Review article



# Methylmalonic Acidemia Diagnosis by Laboratory Methods

Fatemeh Keyfi<sup>1, 2</sup>, Saeed Talebi<sup>3</sup>, Abdol-Reza Varasteh<sup>\*2, 4, 5</sup>

## Abstract

Methylmalonic acidemia (MMA) is usually caused by a deficiency of the enzyme methylmalonyl-CoA mutase (MCM), a defect in the transport or synthesis of its cofactor, adenosyl-cobalamin (cblA, cblB, cblC, cblF, cblD, and cblX), or deficiency of the enzyme methylmalonyl-CoA epimerase. A comprehensive diagnostic approach involves investigations of metabolites with tandem mass spectrometry, organic acid analysis with gas chromatography, enzymatic studies with fibroblast cell culture, and finally, mutation analysis. With biochemical techniques and enzymatic assay the reliable characterization of patients with isolated MMA for mutation analysis can be achieved. Reliable classification of these patients is essential for ongoing and prospective studies on treatments, outcomes, and prenatal diagnoses. This article reviews the diagnostic techniques used to characterize patients with MMA.

Keywords: Diagnostic techniques, Enzyme assay, Methylmalonic acidemia, Mutation analysis, Organic acid analysis, Tandem mass spectrometry

## Introduction

Methylmalonic acidemia (MMA) is usually caused by a deficiency of the enzyme methylmalonyl-CoA mutase (MCM, EC 5.4.99.2), a defect in the transport or synthesis of its cofactor, adenosyl-cobalamin (cblA, cblB, cblC, cblF, cblD and cblX), or a deficiency of the enzyme methylmalonyl-CoA epimerase. MCM is a mitochondrial enzyme that catalyzes the isomerization of methylmalonyl-CoA to succinyl-CoA.

Isolated MMA is found in patients with mutations in MUT, located on chromosome 6p21, causing partial (mut<sup>-</sup>) or complete (mut<sup>0</sup>) enzyme deficiency (1). In general, the mut forms of MMA is unresponsive to vitamin B12 therapy. MCM activity requires 5-primedeoxyadenosylcobalamin (AdoCbl), a coenzyme form of vitamin B12. Patients with defects in the synthesis of AdoCbl are usually responsive to vitamin B12 therapy and are classified as 'cbl' type. The cblA type is caused by mutations in the MMAA gene on 4q31. MMAA is involved in the synthesis of adenosylcobalamin (AdoCbl), a coenzyme for MCM. The cblB type is caused by mutations in the MMAB gene on 12q24. MMAB encodes cobalamin adenosyl transferase (ATR), which catalyzes transfer of an adenosyl group from ATP to cobalamin (I) to form AdoCbl (2, 3).

Combined MIMA and homocystinuria is a genetically heterogeneous disorder of cobalamin (cbl; vitamin B12) metabolism. The defect causes decreased levels of the coenzymes adenosylcobalamin (AdoCbl) and methylcobalamin (MeCbl), which result in decreased activity of the respective enzymes MCM and methyltetrahydrofolate homocysteine methyl transferase, also known as methionine synthase (MTR). Different forms of the disorder have been classified according to complementation groups of cells in vitro: cblC, cblD, cblX and cblF. Members of

Downloaded from rbmb.net on 2024-05-03

<sup>1:</sup> Immunobiochemistry Lab, Immunology Research Center, School of medicine, Mashhad University of Medical Sciences, Mashhad, Iran. 2: Pardis Clinical and Genetic Laboratory, Mashhad, Iran.

<sup>3:</sup> Department of Medical Genetics, Faculty of Medicine, Tehran University of medical sciences, Tehran, Iran

<sup>4:</sup> Immunobiochemistry Lab, Allergy Research Center, Mashhad University of Medical Sciences, Mashhad, Iran

<sup>5:</sup> Varastegan Institute for Medical Sciences, Mashhad, Iran.

