Identification of Novel Mutations in the \textbf{MMAA} and \textbf{MUT} Genes among Methylmalonic Aciduria Families

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OPEN ACCESS

Received: 27 June 2022  
Accepted: 8 February 2023  
Published online: 12 February 2023

ABSTRACT

\textbf{Background:} Methylmalonic aciduria is a rare inherited metabolic disorder with autosomal recessive inheritance pattern. There are still MMA patients without known mutations in the responsible genes. This study aimed to identify mutations in Iranian MMA families using autozygosity mapping and NGS.

\textbf{Methods:} Multiplex PCR was performed on DNAs isolated from 12 unrelated MMA patients and their family members using 19 STR markers flanking \textit{MUT}, \textit{MMAA}, and \textit{MMAB} genes, followed by Sanger sequencing. \textit{WES} was carried out in the patients with no mutation.

\textbf{Results:} Haplotype analysis and Sanger sequencing revealed two novel mutations, A252V*5 and G87R, within the \textit{MMAA} and \textit{MUT} genes, respectively. Three patients showed no mutations in either autozygosity mapping or NGS analysis.

\textbf{Conclusion:} High-frequency mutations within exons 2 and 3 of \textit{MUT} gene and exon 7 of \textit{MMAB} gene are consistent with the global expected frequency of genetic variations among MMA patients. \textbf{DOI: 10.61186/ibj.3782}

Keywords: Autozygosity mapping, Genotype, Methylmalonic acidemia

INTRODUCTION

Methylmalonic aciduria is a classic form of organic acidemia caused by mutations in genes encoding methylmalonyl-CoA mutase and its co-enzyme, 5’-deoxyadenosylcobalamin. Prevalence of this disease varies worldwide and has recently been reported as 1.14 per 100,000 neonates\textsuperscript{[1]}. According to reports of mutations disabling the critical genes, clinical presentations and onset of the MMA may vary among different patients and populations. MMA can be caused by the deficiency in either MUT enzyme or cobalamin (vitamin B12). Cobalamin deficiency occurs as the result of disturbances in the process of synthesis of adenosylcobalamin. MMA is also responsive to vitamin B12 supplement therapy and can be induced by mutations in \textit{MMAA} (4q31), \textit{MMAB} (12q24), and \textit{MADHIC} genes or so-called cblA, cblB, and cblDv2 types, respectively.

Current diagnosis of MMA is based on a combination

List of Abbreviations:  
NGS: next generation sequencing; MMA: methylmalonic aciduria; PCR: polymerase chain reaction; STR: short tandem repeat; WES: whole exome sequencing
of biochemical and genetic analyses. Genetic counseling for the suspected families can not only increase overall detection rate of the disease but also decrease the time and cost of unnecessary genetic tests[2,3]. Biochemical analysis includes detecting the level of urinary methylmalonic acid, 3-hydroxypropionate acid, citric acid, plasma glycine, valine, threonine, isoleucine, carnitine, and methionine. Genetic analysis often entails genotyping of MMAA (4q31), MMAB (12q24), and MUT (6p21) genes[4]. However, some of MMA patients have no known mutations in responsible genes (MUT, MMAA, and MMAB) in the patients with typical biochemical and clinical characteristics of MMA sometimes have no known mutation[5].

A recent study has reported five novel mutations, including c.805delG, c.693delC, c.223A>T, c.668A>G, and c.976A>G, within the MUT gene among Iranian patients, who are clinically diagnosed as MMA[6]. In another study, PCR sequencing analysis of MMAA gene in one family with two MMA siblings detected a homozygous deletion, c.674delA, in exon 4 of MMAA in both affected cases[7]. A separate study investigating three unrelated MMA patients by PCR sequencing reported one homozygous nucleotide change (c.2125-3

C>G) in the intron 12 of the MUT gene[8]. A research group found a novel C to G variation at the position -3 in the intron 12 of the MUT gene in two probands with definite diagnosis of MMA, which were associated with the retention of intron 12 in the final transcript product[9]. Herein, we performed autozygosity mapping using 19 STR markers flanking MMAA, MMAB, and MUT genes to identify candidate genes for MMA disease among the selected families. The identified genes were then subjected to PCR and Sanger sequencing and the negative cases were further analyzed by WES.

MATERIALS AND METHODS

Selection of patients

Twelve unrelated Iranian patients with the same ethnicity, including six girls and six boys, affected by the isolated form of MMA were referred to Kawsar Human Genetics Research Center by pediatric endocrinologists. All the patients had a consanguineous marriage (first cousin). Diagnosis of the disease was based on the clinical presentations and the metabolite levels, methylmalonic acid and methylicric acid.

