The biallelic novel pathogenic variants in AGL gene in a Chinese patient with glycogen storage disease type III

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Abstract
Background: Glycogen storage disease type III (GSD III) is a rare autosomal recessive glycogenolysis disorder due to AGL gene variants, characterized by hepatomegaly, fasting hypoglycemia, hyperlipidemia, elevated hepatic transaminases, growth retardation, progressive myopathy, and cardiomyopathy. However, it is not easy to make a definite diagnosis in early stage of disease only based on the clinical phenotype and imageology due to its clinical heterogeneity.

Case presentation: We report a two-year-old girl with GSD III from a nonconsanguineous Chinese family, who presented with hepatomegaly, fasting hypoglycemia, hyperlipidemia, elevated levels of transaminases. Accordingly, Sanger sequencing, whole-exome sequencing of family trios, and qRT-PCR was performed, which revealed that the patient carried the compound heterogeneous variants, a novel frameshift mutation c.597delG (p. Q199Hfs*2) and a novel large gene fragment deletion of the entire exon 13 in AGL gene. The deletion of AGL was inherited from the proband’s father and the c.597delG variant was from the mother.

Conclusions: In this study, we identified two novel variants c.597delG (p. Q199Hfs*2) and deletion of the entire exon 13 in AGL in a Chinese GSD III patient. We extend the mutation spectrum of AGL. We suggest that high-throughput sequencing technology can detect and screen pathogenic variant, which is a scientific basis about genetic counseling and clinical diagnosis.

Keywords: Glycogen storage disease type III, AGL gene, Frameshift variant, Novel variant, Whole-exome sequencing

Background
Glycogen storage disease type III (GSD III; OMIM #232,400) also known as “limit dextrinosis”, “Cori” or “Forbes” disease, which was discovered by Barbara Illingworth and Gerty Cori in 1952, is a rare autosomal recessive glycogenolysis disorder with an incidence of 1:100,000 [1, 2]. Owing to the pathogenic variations in amyl glucosidase (AGL) gene (MIM #610,860), the glycogen debranching enzyme (GDE) is genetic deficiency or reduced activity, causing an accumulation of limit dextrin-like molecules in the cytoplasm of hepatocytes, myocytes, and other tissues [3].

The AGL gene has been isolated on chromosome 1p21 since 1991, and has been known as 85 kb in length with 35 exons encoding a 1532 amino acids protein (GDE) [4]. There are three important domains in AGL, the putative transferase catalytic residues, the putative glucosidase catalytic residues and the putative glycogen-binding domain (Fig. 1) [5]. Therefore, variants in these regions are more likely to influence the normal structure and function of AGL, resulting in the occurrence of GSD III. Due to alternative exon splicing and differential transcription of AGL, it has fix-transcript isoforms [6]. Isoform 1 (NM_000642) is the major isoform, which

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consists of 34 exons and widely expresses in liver, muscle, kidney, and lymphoblastoid cells [7]. GDE is a 175-kD monomeric protein containing two independent catalytic domains, amylo-1-6-glucosidase (EC 3.2.1.33) and 1,4-α-D-glucan 4-α-D-glycosyltransferase (EC 2.4.1.25). It catalyzes one of the last steps in the conversion of glycogen to glucose-1-phosphate, playing an indispensable role in the glycogen degradation [5]. To date, over 180 different AGL gene pathogenic variants have been reported worldwide (Clinvar accessed in May 2021), reflecting a high degree of genetic heterogeneity. Among these variants, the majorities are nonsense variants followed by frameshift, splice site, and missense variants (https://www.ncbi.nlm.nih.gov/clinvar).