<sup>\*</sup>Corresponding author: Abdol-Reza Varasteh; Tel: +98 51-38 44 20 16; Fax: +98 51-3845 22 36; E-mail: varasteha@mums.ac.ir Received: Nov 22, 2015; Accepted: Jan 20, 2016

complementation group cbID all contain homozygous or compound heterozygous mutations in the MMADHC gene, located on chromosome 2q23. The cbIC type of combined MMA and homocystinuria is caused by homozygous or compound heterozygous mutations in the MMACHC gene located on chromosome 1p34. MMA and homocystinuria, cbIC type, is the most common inborn error of cobalamin metabolism, with about 250 known cases (4). The cbIF type is caused by homozygous or compound heterozygous mutations in the LMBRD1 gene on chromosome 6q13 (5). The cbIX type is an X-linked, Xq28, recessive metabolic disorder that is caused by mutations in the HCFC1 gene.

Also deficiency of the enzyme methylmalonyl-CoA epimerase and ADP-forming succinyl-CoA synthetase (SCS-A, EC 6.2.1.5) is found in patients with MMA. Deficiency of the enzyme methylmalonyl-CoA epimerase is caused by mutation in the MCEE gene on chromosome 2p13.3 and deficiency of the SCS is caused by mutation in the SUCLA2 gene on chromosome 13q14.2. SCS is a mitochondrial matrix enzyme that catalyzes the reversible synthesis of succinyl-CoA from succinate and CoA. Fig. 1 shows the genes involved in the propionyl-CoA to succinyl-CoA conversion pathway.

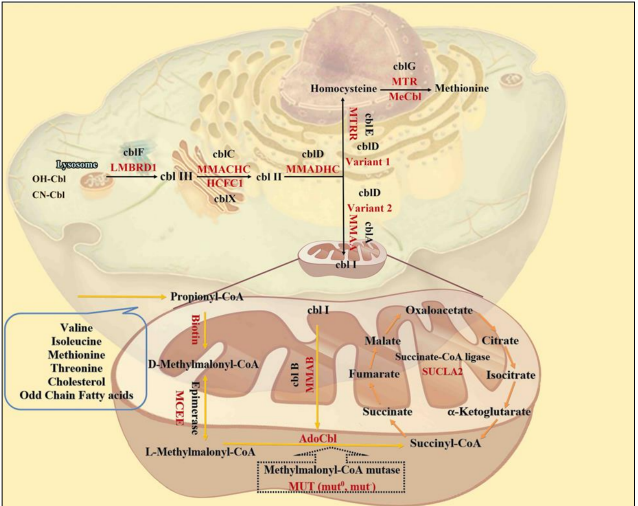


Fig. 1. The propionyl-CoA to succinyl-CoA conversion pathway

Onset of the manifestations of isolated MMA ranges from the neonatal period to adulthood. All phenotypes demonstrate intermittent periods of relative health and metabolic decompensation, usually associated with intercurrent infections and stress. In the neonatal period the disease can present with lethargy, vomiting, hypotonia, hypothermia, respiratory distress, severe ketoacidosis, hyperammonemia, neutropenia, and thrombocytopenia, and can result in death. In the infantile/non-B12-responsive phenotype, the most common form, infants are normal at birth but develop lethargy, vomiting, dehydration, hepatomegaly, hypotonia, and encephalopathy. An intermediate B12responsive phenotype occasionally presents in neonates, but usually presents in the first months or years of life; affected children exhibit anorexia, failure to thrive, hypotonia, and developmental delay, and sometimes have protein aversion and/or vomiting and lethargy after protein intake.

Atypical and "benign"/adult MMA are associated with increased, albeit mild, urinary excretion of methylmalonate; however, it is uncertain whether individuals with these conditions will develop symptoms. Major secondary complications of MMA include developmental delay (variable), tubulointerstitial nephritis with progressive renal failure, "metabolic stroke" (acute and chronic basal ganglia involvement), disabling movement disorder with choreoathetosis, dystonia and para/quadriparesis, pancreatitis, growth failure, functional immune impairment, and optic nerve atrophy. This review describes the biochemical and molecular genetics methods used to diagnosis isolated MMA.