Molecular genetic studies

After genetic counseling for the patients’ families, 5 ml of whole blood was collected from all the cases and kept in EDTA-containing tubes. DNA was isolated from blood samples using the QIAamp DNA Mini Kit (Qiagen, Germany) and stored at -20°C until further steps. Finding STRs of the genes was performed using tandem repeat finder software. To consider the STR markers with higher heterozygosity, each marker was examined in 10 random samples. The designed primer pair sequences (using Primer3 software) with fluorescent primer binding site were employed for the amplification of STRs[6]. Selected STR markers were amplified in two independent multiplex PCR reaction, followed by fragment analysis by the ABI 3130 XL Genetic Analyzer (Thermo Fisher Scientific, USA). Haplotype maps were then separately drawn for all the families enrolled in the study with positive mutation. Cases with similar mutations in their family members were selected to be further analyzed by WES and carried out with targeted depth of 100x. Samples were prepared according to TruSeq Nano DNA library preparation and sequenced by an Illumina HiSeq X Ten platform (Centogene, Rostok, Germany). After aligning the sequencing reads to the NCBI Build 38 of the human reference sequence and merging the alignments into a single BAM file, variants were defined through Genome Analysis Toolkit and then annotated using Ensemble Variant Effect Predictor. Positive cases, the only carriers of the mutation within their families, were subjected to further analysis by Sanger sequencing using specific primer pairs previously designed for all the exons of MMAA, MMAB, and MUT genes[6]. Sanger sequencing was performed on PCR products using a BigDye Terminator Kit (Thermo Fisher Scientific) according to the manufacturer’s protocol and then resolved on the ABI 3130xl genetic analyzer (Thermo Fisher Scientific) at the Kawsar Biotech Center facility. The identified novel mutations were genotyped by PCR Sanger sequencing in 100 Iranian healthy controls aged 17-60 years old, who were randomly referred to a cosmetic clinic to further determine their pathogenicity.

In silico analysis

The pathogenicity of the identified variants was investigated using different software, including Human Gene Mutation Database, MutationTaster, DynaMut, Fathmm, PolyPhen, and ClinVar.

RESULTS

Autozygosity mapping and Sanger sequencing demonstrated homozygosity, including c.668A>G (K223R), c.259G>A (G87R), c.322C>T (R108C), c.1106G>A (R369H), and c.454C>T (R152X), for the MUT gene in five patients, while c.569G>A (R190H), IVS2-1G>T, and c.557G>A (R186Q) substitutions were
found in other three patients in MMAB gene (Fig. 1). Moreover, c.749_750insGT on (A252Vi*5) alteration was found in MMAA gene (Table 1: Supplementary Figs. 1 and 2). Three MMA patients revealed no homozygosity in none of the studied genes, and they were then subjected to further analysis by WES. In one of the patients examined by WES, c.380C>A (A127D) was found in the MMAB gene. In the second patient, c.628A>C (K210Q), c.901A>T (M301L), and c.1084A>T (M362L) were identified in the ACSF3 gene. No mutation was found in the third patient. The identified mutations were examined in family members by Sanger Sequencing to determine their role in the pathogenicity of MMA disease (Supplementary Fig. 3). The c.380C>A mutation was detected in homozygous status in the healthy brother of proband, and, therefore, its pathogenicity was ruled out. Owing to the presence of ACSF3 gene mutations in other family members, the pathogenicity of those mutations was also ruled out (Supplementary Figure 3). First cousin of the latter case suffered from MMA. In the WES analysis, he demonstrated no mutation, including mutations in ACSF3 gene. Two other patients with homozygosity in the MMAB gene did not show any pathogenic mutations, and one patient showed homozygosity, c.749_750insGT on, in the MMAA gene mutation. There are no previous reports on c.259G>A and c.749_750insGT on mutations; therefore, they were introduced as novel cases. Parents of all the patients showed mutations in heterozygote form. There was no familial history among the enrolled families in terms of the inborn errors of metabolism, except for one family who had a male first cousin affected by MMA. He has previously been subjected to NGS analysis, and no mutation was found. It is worth to note that no novel mutation was identified in the control group.
## DISCUSSION

