Glutaric Aciduria Type 1: Clinical and Molecular Study in Iranian Patients, 3 Novel Mutations


Abstract

Objective

Glutaric aciduria type 1 (GA1), is a rare, treatable neuro metabolic disease, due to glutaryl-CoA dehydrogenase (GCDH) gene mutation. In regions without neonatal blood screening (NBS), patients are diagnosed in symptomatic period. This study was carried out to assess patients with GA1 for clinical, biochemical, neuroimaging findings and GCDH gene mutations analysis.

Materials & Methods

In this cross-sectional study, clinical manifestation, neuroimaging and metabolic findings of eleven Iranian GA1 patients of Mofid Children’s Hospital, Tehran, Iran between 2001 and 2011, were evaluated. Mutational analysis of the GCDH gene was performed on genomic DNA. Genomic DNA was extracted from peripheral lymphocytes using QIAamp DNA Micro Kit (Qiagen). All 11 exons and flanking intronic regions of the GCDH gene were amplified by polymerase chain reaction (PCR).

Results

All patients were diagnosed before 32 months old. Clinical presentations of GA1 include acute encephalopathic crisis and/or developmental delay and macrocephaly. Seven GCDH gene mutations were detected in our patients. The most frequent GCDH mutations occurred in exon7 then exon8, 10 and 11. G244 C in exon7, R294 Q in exon8 and N373 S in exon 10 were three novel mutations. There was no correlation between of genotype and phenotype in our patients.

Conclusion

Physician must remember GA1 in differential diagnosis of acute encephalopathic crisis, macrocephaly, developmental delay, movement disorders such as dystonia and dyskinesia. Early detection, proper treatment and selective screening of patients’ siblings can prevent neurologic disabilities.

Keywords: Glutaricaciduria type1; Glutaryl co-A dehydrogenase; GCDH mutation; Iran
Introduction

Glutaric aciduria type 1 (GA1, #231670) was introduced as a new neurodegenerative disorder in 1975 (1). This progressive neurometabolic disorder with characteristic clinical presentation usually manifests with macrocephaly, acute encephalopathic crisis after upper respiratory/gastrointestinal infection, fever, dehydration, vaccination or in lesser extent with insidious-onset presentation such as movement disorder (dystonia, ataxia), developmental delay (2-4). Occasionally child physical abuse has been suspected because of presence of subdural effusions (5). Life expectancy is greatly reduced in symptomatic patients with complex movement disorder (6). The peak age of presentation is 9 months and most symptomatic patients present before 3 yr of age; although rare symptomatic neurological presentation (late onset) are reported after 5 yr old of age (7,8).

Frequent neuroimaging findings in GA1 are described in the different medical literature include widened Sylvian fissures, frontotemporal volume loss, ventriculomegaly, subdural hematomas, delayed myelination demyelination and basal ganglia lesions (9,10).

Deficient activity of mitochondrial enzyme glutaryl-CoA dehydrogenase (GCDH; EC.1.3.99.7) that is essential for degradation of lysine, hydroxylysine and tryptophan pathway is responsible for this autosomal recessive disorder of inborn error of metabolism. Thus, the biochemical diagnosis is based on detection of increased levels glutaric acid, 3-hydroxyglutaric acid in urine, other physiologic body fluid and elevation glutarylcarnitine (C5DC) level in plasma or urine (11). Suspected GA1 patients based on clinical, biochemical, neuroimaging finding is confirmed by GCDH gene mutations or GCDH enzyme activity.

GCDH gene encodes GCDH enzyme located on chromosome 19p13.2 and is composed of 11 exons and 10 introns (12,13). More than 200 mutations have been reported in the GCDH gene (14). However, some GCDH gene mutations are frequent in specific population. This study was carried out to assess patients with GA1 for clinical, biochemical, neuroimaging findings and GCDH gene mutations analysis.