#### Diagnosis of MMA

Acidosis, ketosis, hyperammonemia, hypoglycemia, hyperglycemia, and neutropenia are symptoms of methylmalonic and propionic acidemia (PA). Laboratory findings suggestive of MMA and PA include low bicarbonate levels less than 22 mmol/l in infants and less than 17 mmol/l in neonates, ketones in the urine, blood ammonia levels greater than150 µg/dl in neonates, 70 µg/dl in infants, and 35-50 µg/dl in older children and adults, blood glucose levels less than 40 mg/ml in infants and less than 60 mg/ml in children, and absolute neutrophil counts less than 1,500/mm<sup>3</sup> (6). Acylcarnitine levels in patients with any of the listed laboratory findings should be determined by tandem mass spectroscopy (MS/MS). Patients should also be screened for PA/MMA based on C3 and C3:C2 ratios. Patients with suspected disorders of cobalamin or propionate metabolism have C3 values greater than 7 µmol/l and C3:C2 ratios greater than 0.2; however, such screening cannot distinguish between MMA and PA. Table 1 shows the differential diagnoses of MMA and PA.

	Amino Acids pathway affected	Enzyme	Clinical Symptoms	Routine Laboratory Investigation	Plasma Amino Acids (Acylcarnitine profile)	Urine Organic Acids
Methyl malonic Acidemia (MMA)	Isoleucine, valine, methionine, threonine	Methylmalo nyl CoA mutase	Acidosis, ketosis, hyperammonemi a, hypoglycemia, hyperglycemia, and neutropenia	Low bicarbonate level, positive ketones in the urine, blood ammonia levels, low blood glucose levels	Methylmalonic acid in blood, Acyl carnitines, increased glycine in blood	Methylmalonic acid and methyl citrate in urine
Propionic Acidemia (PA)	Isoleucine, valine, methionine, threonine	Propionyl CoA carboxylase	Acidosis, ketosis, hyperammonemi a, hypoglycemia, hyperglycemia, and neutropenia	Low bicarbonate level, positive ketones in the urine, blood ammonia levels, low blood glucose levels	Propionylcarnitine, increased glycine in blood	Propionic acid, 3- OH propionic acid, methyl citrate, propionyl glycine in urine

Table 1. Differential diagnosis of MMA and PA

A definitive diagnosis of the disorder is based on urine organic acid analysis using gas chromatography/mass spectrometry (GC/MS) (6, 7). Organic acids can be measured using any body fluid, but urine is the most efficient for determining the type of disorder. The determination of organic acids and glycine conjugates in urine is key for the diagnosis and follow-up of MMA. Urine collected over 24 h allows for variations in volume excretion during the day. The impracticality of 24 h collection is, however, such that a random specimen, preferably the first morning voiding, an acceptable alternative. Intra-individual variations will occur with respect to the time of sampling, the patient's clinical status and diet, and whether the sample is collected when the patient is fasted or fed. Sampling during fasting or metabolic decompensation is often considered to be most valuable because, in most cases, metabolites of interest are then excreted selectively or at higher concentrations. Two factors can increase excretion of organic acids; first, an increase in excretion may be nonspecific because some metabolites are reported to be abnormally excreted in conditions not attributable to IEM, such as drug therapy, diet, non-IEM diseases, or physiologic conditions. A second common

misinterpretation may arise from bacterial metabolism. Of possible endogenous origin, such as in intestinal infections, is the abnormal excretion of d-lactate, methylmalonate, phydroxyphenylacetate, p-hydroxyphenyllactate, glutarate, benzoate, and hippurate. A urinary organic acid profile is nearly always abnormal during the acute illness phase, but it is commonly barely detectable before or between crises; therefore, it is important to obtain a urine sample for testing during the peak of a crisis. Urine organic acid levels from patients with MMA contain relatively high concentrations of methylmalonic acid and methyl citrate, whereas urine from patients with PA will show relatively high concentrations of metabolites of propionyl CoA, including propionic acid, methyl citric acid, 3-OH propionic acid, and propionyl glycine (6). Table 2 shows methylmalonic acid concentrations in urine for different subtypes of MMA.