Herein, two novel mutations, c.259G>A (G87R) and c.749_750insGTTT insertion mutation, were found in the *MUT* and *MMAA* genes, respectively. Protein structure and function analysis of the c.259G>A mutation revealed that arginine substitution can cause deletion of 6 β-sheets, 32 α-helix, and B12-binding domain, which were associated with reduced protein stability. We also found a 15-month-old male patient with severe clinical presentations, including developmental delay, recurrent vomiting, and acidosis, which was compatible with the deleterious effect of mutation on protein function. Third novel mutation, 749_750insGTTT, was predicted to be associated with reduced enzyme activity, deletion of some peptides in alpha helix, and GTP-binding site of MMAA. This protein has a critical role in the exchange of co-factor of enzyme; thus, 749_750insGTTT mutation can impede recycling of coenzyme, which in turn decreases the enzyme activity[10,11]. Reduced enzyme activity with residual function is consistent with less severe and relatively late onset (14 months) presentation of our patient, including hypotonia as the major symptom in physical examination. Remaining mutations, i.e. c.569G>A, c.557G>A, and IVS2-1G>T in *MMAB* gene and c.1106G>A, c.454C>T, and c.322C>T in *MUT* gene have previously been reported in different studies. The c.569G>A substitution was found in a three-year-old female patient with major clinical presentations of poor feeding and agitation. Also, it has previously been reported in two separate studies and primarily found in two MMA type cblB patients with the early onset of the disease in both homozygote and heterozygote status[11,13]. Zhang et al.[12] have demonstrated that histidine substitution at the 190 residue position can dramatically change the affinity of MMAB to adenosylcobalamin. Finding this mutation and displaying the clinical presentations in either homozygote or heterozygote status could be a further confirmation on the pathogenicity of this mutation. The c.557G>A mutation was the next recurrent mutation found as homozygote in a one-year-old male patient with severe developmental delay and acidosis. This mutation has been formerly been reported in a white 14-year-old male patient with undefined clinical presentations, which has been predicted to affect enzyme activity through disrupting active site of protein and disabling its interaction with ATP in spite of normal enzyme production[11]. Furthermore, it has been found in two Canadian cblB type patients in both heterozygote and homozygote status[13]. While the c.557G>A mutation has been assigned as uncertain significance in ClinVar website, its replication in our patient with severe disease.

<table>
<thead>
<tr>
<th>Patient No.</th>
<th>Age at diagnosis</th>
<th>Sex</th>
<th>Current age (y)</th>
<th>Clinical symptoms at diagnosis</th>
<th>Mutation at nucleotide level</th>
<th>Mutation at protein level</th>
<th>Exon</th>
<th>Gene</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2 years</td>
<td>F</td>
<td>1</td>
<td>Severe developmental and growth delay</td>
<td>c.322G&gt;T</td>
<td>R108C</td>
<td>2</td>
<td>MUT</td>
</tr>
<tr>
<td>2</td>
<td>1 years</td>
<td>M</td>
<td>2</td>
<td>Severe developmental and growth delay, Frequent vomiting and acidosis</td>
<td>c.557G&gt;A</td>
<td>R186Q</td>
<td>7</td>
<td>MMAB</td>
</tr>
<tr>
<td>3</td>
<td>15 months</td>
<td>M</td>
<td>3</td>
<td>Developmental and growth delay, Frequent vomiting and acidosis</td>
<td>c.259G&gt;A</td>
<td>G87R</td>
<td>2</td>
<td>MUT</td>
</tr>
<tr>
<td>4</td>
<td>5 months</td>
<td>M</td>
<td>1</td>
<td>Hypotonia, developmental delay</td>
<td>c.668A&gt;G</td>
<td>K223R</td>
<td>3</td>
<td>MUT</td>
</tr>
<tr>
<td>5</td>
<td>10 months</td>
<td>F</td>
<td>2</td>
<td>Frequent vomiting and acidosis</td>
<td>c.454C&gt;T</td>
<td>R152X</td>
<td>3</td>
<td>MUT</td>
</tr>
<tr>
<td>6</td>
<td>6 months</td>
<td>F</td>
<td>3</td>
<td>Agitation, poor feeding</td>
<td>c.569G&gt;A</td>
<td>R190H</td>
<td>7</td>
<td>MMAB</td>
</tr>
<tr>
<td>7</td>
<td>3 days</td>
<td>M</td>
<td>7</td>
<td>Poor feeding, hypotonia</td>
<td>c.1106G&gt;A</td>
<td>R369H</td>
<td>6</td>
<td>MUT</td>
</tr>
<tr>
<td>8</td>
<td>14 months</td>
<td>F</td>
<td>17</td>
<td>Hypotonia</td>
<td>749_750insGTTT*</td>
<td>A252Vf*5</td>
<td>5</td>
<td>MMAB</td>
</tr>
<tr>
<td>9</td>
<td>2 years</td>
<td>F</td>
<td>16</td>
<td>Frequent vomiting and acidosis</td>
<td>IVS2-1G&gt;T</td>
<td>-</td>
<td>2</td>
<td>MMAB</td>
</tr>
</tbody>
</table>

*Novel mutation*
manifestations may clinically classify it as a definite pathogenic variant.