Individuals with GSD III are characterized by progressive myopathy and liver disease. Based on differences in tissue expression level of the GDE, there are four subtypes of GSD, including GSD IIIa, IIIb, IIIc and IIId [8, 9]. GSD IIIa is the most common subtype, accounting for about 85% [3]. It mainly affects liver, skeletal, and cardiac muscle. Patients with GSD IIIb, which affects only the liver and account for about 15% of all GSD III patients [10]. It has been confirmed that selective loss of glucosidase activity or transferase activity is the pathogenic mechanism of GSD IIIc and GSD IIId, respectively [11]. Both GSD IIIc and GSD IIId are extremely rare. GSD III is usually developing in infancy and early childhood, presenting as hepatomegaly, growth retardation, fasting hypoglycemia, hyperlipidemia, significantly elevated levels of transaminases. Hepatomegaly was confirmed with abdominal ultrasound and abdominal computed tomography enhanced scan. These clinical phenotypes suggested GSD III. However,

Here, we reported a patient whose clinical diagnosis was considered as glycogen storage disease III. Whole exome sequencing (WES), Sanger sequencing, and quantitative real time polymerase chain reaction (qRT-PCR) was performed on the proband and her parents to make an accurate diagnosis. Our findings provide a wider range of AGL gene variants spectrum in patients with GSD III in the Chinese population.

Case presentation

A two-old-girl was admitted to Tianjin Children’s Hospital due to progressive enlargement of the abdomen resulting from hepatomegaly. The parents of the patient are non-consanguineous. The pedigree of the family is depicted in Fig. 2. Laboratory examinations revealed nocturnal fasting hypoglycemia, hyperlipidemia, significantly elevated levels of transaminases. Hepatomegaly was confirmed with abdominal ultrasound and abdominal computed tomography enhanced scan. These clinical phenotypes suggested GSD III. However,
electromyography excluded myopathy, cardiomyopathy was ruled out by electrocardiogram and echocardiography, and physical examination showed no obvious growth retardation.

Biochemical, imaging, clinical and diet therapy data at baseline, 8 days and 20 days are shown in Table 1.

To make a definite diagnosis, the peripheral blood of the patient and parents were sent to MyGenostics company (Beijing, China) for whole-exome sequencing (WES), Sanger sequencing, and qRT-PCR. Genetic analysis revealed a biallelic novel pathogenic variants in AGL gene. Sanger sequencing identified a previously unreported heterozygous variant in exon 5 of the AGL gene: c.597delG (p. Q199Hfs*2) (NM_000642), which inherited from the mother (I-2) (Fig. 3). This variant causes the glutamine at position 199 is changed to histidine, leading to the insertion of a stop codon at position 200. Furthermore, the heterozygous deletion of exon 13 found by WES was also a previously unreported variant, which inherited from the father(I-1). The results of qRT-PCR showed that the relative copy number of exon 13 in the control and the patient’s mother were normal, while the proband and his father were half of the normal, confirming the absence of exon 13 (Fig. 4). The deletion of exon 13 causes deletions of amino acid from position 538 to 578, leading to the termination transcription at the 540th amino acid. Both variants

Table 1 Biochemical, imaging, clinical and diet therapy data of the patient

| Clinical data | Before | 8 days later | 20 days later |
|---------------|--------|-------------|--------------|
| Age (years)   | 2      | 2           | 2            |
| Height (cm)   | 85     | 85          | 85           |
| Weight (kg)   | 14.2   | 14.2        | 14.2         |
| Abdominal circumference(cm) | 59.5   | 60.0        | 59.0         |
| Right subcostal liver(cm)    | 10     | 10          | 10           |

| Biochemical data | | | |
|------------------|--|--|--|
| Pre-prandial fasting blood glucose (mmol/L) | 4.64 | 5.67 | 6.1 |
| Nocturnal blood glucose(mmol/L) | - | 1.3 | 5.6 |
| Morning fasting blood glucose (mmol/L) | 2.5 | 4.1 | 6.5 |
| Lactate (normal range 0.5–2.22mmol/L) | 5.54 | - | - |
| Triglycerides (normal range 0–2.26 mmol/L) | 5.45 | 3.71 | 2.4 |
| Cholesterol (normal range 0–5.2 mmol/L) | 5.50 | 4.96 | 5.05 |
| Creatine kinase (normal range 40–200 U/L) | 213 | 141 | - |
| Aspartate transaminase (normal range 13–35 U/L) | 1154.6 | 452 | 532 |
| Alanine transaminase (normal range 7–40 U/L) | 781.6 | 282 | 165 |
| γ-glutamyl transpeptidase (normal range 7–45 U/L) | 98 | 101 | 84 |
| Troponin T (normal range 0–0.0223 ng/mL) | 21.0 | - | - |
| Myoglobin (normal range 25–58 ng/mL) | 0.01 | - | - |