Materials & Methods

We studied 11 Iranian patients with GA1 previously diagnosed in Neurology and Endocrinology Clinics of Mofid Children’s Hospital, Tehran, Iran between 2001 and 2011. Fourteen Iranian GA1 patients were diagnosed during these 10 yr. One patient died. One family did not accept to test their child for GCDH gene mutation. One patient’s family immigrated to another country. Diagnoses of these symptomatic patients were based on glutarylcarnitine analysis in their dried blood spots or urine and/or their urinary organic acid profiles. They were invited to hospital for interview, physical examination, collecting their urine and blood laboratory information, neuroimaging findings and performing GCDH gene mutation study. GCDH gene mutation studies were done in Special Medical Center, Tehran, Iran.

Mutational analysis of the GCDH gene was performed on genomic DNA. Genomic DNA was extracted from peripheral lymphocytes using QIAamp DNA Micro Kit (Qiagen). All 11 exons and flanking intronic regions of the GCDH gene were amplified by PCR. Primer sequences are given in Table 1 (13). PCR was performed in a total volume 25 µl containing 1x PCR buffer, 1.5 mM MgCl2, 8 pmol each primer, 0.2 mM each dNTP, 50 ng template DNA, and 0.3 U Taq polymerase (SinaClon, Iran). Five percent DMSO was also added for PCR of exon 2. Amplification was performed with initial denaturation at 94°C for 5 min, 32 cycles of 1 min at 94°C, 1 min at 55-62°C at annealing according to Table 1, and 1 min at 72°C and final extension were 5 min at 72°C. The PCR products were sequenced on ABI 3700 sequencer (Kosar Company, Tehran). Sequencing results were compared with ref sequences in NCBI. Informed consent to perform DNA analysis was obtained from the parents of the patients. The Ethics Committee of the Faculty of Medicine, Shahid Beheshti University of Medical Sciences, Tehran, Iran approved our study protocol.

Results

The clinical and molecular findings of 11 Iranian GA1 patients are summarized in Table 1.
<table>
<thead>
<tr>
<th>Patient ID</th>
<th>Age at diagnosis (months)</th>
<th>Age at mutation analysis (months)</th>
<th>Presentation</th>
<th>Macrocephaly after birth</th>
<th>Exon</th>
<th>Nucleotide substitution</th>
<th>Amino-acid change</th>
<th>Mutation</th>
<th>Movement disability</th>
<th>Reference</th>
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<td>1</td>
<td>9</td>
<td>52</td>
<td>Developmental delay, dystonia,</td>
<td>No</td>
<td>7</td>
<td>C&gt;T</td>
<td>CCC&gt;CTC</td>
<td>Pro&gt;Leu</td>
<td>P 248 L</td>
<td>severe</td>
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<td>2</td>
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<td>13</td>
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<td>”</td>
<td>”</td>
<td>moderate</td>
<td>”</td>
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<td>”</td>
<td>severe</td>
<td>”</td>
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<td>34</td>
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<td>7</td>
<td>T&gt;C</td>
<td>TTC&gt;CTC</td>
<td>Phe&gt;Leu</td>
<td>F236L</td>
<td>severe</td>
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<tr>
<td>5</td>
<td>12</td>
<td>23</td>
<td>Acute: seizure</td>
<td>Yes</td>
<td>7</td>
<td>G&gt;T</td>
<td>GGT&gt;TGT</td>
<td>Gly&gt;Cys</td>
<td>G244C</td>
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<td>C&gt;T</td>
<td>ACG&gt;ATG</td>
<td>Thr&gt;Met</td>
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<tr>
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<td>10</td>
<td>A&gt;G</td>
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<td>65</td>
<td>Acute: neonatal and postvaccinal seizure,</td>
<td>Yes</td>
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<td>”</td>
<td>”</td>
<td>”</td>
<td>mild</td>
</tr>
<tr>
<td>10</td>
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<td>15</td>
<td>Acute: irritability encephalopathy-</td>
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<td>8</td>
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<td>CGG&gt;CCG</td>
<td>Arg&gt;Pro</td>
<td>A294P</td>
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<td>15</td>
<td>102</td>
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<td>8</td>
<td>”</td>
<td>”</td>
<td>”</td>
<td>”</td>
<td>severe</td>
</tr>
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</table>
Demographic and clinical findings
Eleven symptomatic Iranian patients with GA1 were diagnosed from 10 families. The only sibling patients were two brothers, delivered as triple pregnancy. All of patients were offspring of consanguineous marriages. Ethnic origin and gender frequency of four patients are shown in Figure 1.