The c.197-G>T alteration was identified in a two-year-old female patient with vomiting and acidosis. It has also previously reported in two Arab patients and validated as pathogenic mutation[14,15]. The c.1106G>A mutation was determined in a three-day-old male neonate with poor feeding and lethargy. This mutation has been recognized as pathogenic variant and frequently reported in the patients with Caucasian origin in two earlier studies[16,17]. It was also characterized as catalytic mutation, which disrupts the highly conserved residue (arginine) at the position of 369. Additionally, the c.1106G>A substitution occurs in a CG-rich hot spot and it is always found as homozygote and associated with mut[6] phenotype, which is in line with our findings[16]. The c.454C>T (R152X) mutation was detected in our 10-month-old female patient with severe clinical presentations, including recurrent vomiting and acidosis. It is one of the most recurrent MUT gene mutations frequently reported in patients with the early onset severe clinical manifestations, which is in agreement with our patient’s phenotype[16,18-20].

Data regarding the premature stop codon and truncated protein as the result of c.454C>T (R152X) mutation is also compatible with the early and severe disease manifestations, which has been further replicated in the present study.

Prediction of the enzyme structure through online software demonstrated that c.668A>G substitution and replacement of lysine with arginine result in the reduced enzyme activity, as well as deletion of 26 alpha helix and vitamin B12-binding domain, which in turn disrupts the final protein structure and function. This mutation was found in a five-month-old male patient that his major clinical presentations were hypotonia, lethargy, and developmental delay, which are consistent with the severity of the identified mutation. The c.668A>G has previously been identified in a compound heterozygote Iranian MMA patient. Habibzadeh et al.[21] have also found another truncating mutation (c.1055A>G) in exon 5 of MUT gene of a 15-month-old patient presenting the same clinical manifestations, as well as pneumonia aspiration.

All the identified mutations, except for one patient, were in homozygous status, which revealed as compound heterozygote, including c.628A>C, c.901A>T, and c.1084A>T. She was a 16-year-old patient who was absolutely normal till the 15 years of age and was then affected by sudden aphasia and hearing loss. Investigation of the mutations found within the ACSF3 gene in her parents indicated that they had no important defect in ACSF3 protein structure and function. Hence, severe late onset neurologic demonstrations might rely on the presence of possible mutation/s within MMACHC gene[22].

One of the remarkable points of the present study is the clear genotype-phenotype correlation, which was found in most of the identified mutations, regardless of the patient number 2 with 557G>A mutation. Minimal thermal and stability changes in the protein structure, in spite of severe neurological manifestations, may indicate the presence of mutations in other responsible genes, including MMACHC and MCEE[23].

Herein, we demonstrated that autozygosity mapping using selected STR markers can dramatically decrease the time and cost of the analysis for patients, particularly in those who were born from consanguineous marriage[6]. We could find pathogenic mutations in 9/12 (75%) of the patients. Regarding the patients with no mutations in the assayed exons, the most remained probability was mutations in other MMA responsible genes or cryptic mutations within the introns. There are limited reports on pathogenic intron mutations within three major MUT, MMAA, and MMAB genes, which were almost associated with splicing defect[8,9,19]. In addition, frequency of the mutations in the three main MMA genes has been estimated as 97%. Therefore, finding a mutation in two remaining genes may change the overall frequency of gene alterations at least in our population[24]. The frequency of the mutations was higher in exons 2 and 3 of the MUT gene and exon 7 of MMAB gene compared to other exons and responsible genes, which strongly corroborates most of the MMA genotyping studies performed previously[11,17,25,26].

Herein, two novel mutations, 749_750insGTTT and c.259G>A, were introduced within the MMAA and MUT genes, respectively. In silico analysis of the effects of the mutations on their corresponding protein structure and function was in agreement with clinical manifestations of the patients. However, we could not find any pathogenic mutations in the key MMA genes, in three patients. Further studies are warranted to determine the responsible mutations in other susceptible genes and the introns of MUT, MMAA, and MMAB genes to improve genetic counseling of patients with MMA disease.

DECLARATIONS

Ethical statement

The study protocol was approved by the Ethics Committee of Kaw sar Human Genetic Research Center, Tehran, Iran (ethical code: 98/6301). All the patients provided their written consents.
Data availability
Data supporting this article are included within the article and supplementary file.

Author contributions
MJ: collected samples and performed laboratory assessment; FK: supervised the project, wrote the manuscript draft and performed the final manuscript revision; AS, AR, HB, MRA, FR: collected samples and analyzed the data; SZ: supervised the project and performed final manuscript revision.

Conflict of interest
None declared.

Funding/support
There is no funding for this project.

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