| Imaging data | | | |
|---------------|--|--|--|
| Abdominal ultrasound | Hepatomegaly | - | Hepatomegaly Below umbilical level 55 mm |
| Abdominal computed tomography enhanced scan | Hepatomegaly No obvious abnormal enhancement | - | - |
| Liver or muscle biopsy | - | - | - |
| Electrocardiography | normal | - | - |
| Electrocardiogram | normal | - | - |
| Echocardiography | normal | - | - |
| Chest computed tomography | normal | - | - |
| Head computed tomography plain scan | - | - | - |
| Skeletal system x-rays | - | - | - |

| Diet therapy | Regular diet | Start uncooked cornstarch treatment 1-2 g/kg/d | Adhere to uncooked cornstarch treatment 1-2 g/kg/d |

'-means no done
could cause the early termination of gene transcription. The patient (II-2) inherited both variants and was therefore a compound heterozygote. The sister (II-1) had no symptoms of GSD III, who did not performed WES.

**Discussion and conclusions**

In our study, we reported a two-year-old girl (Fig. 2, II-2) from the region of China. She came to hospital for treatment mainly because of hepatomegaly. All examinations performed that she had hepatomegaly, nocturnal...
fasting hypoglycemia, hyperlipidemia, significantly elevated levels of transaminases, and midface hypoplasia with a depressed nasal bridge. However, she did not have any symptoms or exam evidence of myopathy or cardiomyopathy.

The cardinal clinical features of GSD III are hepatomegaly, fasting hypoglycemia, elevated serum concentrations of transaminases and CK [17]. The diagnosis is established by identification of biallelic pathogenic variants in AGL gene using molecular genetic testing such as single-gene testing or whole exome sequencing [18]. Liver or muscle biopsy can certify deficient glycogen debranching enzyme activity and accumulation of limit dextrin-like molecules in either liver or muscle biopsy specimen, so they are always used to distinguish different subtypes [19]. However, biopsy is more invasiveness and not required to make the diagnosis of GSD III, it is rarely performed. In our case, due to the typical clinical symptoms, the patient shown high clinical suspicion of GSD III. To make clear diagnosis, we performed genetic analysis technology on the proband and her parents. The result laid out that the proband had a biallelic likely pathogenic variants in AGL gene. Thus, the patient was diagnosed as GSD III.

Unfortunately, due to the invasive of biopsy the patient’s family disagreed with the operation, and no further detection of liver or muscle branch enzyme activity was carried out. Hence, the clinical subtype of the patient could not be confirmed. Since there was no obvious growth retardation in the patient, no further evaluation was undertaken, such as imaging examination of the skeletal system, etc. Therefore, follow-up in the later stage is very necessary.