 Table 2. Methylmalonic acid concentrations in urine for different subtypes of MMA.

	Methylmalonic acid concentration				
Subtype	Urine (n	Blood (µM)			
mut <sup>-</sup> , mut <sup>0</sup>		1000-10000	100-1000		
cblB, cblA, cl	bID	10-100	5-100		
MCEE defic SUCLA2	iency,	50-1500	7		
Normal		<4	<0.27		

Plasma amino acids in MMA contain carnitines, whereas those in PA contain propionylcarnitine. Both can have high glycine even when well controlled (6). Also, plasma homocysteine can be measured to identify gene types involved in MMA. After urine organic acid analysis and determination of plasma homocysteine concentration, patients are diagnosed based on one of the following criteria:

a) Patients with very high concentrations of methylmalonic acid in urine, but normal homocysteine, have mutations in at least one of the MUT (mut<sup>-</sup>, mut<sup>0</sup>), cblB, cblA and cblD (var 2) subtypes. MMA subtypes are diagnosed by enzyme assay analysis and/or molecular studies. Molecular genetics techniques are available for carrier testing of family members to aid in reproductive decision making and to determine whether prenatal testing is necessary (8).

b) Patients with slightly elevated methylmalonic acid in urine, but normal homocysteine, have mutations in at least one of the MCEE, SUCLA2 and benign MMA subtypes. Subtypes are diagnosed by enzyme assay analysis and/or molecular study.

c) Patients with abnormally high concentrations of methylmalonic acid in urine and homocysteine in plasma have mutations in at least one of the cblC, cblF, or cblD (var 1) subtypes. Subtypes are diagnosed by mutation analyses of MMACHC, MMADHC, and LMBRD1. Fig. 2 shows a flowchart for different stages of MMA diagnosis.

## 1. Acylcarnitine profile analysis

In a subset of newborn diseases with severe metabolic disorders, irreversible damage with adverse lifelong consequence may occur. For some of these diseases, a diagnostic method may help to prevent such damages. A critical screening technique used to detect many of these metabolic disorders in newborns is tandem mass spectrometry (MS/MS). MS/MS has the potential to simultaneously detect and quantify many physicochemical metabolites with similar properties. This constitutes a dramatic advance over the classical methods used for newborn screening. MS/MS has improved the detection of inborn errors of metabolism in newborns by making the analysis more sensitive, specific, and reliable than was previously possible. Its inherent ability to detect and quantify multiple metabolites in a single sample permits wide recognition of amino acid, fatty acid, and organic acid disorders.

Many patients have less severe symptoms if diagnosed and treated early. In addition, early diagnosis can decrease medical expenses and allow family planning to be considered before other affected siblings are born. Therefore, MS/MS can provide considerable benefits to patients and their families if integrated into newborn screening programs, provided that adequate funding is made available to cover the costs of the additional medications and foods. One of the disorders that can be diagnosed following MS/MS is MMA. The screening program identifies children that may be considered at risk for these disorders. Newborn screening for PA/MMA based on C3 and C3:C2 was begun in New York State in November 2004. The diagnoses are made at specialty care centers based on test results and evaluation by metabolic specialists. In May 2005, methylmalonyl carnitine (C4DC) was added to the newborn screening panel as a secondary marker for PA/MMA. In some instances, samples were referred to rule out disorders of propionate metabolism on the basis of persistently elevated C4DC. Until 2008, a protocol was used in New York State that referred patients with suspected disorders of cobalamin or propionate metabolism with C3 concentrations greater than7 µmol/l and C3:C2 ratios greater than 0.2 (9).

