficiency within an extended inbred family. We identified three patients with a complete deficiency (homozygous for a null allele), one patient heterozygous for this variant, and numerous miscarriages and fetal deaths, probably related to the disease. This report illustrates the overall picture of the disease, its management, and the value of early prenatal or pre-symptomatic diagnosis.

2. Patients and methods

We describe infants from a sizeable inbred family of Afghan descent (Fig. 1). All fathers were brothers, and the two mothers were sisters and first cousins of two of the fathers. We divided the whole family into two branches, each one for mother.
3. Reports and results

3.1. The first branch of the family

The mother of the index case reported one neonatal death in 2004 (IV-1), a miscarriage in 2005 (IV-2), and gave birth to a healthy boy (IV-3) in 2005. She then married a second brother among the siblings. She gave birth in 2013 to a baby girl (IV-4) (Fig. 1; Proband), at 39 weeks' gestation after pregnancy without any particular complications. The child's birth weight was 2770 g. At the age of 2 days, the neonate presented axial and peripheral hypotonia, weight loss of more than 10%, with alkalosis, hyperlactatemia, and major hyperammonemia (676 μmol/L; Table 1). Dialysis was immediately proposed, and she was treated with sodium phenylacetate and sodium benzoate at 250 mg/kg/day (Table1). After 24 h, ammonia fell below 200 μmol/L, treatment was adjusted (sodium benzoate 200 mg/kg/day), and Carbaglu® treatment was initiated at 100 mg/kg/day, with carnitine, arginine, and B12 vitamin supplementation. Continuous enteral nutrition without protein, with glucose and lipids, was implemented (Table1). At day-5, ammonia blood concentration normalized, and dietary proteins were progressively included in her regimen to 5 g per day. The plasma amino acid profile revealed a urea cycle disorder (Table 1), and urinary organic acid chromatography showed the absence of orotic acid. NAGS deficiency was then suspected, and the patient was found to be homozygous for the p. Thr439Hisfs*52 (NM_153006.2: c.1313dupG) NAGS variant, which was previously described [6]. MRI showed a discrete T2 hyper signal on palladium, and punctiform hyper signals on the periventricular white matter when the child reached the age of 7 months. She experienced a single decomposition episode and hospitalization for varicella at one year of age. Now aged six years, despite the massive hyperammonemia at birth, she presents good stature and weight and normal neurodevelopment. She receives a normal protein intake at 3.5 g/kg/day and Carbaglu® at 50 mg/kg/day.

3.2. The second branch of the family

In the second branch of the family, the parents are also consanguineous (first cousins): the father was another of the siblings mentioned above, and the mother was the sister of the index case's mother. These parents reported a baby girl who died abroad on day-12 from hypotonia and coma (IV-7; 2014). Her brother (IV-8) was born in Iran in 2015 by a cesarean section (birth weight 3130 g). He presented mild hyperammonemia at six months in a context of severe fatigue and frequent crying (Table 1). Because he was heterozygous for the p. Thr439Hifs*s*52 variant of NAGS, we supposed his mild hyperammonemia was likely due to difficult blood sampling or delay in processing the sample. He is now aged five years and exhibits good stature, weight, and neurological development and has no specific treatment or diet.

Two years later, a baby girl was born at 39 weeks' gestation (birth weight 2844 g; IV-9) with a physiological adaptation to extra-uterine life. No prenatal diagnosis was performed because the parents initially failed to mention the family history to the physician until birth. At 24 h of life, she presented moderate hyperammonemia without alkalosis or clinical signs (Table 1). She was immediately treated with Carbaglu®, sodium benzoate, arginine and low protein diet (Table1). Before treatment, amino-acid chromatography was performed and presented abnormalities of urea cycle disorders (Table 1). The genetic diagnosis revealed homozygosity for the p.(Thr439Hisfs*52) variant of NAGS. She was treated with Carbaglu® (50 mg/kg/day), and arginine supplementation (58 mg/kg/day), and a low-protein regimen (2 g/kg/day) in the early stages of treatment, which was subsequently adjusted according to age and development. She presented an acute episode of hyperammonemia (153 μmol/L) at 16 months, during acute infectious disease. Now aged 20 months, she exhibits good stature, weight, and neuro-developmental evolution with no protein restriction and is...
4. Discussion

We present here several patients with various features of NAGS deficiency, the rarest urea cycle disorder, with an estimated incidence of 1:3,500,000–7,000,000 [3]. In our description, despite major hyperammonemia, the index case presented a favorable long-term outcome, mainly as a result of early care and treatment with N-carbamoyl glutamate. Her sister had the benefit of a prenatal diagnosis enabling management at birth to avoid hyperammonemia, without any neurological damage. To our knowledge, we report here the first prenatal molecular diagnosis that enabled successful management from birth.

The three homozygous patients reported very few decompensations affected by NAGS deficiency: novel mutations of the N-Acetylglutamate synthase family: structures, function and mechanisms, Int. J. Mol. Sci. 16 (2015) 13004–13022.

The authors have no disclosures to declare.

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