To date, there are over 180 different AGL gene pathogenic variants registered in the Clinvar database (https://www.ncbi.nlm.nih.gov/clinvar). It includes all variant types: missense, nonsense, splice site, frameshift, and large gene fragment deletions and duplications. The majorities are frameshift (39.0%), nonsense (36.2%),
and splice-site (10.4%) variants. Depending on ethnic groups, the prevalent mutations of AGL vary in GSD III. Information on the spectrum of AGL variants will improve molecular diagnosis of GSD III in these populations and genetic counseling. For example, the most frequent variant among Italian patients is splice-site variant (c.2681+1G>A), the common nonsense variant in the Faroe Islands is p.R408X, and the prevalent variants in the United States are p.R864X and c.3964delT [20]. In Asian, c.1735 +1G>T shows the highest frequency in Japanese, Korean and Chinese patients [20]. The common variant in Turkish patients is p. W1327X [21]. Despite these, there is still high heterogeneity of AGL variants worldwide and pathogenic variants scattered throughout the gene [22]. In our case, the patient carries two compound-heterozygous variants (c.597delG and deletion of exon 13) in AGL. According to the ACMG standards and guidelines for the interpretation of sequence variants [23], the maternal frameshift variant c.597delG is a 1 bp deletion encompassed in exon 5, which causes a shift in the open reading frame and insertion of a premature termination codon (p. Q199Hfs*2), resulting in a truncating protein. This will lead to the loss of protein function and the occurrence of GSD III (PVS1). The variant is absent from controls in Exome Sequencing Project, 1000 Genomes or ExAC (PM2). Therefore, variant c.597delG is considered a likely pathogenic variant. Otherwise, the large gene fragment deletion in AGL gene with a paternal origin is a 124 bp deletion of the entire exon 13. It results in a premature truncation of the protein, leading to the loss of protein function (PVS1). The variant is absent from controls in Exome Sequencing Project, 1000 Genomes or ExAC (PM2). Thus, variant deletion of exon 13 is classified as likely pathogenic variant of AGL. Those two variants are novel and no registered in the Clinvar or the HGMD. The patient was diagnosed with GSD III according to the typical clinical features of liver disease hepatomegaly, nocturnal fasting hypoglycemia, hyperlipidemia, significantly elevated levels of transaminases, with or without elevated serum concentrations of CK. The increased serum CK is a suggestive and nonspecific marker of muscle breakdown. There are correlations between premature termination codon (PTC) variants and elevated serum CK value. Otherwise, the missense and small in-frame deletion mutations are associated with a normal serum CK level in GSD III [20]. However skeletal muscle and cardiologic complications were usually minimal during childhood, progressive myopathy can develop after the third or fourth decade of life [30]. In our study, the patient manifests as liver damage, and has no evidence for myopathy or cardiomyopathy up to now, which may associate with the age at diagnosis, disease classification or heterozygous variants of AGL. We considerate the patient probably GSD IIIb (though not confirmed) as clinical and biochemical data of the patient presented here rule out muscular manifestations. So, if there is no muscle damage in the patient during the late follow-up or confirmed by muscle biopsy as GSD IIIb, our report and the previous report [29] provide evidence that variants in exons other than exon 3 could be responsible for the GSD IIIb phenotype. Hyperlipidemia may relate to frameshift variant (c.597delG). Although previous reports suggest that PTC is associated with elevated levels of serum CK, our study shows a normal serum CK level, which may due to the patient's age or measurement in resting state [7]. These differences indicate the vary widely heterogeneity of phenotype. The follow-up date and further research is needed to clarify the relationship between PTC and CK. It also helps to understand the disease development and guide treatment in the future.

The treatment of GSD III is still controversial, and the recognized mainstay treatment is dietary management to maintain euglycemia, including uncooked cornstarch therapy with a nocturnal high-protein supplementation [33]. Recently, high fat diet has been confirmed a benefit effect to the progress of cardiomyopathy and myopathy [34]. In the past few years, significant progress has been made in gene therapy, which is expected to be more effective in the future for the treatment with GSD III [35]. In this research, the patient was treated with uncooked cornstarch 1-2 g/kg/d after being diagnosed with GSD III 8 days later. After a period of dietary control, her nocturnal fasting blood glucose returned to normal. 20 days after admission, compared to the situation at the time of