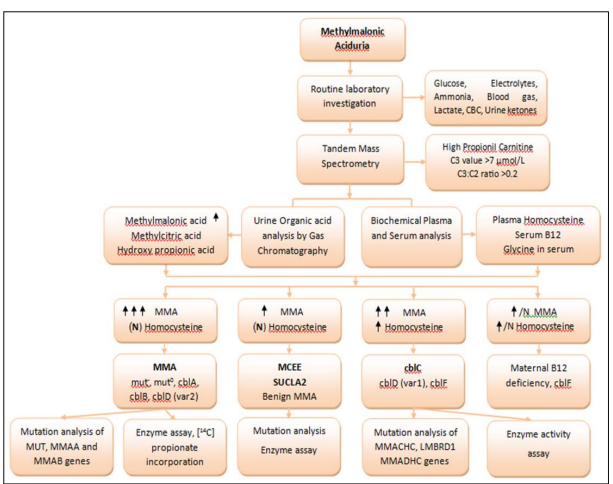


Fig. 2. A flowchart for different stages of MMA diagnosis

#### 2. Organic acid analysis

In the hereditary diseases known as organic acidemia, an enzyme or co-factor defect in a metabolic pathway leads to the accumulation and increased excretion of one or more of these acidic metabolites in urine. Therefore, human urine contains numerous organic acids and other chemical compounds at a variety of concentrations (10) and urinary organic acid analysis has become an important tool for laboratories involved in the diagnosis of these inherited metabolic disorders. Gas chromatography (GC) is the technique of choice to separate and identify more than 250 normal and pathological acidic metabolites detected in these diseases (11-13).

In 1963, Cox and White demonstrated an increase in urinary excretion of methylmalonic acid in patients with vitamin B12 deficiencies (14). Since then numerous methods have been described for the determination of urinary methylmalonic acid using colorimetry or GC (15, 16). Some of these methods utilize pre-purification of methylmalonic acid by thin-layer chromatography or ion-exchange resins in conjunction with calorimetric procedures or GC.

They were sensitive enough to measure a greatly increased amount of methylmalonic acid, but accurate determination of small amounts of methylmalonic acid in biological specimens has been difficult (17).

Millar et al (1974) described an improved method based on gas-liquid chromatography (GLC) of the butyl ester of methylmalonic acid, which was produced by reacting a diethyl ether extract of urine or freeze-dried urine with a mixture of boron trifluoride and butanol. Therefore, methylmalonic acid was directly extracted from urine and measured as its butyl ester (18). Tanaka et al. (1980) described a practical gas-chromatographic method of urinary organic acid analysis that was designed to be used in organic acidemia screening programs. This method involves extraction of urine with ethyl acetate, dehydration of the extracted residues. trimethylsilylation, and use of the list of retention indices to identify the organic acids (19).

In 1981 Maties et al. modified Millar's method and adapted it for use with urine specimens absorbed into filter paper. The advantage of this method was that methylmalonic acid was quantitated with acceptable confidence from the small amount of urine present on filter paper specimens that were easily collected and mailed to a central testing laboratory. This technique was also applicable to the detection of other types of organic acidemias (20).

In 1983, Hyman et al. described a rapid method for MMA detection that utilizes DEAE-cellulose paper for sample collection and diazotized pnitroaniline for color development. Using anionexchange filter paper, urine organic acids are selectively adsorbed to the paper. The color reaction with diazotized p-nitroaniline, which takes place in situ, is more sensitive than a reaction in solution, and substances producing interfering colors with the reagent can be rinsed from the disc prior to the assay, making the assay more accurate (21).

Nakamura et al. (1987) described a method for microanalysis of short chain dicarboxylic acids including methylmalonic, succinic, and methylsuccinic acids, which consists of prefractionation of the dicarboxylic acid fraction by ion-exchange chromatography, extraction of the eluate with ethyl acetate, and analyses of dicarboxylic acids as dimethyl esters by GC. They suggested that small amounts of these dicarboxylic acids in normal human urine, amniotic fluid, and serum can be accurately measured with this method (17).

