SHORT COMMUNICATION

Nineteen-year follow-up of a patient with severe glutathione synthetase deficiency

Paldeep S Atwal1,2, Casey R Medina1, Lindsay C Burrage1 and V Reid Sutton1

Glutathione synthetase deficiency is a rare autosomal recessive disorder resulting in low levels of glutathione and an increased susceptibility to oxidative stress. Patients with glutathione synthetase deficiency typically present in the neonatal period with hemolytic anemia, metabolic acidosis and neurological impairment. Lifelong treatment with antioxidants has been recommended in an attempt to prevent morbidity and mortality associated with the disorder. Here, we present a 19-year-old female who was diagnosed with glutathione synthetase deficiency shortly after birth and who has been closely followed in our metabolic clinic. Despite an initial severe presentation, she has had normal intellectual development and few complications of her disorder with a treatment regimen that includes polycitra (citric acid, potassium citrate and sodium citrate), vitamin C, vitamin E and selenium.

INTRODUCTION

Glutathione (GSH) is a ubiquitous tripeptide (γ-glutamylγ-cysteinylglycine) known to function in many aspects of cellular activity including protein and DNA synthesis, transport, detoxification of xenobiotics and carcinogens, metabolism, and defense against free radicals and oxidative stress.1 It is synthesized in the α-glutamyl cycle (Figure 1) by GSH synthetase (GS). GSH is most commonly found in its reduced form with only 1–5% existing in the oxidized form GSH-disulphide. GSH-disulphide is reduced back to GSH by GSH-disulphide reductase requiring one equivalent of NADPH.2 Biallelic, pathogenic variants in the GSS gene result in GS deficiency (GSSD).

GSSD (MIM:266130) is a rare autosomal recessive disorder that has only been described in approximately 70 individuals worldwide.3 Erythrocytes have some of the highest concentrations of GSH in the body, and thus, erythrocytes from patients with GSSD are more susceptible to oxidative stress. As a result, hemolytic anemia is a common problem in GSSD.4 Depending on their clinical manifestations, patients can be divided into mild, moderate or severe phenotypes. Patients with the mild GSSD present with isolated hemolytic anemia as their only clinical symptom, whereas patients with the moderate form of the disorder present with hemolytic anemia and metabolic acidosis in the neonatal period. In addition to these symptoms, patients with severe disease have neurologic findings including motor disturbances and developmental delay.5 Some severely affected patients show an increased susceptibility to bacterial infections, thought to be due to defective granulocyte function. To date, there is only one severely affected individual who was treated with antioxidants from birth and long-term follow-up on that individual is not reported. 5-Oxoprolinuria can also be found in all patients with GSSD due to the accumulation of γ-glutamylcysteine being hydrolyzed to 5-oxoproline and cysteine by γ-glutamylcyclotransferase. This accumulation of 5-oxoproline exceeds the capacity of 5-oxoprolinase leading to high urinary excretion of 5-oxoproline that is detectable on urine organic acid analysis.6 In the present report, we describe the long-term follow-up of a case of severe GSSD.