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| Reference       | Age/sex | Ethnicity or origin of study | Hepatomegaly | Liver biopsy | Cardio myopathy | Myopathy | CK | Hypertriglyceridemia | AST, ALT | Hypoglycemia | Sub type | AGL mutation                        |
|-----------------|---------|-------------------------------|--------------|--------------|-----------------|----------|----|---------------------|----------|--------------|---------|-----------------------------------|
| Crushell et al. 2010 [24] | 18Y     | Irish                         | +            | N/A          | -               | -        | N/A| +                   | +        | -            | IIb     | c[1879G>T]+[4197delA] exon 15/exon 32 |
|                  | 18Y     |                               | +            | N/A          | -               | -        | N/A| +                   | +        | -            | IIb     | c[1879delGA]+[3662C>T] exon 3/exon 28 |
|                  | 20Y     |                               | +            | N/A          | -               | -        | N/A| +                   | +        | -            | IIb     | c[1879delGA]+[3980G>A] exon 3/exon 31 |
|                  | 17Y     |                               | +            | N/A          | -               | -        | N/A| +                   | +        | -            | IIb     | c[1879delGA]+[3980G>A] exon 3/exon 31 |
| Lee et al. 2011 [25] | 16Y/F   | Nicaraguan                    | +            | +            | -               | -        | N/A| +                   | +        | -            | IIb     | 17delAG/Hom                           |
| Mili et al. 2012 [28] | 2Y/M    | Tunisian                      | +            | N/A          | N/A             | -        | -  | N/A                 | +        | -            | IIb     | IVS32-12A>G/ Hom Intron 32            |
| Minen et al. 2012 [31] | 3Y/M    | Italian                       | +            | +            | -               | -        | -  | +                   | +        | -            | N/A     | c.664+3A>G/Hom Intron 6               |
| Ben et al. 2013 [32] | 11Y/M/Con | Tunisian                      | +            | +            | +               | -        | -  | +                   | +        | -            | N/A     | c.2682-8A>G/Hom Intron 21             |
| Sentner et al. 2013 [26] | 3Y/M    | Caucasian                     | +            | +            | -               | -        | -  | +                   | +        | -            | N/A     | c.655A>G+[4529dupA] exon 6/exon 35    |
|                  | 30Y/M   |                               | +            | +            | -               | -        | -  | +                   | +        | -            | IIb     | c.16C>T/Hom Intron 3                 |
|                  | 41Y/F   |                               | +            | +            | -               | -        | -  | +                   | +        | -            | IIIb    | c.1027C>T+IVS32-12A>G/ Hom Intron 32 |
| Ko et al. 2014 [30] | 5Y/F    | Korean                        | +            | +            | -               | -        | -  | +                   | N/A      | -            | N/A     | p.[81285X]+[81675W]                   |
| Basit et al. 2014 [29] | 12Y/M   | Saudi                         | +            | +            | -               | -        | -  | +                   | +        | -            | IIIb    | VS32-12A>G/ Hom Intron 32            |
|                  | 5Y/M    |                               | +            | +            | -               | -        | -  | +                   | +        | -            | IIIb    | VS32-12A>G/ Hom Intron 32            |
|                  | 9Y/F    |                               | +            | N/A          | -               | +        | N/A| +                   | +        | -            | IIIb    | VS32-12A>G/ Hom Intron 32            |
|                  | 11Y/F   |                               | +            | N/A          | -               | -        | N/A| N/A                 | +        | -            | IIIb    | No detect any pathogenic variant     |
|                  | Con A   | Saudi                         | +            | N/A          | -               | +        | N/A| +                   | +        | -            | IIIb    | VS32-12A>G/ Hom Intron 32            |
|                  | Con B   | Saudi                         | +            | N/A          | -               | +        | N/A| +                   | +        | -            | IIIb    | VS32-12A>G/ Hom Intron 32            |
| Reference          | Age/sex | Ethnicity or origin of study | Hepatomegaly | Liver biopsy | Cardiomyopathy | Myopathy | CK | Hypertriglyceridemia | AST, ALT | Hypoglycemia | Subtype | AGL mutation |
|--------------------|---------|-------------------------------|--------------|--------------|----------------|----------|----|---------------------|----------|-------------|---------|--------------|
| Oterdoom et al. 2015 [27] | 63Y/M   | Dutch                         | +            | +            | -              | -        | -  | -                   | -        | -           | IIIb    | VS32-12A>G/ Hom Intron 32 |
| Perveen et al. 2020 [5]   | 3Y/M    | Indian                        | +            | N/A          | N/A            | N/A      | N/A| +                   | -        | +           | +       | c.[16C>T]+[1013_1014dupAT] |
|                      | 3Y/F    |                               | +            | +            | N/A            | N/A      | -  | +                   | +        | N/A         | N/A     | c.1880A>G/ Hom |
|                      | 4Y/F    |                               | +            | N/A          | N/A            | N/A      | -  | +                   | +        | N/A         | N/A     | c.4331A>G/ Het |
|                      | 3Y/M    |                               | +            | +            | N/A            | N/A      | -  | +                   | +        | N/A         | N/A     | c.309G>A/ COH |
|                      | 6Y/F    |                               | +            | +            | N/A            | N/A      | -  | +                   | +        | N/A         | N/A     | c.4334G>T/ COH |
|                      | 2.5Y/F  |                               | +            | +            | N/A            | N/A      | -  | +                   | +        | N/A         | N/A     | c.3083+1G>A/Het |
|                      | 3.5Y/M  |                               | +            | +            | N/A            | N/A      | -  | +                   | +        | N/A         | N/A     | c.3432-2392dup31/ COH |
|                      | 4Y/M    |                               | +            | +            | N/A            | N/A      | -  | +                   | +        | N/A         | N/A     | c.[2362-2392dup31]+[4334A>G] |
|                      | 3Y/M    |                               | +            | +            | N/A            | N/A      | -  | +                   | +        | N/A         | N/A     | c.2362-2392dup31+[4334A>G] |