The mean age at diagnosis was 14.8 (range: 5-32) months. The mean age at study time was 44.5 (range 15-102) months. Acute encephalopathic crisis and seizure following URI or vaccination were the first clinical presentations in six cases. Macrocephaly, developmental delay and extrapyramidal signs were the first presentations in other five patients. Birth head circumferences of all patients were under percentile 95 (percentile 25-90). Patients were treated by low protein (lysine, tryptophan, etc.) diets and carnitine supplementation. Treatments were discontinued in five patients because their families did not fill any improvement in their child symptoms and signs. Other families tried to continue this treatment, although these patients were diagnosed, recently.

Neuroimaging findings
Included widened Sylvian fissure in 100%, bitemporal arachnoid cysts in 60%, white matter signal changes in 33% and basal ganglia lesions in 27% of 25 times brain neuroimaging. Neuroimaging studies include 16 brain CT scan and 9 brains MRI in eleven patients (Figure 2-4).
Metabolic analysis finding

Our patients’ diagnoses were based on blood and/or urine analysis. Bloodglutarylcarnitine and glutarylcarnitine/free carnitine ratio were increased in 11 patients. Urines of 5 patients were tested for organic acid. Increased excretions of glutaric acid and/or 3-hydroxyglutaric acid (organic acids) were reported in these 5 patients.

GCDH mutation

We found 7 GCDH gene mutations in eleven patients, three of them were not reported before. Five (45%) patients had mutations in exon 7. P 248 L mutation in exon 7 was detected in 3(27%) patients. Three novel mutations were G244C (exon7), A294P (exon8) and N373SMutation in exon10. Four different mutations were seen in 6 Turkish patients. More information about GCDH mutations is shown in Table1, too.

Discussion

Heterogeneous clinical manifestations, biochemical findings, and GCDH gene mutations are seen in GA1. A severe and treatable inborn error of metabolism usually diagnosed in some countries in neonatal period through expanded newborn screening program using analysis of glutarylcarnitine in dried blood spots by tandem mass spectrometry(15,16). In areas without newborn screening for acylcarnitine profile such as Iran at present, patients with GA1 usually are diagnosed in symptomatic period or as a selective screening in siblings of a known GA1 patient (17). In most neonates diagnosed with presymptomatic neonatal period, striatal necrosis and movement disorder can be prevented by low lysine diet and carnitine supplementation (18). An emergency treatment protocol in catabolic states such as fever, gastroenteritis, surgery, immunization, including natural protein stopping, proper hydration, increase carnitine supplementation, appropriate glucose and caloric supply, brisk alkaline urine output, proper antibiotics and antipyretic drugs also are important to prevent neurological sequels(19,20).

Acute encephalopathic crisis was observed as equally as insidious presentation (developmental delay dystonia) in our patients. Acute encephalopathic crisis was reported in majority of patients with GA1 (4) and most patients were presented insidious with developmental delay, macrocephaly (21,22). The discrepancy is due to likely different factors such as the misdiagnosis of initial acute crises by poorly trained physicians or perhaps different GCDH gene mutations.

Enlarge head circumference was described in most neonates with GA1 as the only early sign of this disorder (19).Macrocephaly (occipito-frontal circumference above 95 percentile) was not present at birth in our patients. However, macrocephaly was observed at interview time in our seven (63%) patients.
Dietary management was not continued in five of our severe disabled patients because they did not fill any improvement with this type of therapy. No detectable beneficial effects of long-term dietary treatment were reported in patients with obvious neurological symptoms (6). Although strict low lysine diet is recommended especially in asymptomatic patients up to age of 6 yr old, continuing less strict diet therapy is advised after 6 yr old, too. Carnitine supplementation must be continued lifelong (23). Detection of patients in asymptomatic period and appropriate prospective care reduces clinical and neuroimaging presentation of basal ganglia injury from 90% to 35% (19).