CASE PRESENTATION

The proband is a female born at 40 weeks gestation by Cesarean section weighing 2.970 kg (16th centile) with Apgar scores of 8 and 9 at 1 and 5 min, respectively. At approximately 18 h of life, she was found to be tachypneic with a metabolic acidosis. Arterial blood gas showed pH of 7.32 (normal pH 7.35–7.45), PaCO2 17 mm Hg (normal 35–45 mm Hg), PaO2 94 mm Hg (normal 80–100 mm Hg) and bicarbonate of 8 mmol l−1 (normal 21–32 mmol l−1). She was given 5 mg intravenous (IV) sodium bicarbonate and transferred to a tertiary hospital at which time her metabolic acidosis had worsened to pH 7.15, PaCO2 12 mm Hg, PaO2 94 mm Hg and bicarbonate of 4 mmol l−1. She was treated with IV fluids and IV sodium bicarbonate of 1.1 meq kg−1 h−1. Urine organic acid analysis performed at this time revealed high levels of 5-oxoproline, suggesting a defect in GSH metabolism. Once stabilized, she was started on a combination of citric acid, potassium citrate and sodium citrate (~30 meq kg−1 per day) for metabolic acidosis and vitamin C (250 mg per day) and E (30 international units per day) to help prevent oxidative stress. Her mother was also advised to avoid foods and medications that cause oxidative stress (from a list standardly used for individuals with glucose-6-phosphate dehydrogenase deficiency). She was then discharged home with follow-up in the metabolic clinic.
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At her 2-month follow-up visit at the metabolic clinic, she had remained stable with no intercurrent illnesses or hospitalizations since her discharge. She was tolerating her medications and growth was satisfactory (40th centile for weight, 10th centile for height). Physical exam was normal. Laboratory results showed mild macrocytic anemia with reticulocytosis and hyperkalemia, which became her baseline for most of her life, and a normal bicarbonate of 23 mEq L⁻¹. At 6 months, her red blood cell GSH and GS activity levels were analyzed, and she had a marked decrease in both GSH level and GS activity (see Table 1), consistent with the diagnosis of GSSD. Her mother also had a decrease in GS activity, although normal GSH levels, suggesting heterozygosity.

Throughout early childhood, she had multiple episodes of otitis media, which resolved with antibiotics and multiple minor viral respiratory infections. She also developed an episode of tachypnea and spitting up. Her serum bicarbonate level was reduced with antibiotics and multiple minor viral infections. She also developed an episode of tachypnea and spitting up. Her serum bicarbonate level was normal. At the age of 5 years, she was hospitalized. Her acidosis improved with IV fluids and IV bicarbonate. She was hospitalized again at 18 years of age when a viral gastroenteritis caused decreased oral intake and inability to take medications; her serum bicarbonate level at that time was 7 mEq L⁻¹ and resolved quickly with hydration and IV bicarbonate. Numerous dental caries were reported, and her dental health improved after a switch to a sugar-free combination of citric acid, potassium citrate and sodium citrate.

At approximately 2 years 3 months of age, there was concern that increased oxidative stress might hinder her developmental progress, so she was started on selenium—one crushed tablet daily. A formal developmental evaluation was performed at approximately 2 years 6 months and was reported as normal. However, by 3 years 9 months, developmental delays were evident. She was unable to copy a circle or ride a tricycle and her language was unintelligible. She was also having difficulties with toilet training. Speech therapy led to significant improvement, and by 4.5 years, her speech had greatly improved and her other developmental concerns resolved.

At her 33 months of age, weight loss and deceleration of head growth were noted, and these findings were attributed to chronic metabolic acidosis (bicarbonate of 15–18 mEq L⁻¹), her citric acid, potassium citrate and sodium citrate was adjusted to ~18 mEq kg⁻¹ per day and by three years of age, her acidosis and weight gain had improved.

Ophthalmology evaluation for pigmentary retinopathy, a known finding in patients with GSSD, was normal. At the age of 5 years, she underwent a tonsillectomy and adenoidectomy for chronic snoring and mouth breathing without any complications. She was diagnosed with attention deficit disorder at approximately 7 years of age and was successfully treated with guanfacine. At approximately 10 years, she was diagnosed with a seizure disorder and treated with the anti-epileptic lamotrigine. However, by the age of 15 years, she had no additional seizures, and her electroencephalogram was normal, so she was successfully weaned off lamotrigine. At the age of 16 years, her psychologist diagnosed her with Asperger syndrome. We are unaware of any association with Asperger syndrome with GSSD.

Currently, at 19 years, she is doing well. She graduated from mainstream high school and attends community college. She is currently prescribed guanfacine for attention deficit hyperactivity disorder, citric acid, potassium citrate and sodium citrate syrup (11 mEq kg⁻¹ per day) for metabolic acidosis, and vitamin C (14.5 mg kg⁻¹ per day), vitamin E (150 international units per day) and selenium (50 μg per day) to prevent oxidative stress. Her bicarbonate levels and reticulocyte counts are provided in Figure 2.