**Abbreviations:** AGL amylo-1,6-glucosidase, D day, M month, Y year, /F female, /M male, CK creatine kinase, ALT alanine aminotransferase, AST aspartate aminotransferase; Hom homozygote, Het heterozygote, COH Compound heterozygote, Cons consanguineous, N/A not available/assessed
In summary, we identified two novel variants c.597delG (p.Q199Hfs*2) and deletion of the entire exon 13 in AGL in Chinese GSD III patients. We extend the mutation spectrum of AGL in Chinese patients with GSD III and point out to the heterogenous genetic background of the disease, providing theoretical guidance for genetic counseling and precision therapy. Our results suggest the need for complete AGL gene sequencing to diagnose the disease based on a suggestive clinical feature and provide genetic counseling for young families. However, it is limited to hypothesize on the correlation between genotype and phenotype just based on this patient only. In the future, we will continue to collect cases for cohort studies in order to expand the basic information of the disease and establish genotype-phenotype correlation.

Abbreviations

AGL: Amyl glucosidase gene; GSDII: Glycogen storage disease type II; GDE: Glycogen debranching enzyme; WES: Whole exome sequencing; ALT: Alanine aminotransferase; AST: Aspartate aminotransferase; CK: Creatine kinase; PTC: Premature termination codon.

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Authors’ contributions

YW was involved in the collection and critical analysis of clinical data. YP, YZ and JB contributed to the designing the study, supervision, and critical review of the manuscript. XF, Z and YZ contributed to the supervision and critical analysis of laboratory data, including the interpretation of NGS results. YZ and JB confirmed the authenticity of all the raw data and revised the manuscript. All authors read and approved the final manuscript.

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Availability of data and materials

The raw datasets used and analysed during the current study are not deposited in publicly available repositories because of considerations about the privacy or security of human. However, the datasets can be available from the corresponding author on reasonable request.

Declarations

Ethics approval and consent to participate

Written informed consent was obtained from the patient (or the parent) and the study was approved by the ethics committee of Tianjin Children’s Hospital (Tianjin, China).

Consent for publication

Written informed consent was obtained from the patient’s parents for publication of this Case report and any accompanying images. A copy of the written consent is available for review by the Editor of this journal. Written informed consent was obtained from the patient’s parents for publication.

Competing interests

The authors declare that they have no competing interests.

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