Our patients’ neuroimaging findings showed widened Sylvian fissure (bat wing sign) in all patients. Widened Sylvian fissure as characteristic but not pathogenomic neuroimaging finding was reported in 93% of neuroimaging in symptomatic GA1 patients (9). This finding is reported in asymptomatic patients detected on expanded newborn screening suggestive temporal hypoplasia rather than frontotemporal atrophy (10).

White matter abnormalities were detected in 33% of our patients’ brain neuroimaging. White matter changes did not correlate with clinical manifestation. Abnormal white matter changes in Amish patients were reported less frequently (6%) than non-Amish patients (32%). The exact mechanism of this abnormal signal is not understood (19). We found basal ganglia involvement in 27% of neuroimagings of patients. Acute striatal necrosis was reported as the major cause of movement morbidity and mortality in GA1 patients. Nutritional and pharmacological management of newborn detected in expanded newborn screening in asymptomatic period-reduced incidence of striatal injury. However, it is unclear why despite early asymptomatic detection of GA1 patients and treatment intervention; approximately one-third of them developed striatal injury (19).

Usually suspected GA1 patients based on clinical, neuroimaging, biochemical findings are confirmed by molecular or enzymatic study. GCDH gene mutation is used to confirm GA1, carrier detection, genetic consultation and prenatal diagnosis of this disorder. To date, more than 200 mutations in GCDH gene have been reported. Some mutations are frequent in specific populations. In Ojibway-Cree linguistic group, patients in Manitoba and northwest of Ontario IVS1+5 G > T mutation is prevalent (24). In North Carolina, homozygous mutation E404K (1240 G> A) in exon 10 was reported (25). Selective screening of common GCDH gene mutation could be a useful way to decrease time and cost of molecular gene analysis.

In our eleven patients with GA1, seven different GCDH gene mutations including three reported previously were detected. Near about half (45%) of GCDH mutations occurred in exon 7, other mutations occurred in exon 8, 10, 11 equally. P 248 L mutation in exon 7 as the most frequent GCDH gene mutation in our study was reported in Turkey and Italy before (13). Other reported mutation in exon 7 was F236 L (26). A 433 V mutation in exon 11 was reported from Spain (3). The last mutation T429M in exon 11 was reported (27). G244 C in exon 7, R294 Q in exon 8 and N373 S in exon 10 were novel mutations in this study. GCDH gene mutation in patients with GA1 in Khuzestan Province in southeastern of Iran revealed E181Q, S255L, R402G, in order of frequency, were responsible for this disorder in that region (28). R 402 was the most common mutation in European patients with GA1 was not detected in our patients, too. More than half of our patients were Turkish and we detect four different mutations in their GCDH gene. GA1 is more prevalent in this ethnicity. We did not find any evidence to document genotype and clinical phenotype correlation, owing to high variation of the disease course among patients, as another study (4).

In Conclusion, GA1 is a heterogeneous rapid neurodegenerative disorder with available biochemical and genetic test to detect asymptomatic patients. Expanded newborn screening program is recommended in this country. More patients’ genetic analysis is required to know better about GCDH gene mutation Iranian patients with GA1.

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**Authors’ Contribution**

Pirzadeh Z: Contributions to the design of the work, the acquisition of data for the work, analysis of data for the work, and drafting the work

Houshmand M: Contributions to the conception of the work, the acquisition of data for the work, interpretation of data for the work, and revising the work critically for important intellectual content

NASIRI J: Contributions to the acquisition of data for the work, analysis of data for the work, and drafting the work

Mollamohammadi M: Contributions to the acquisition of data for the work, analysis of data for the work, and drafting the work

Sedighi M: Contributions to the acquisition of data for the work, analysis of data for the work, and drafting the work

Tonekaboni SH: Contributions to the conception of the work, interpretation of data for the work, and revising the work critically for important intellectual content

All authors approved the final version to be published and agreed to be accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved.

**Conflict of interest**

The authors declare that there is no conflict of interest.

**References**