DISCUSSION

GSSD is a metabolic disorder that requires lifelong treatment. Treatment depends on the signs and symptoms of the patient. However, high doses of vitamins C and E are recommended in all patients for protection against oxidative stress. Avoidance of foods and drugs known to cause hemolytic crisis in glucose-6-phosphate dehydrogenase deficiency is also important as these same triggers can cause hemolytic crisis in GSSD. Vitamin E is also used to prevent granulocyte dysfunction, which could cause recurrent infections. Patients with moderate and severe phenotypes typically require treatment for metabolic acidosis. In acute crisis, IV bicarbonate is given for immediate correction, and for long-term management, citrate or trometamol are given. Our patient has required very high doses of citric acid, potassium citrate and sodium citrate (ranging from ~30 mEq kg⁻¹ per day in early infancy to approximately 10–20 mEq kg⁻¹ per day throughout most of her childhood and adolescent years) to maintain a normal serum bicarbonate level (Figure 2). Moreover, her persistently elevated reticulocyte count (Figure 2) reflects ongoing but compensated hemolytic anemia.

Previously, cysteine delivery compounds, specifically N-acetylcysteine, were given because they increase GSH levels in healthy patients. However, N-acetylcysteine has been found to increase intracellular levels of cysteine, which are already elevated in GSSD and these high levels of cysteine are known to be neurotoxic. Selenium was another agent used in our patient to prevent oxidative stress, and to our knowledge, this is the first reported use of selenium in a patient with GSSD. Selenium is found to have strong antioxidant properties through formation of selenoproteins, which are thought to protect against reactive oxygen species.
Early diagnosis and treatment is thought to correlate with a better long-term outcome. If a neonate presents with hemolytic anemia and metabolic acidosis, it is important to consider GSSD. Advanced diagnostic techniques such as antenatal diagnosis can be made by measuring 5-oxoproline in amniotic fluid or a presumptive diagnosis can be made by detecting the elevation of 5-oxoproline in newborn screen blood spots using tandem mass spectrometry. Our patient’s diagnosis was made within a few days of life, based on urine organic acid analysis and thus, treatment with vitamin C and E was started early. Njalsson et al. demonstrated that early initiation of vitamins C and E could prevent the moderate phenotype from progressing to severe phenotype. In a study of 41 patients with GSSD, only 1/18 of the severely affected patients were started on vitamin therapy early in life as compared with 6/17 moderately affected patients. With these data and the fact that there is no significant difference between enzymatic activity in the moderate and severe phenotypes, they posit that early initiation of vitamins could prevent or slow down progression of the disease. There were no specific comments on length of follow-up of this cohort however.

Our patient was diagnosed at birth with moderate disease because of her acidosis and hemolytic anemia. However, at the age of 10 years, she developed seizures, a finding which is more consistent with the severe phenotype. Her seizures were very mild and anti-epileptics were eventually discontinued once she was seizure-free for 5 years. We posit that early initiation of vitamins and combination citric acid, potassium citrate and sodium citrate coupled with good compliance explains her relatively mild course. Another possibility is her seizures were unrelated to her underlying GSSD.

Another area of concern for GSSD patients is growth and developmental delay. Our patient had an episode of weight loss and deceleration in head growth in early childhood that was attributed to chronic metabolic acidosis and that improved with increased doses of her medications. Currently, she has short stature (<5th percentile, Z-score −2.1) with appropriate weight and head circumference. Developmentally, our patient experienced minor speech delays early in life that required speech therapy and there were concerns for mild fine motor delay that resolved without intervention. She had some mild difficulties in school and required extra support in the classroom. However, she is currently performing satisfactorily in high school with plans for attending college.

In conclusion, we describe long-term follow-up of a patient with severe GSSD and good outcome. We attribute early recognition of the disease, initiation of appropriate vitamin therapy and acidosis correction along with excellent compliance from our patient for her success to date. We also believe that the use of selenium as an additional antioxidant has contributed to her relatively mild course. Finally, we recommend testing for GSSD in patients with metabolic acidosis or hemolytic anemia in the newborn period as early recognition and initiation of therapy appears to correlate with better outcomes.
CONFLICT OF INTEREST

The authors declare no conflict of interest.

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