Verhaeghe et al. (1988) developed a method that combines the specificity, reproducibility, and high extraction yield of anion-exchange chromatography with the speed and simplicity of solvent extraction using a gas chromatography-flame ionization detector (GC-FID) to measure urine organic acids. They suggested that this convenient procedure is selective, reproducible, and a suitable alternative to the more cumbersome diethylaminoethyl-Sephadex extraction method (10).

Hoffman et al. (1989) described a procedure for analysis for organic acids in various biological samples that incorporates the O-(2,3,4,5,6pentafluorobenzyl) oxime-trimethylsilyl (O -PFBOxime-TMS) esters of oxoacids, aldehydes, and ketones. The gas-chromatographic properties of the O-PFEOxime-TMS esters have distinct advantages over the commonly used O-ethoxime-TMS esters; each is processed in a manner identical to that for aqueous standards, and requires no deproteinization. They suggested that there are no limitations on sample volumes and it is likely that cerebrospinal fluid and homogenized tissue samples can also be analyzed without further modifications. Proteins and peptides, as well as basic and polar low-molecular mass compounds, such as amino acids, inorganic acids, creatinine, purines, amines, sugars, and urea are retained on the silicic acid column. An additional advantage is that orotate and uracil, compounds valuable in distinguishing some urea-cycle disorders, are efficiently extracted (22).

One of the most critical points in a metabolicprofiling scheme is the isolation of the compounds of interest from the biological matrix. These should be extracted in high, uniform, and reproducible yields, accompanied by as few compounds as possible from other product classes. Because organic acids cover such a wide range of polarity and have different chemical properties related to the various functional groups present, this requirement has been difficult to meet. Moreover, for a procedure to be useful as a routine diagnostic method, one should be able to process several samples simultaneously with reasonable accuracy and speed.

6

Methods currently in use consist of isolating acidic constituents from urine before derivatization and GC. These include anion-exchange chromatography based on organic polymers or cellulose and solvent extraction with ethyl acetate or diethyl ether. The principal advantage of diethylaminoethyl-Sephadexcolumn

chromatography is its specificity and high and reproducible extraction recovery of polar and nonpolar acids. This method is strongly recommended for quantitative monitoring and for recognizing subtle changes in excretion profiles. Disadvantages of this method are that it is a laborious and complex procedure and the profile may be obscured by dominant peaks of some polar acids and inorganic sulfate and phosphate, which can mask some important organic acids in the gas chromatogram. These short comings impede diagnoses of metabolic diseases, especially in instances in which the increases in organic acid metabolites are small, as in vitamin-responsive organic acidopathies. Because of these practical difficulties, many groups have found solvent extraction to be an attractive approach for routine diagnoses of organic acidemias. However, solvent extraction is more widely used because it is fast and simple and yields an adequate recovery of aromatic and less-polar aliphatic acids with minimal coextraction of sulfate and phosphate (10).

Solvent extraction has serious limitations imposed by the low and unreproducible extractability of polar acids such as 3hydroxyisovalerate, 3-hydroxypropionate, methyl citrate, and citrate (19).

In addition, an organic acid profile enables one evaluate metabolic to disorders pathobiochemically on the basis of their relations to one another as precursors or products. Therefore, careful consideration of small changes in organic acid ratios is essential for accurate diagnosis and optimal management for prognosis after treatment in addition to quantitative determination of organic acids. GC-MS is more specific, in that quantification is based on the relative intensities of characteristic fragment ions in a reconstructed ion chromatogram. Mass chromatography usually vields lower precision and sensitivity than GC-FID detection, and single- or multiple-ion monitoring can be used to quantify only a few target compounds. With GC-FID detection, on the other hand, compounds that are 100- to 1000-fold less concentrated than the major components of the sample can still be quantified (10). GC/MS requires expensive instrumentation, and maintenance, operation, and data interpretation require highly-specialized training and technical expertise. In addition, a computer is almost indispensable for data processing. Thus, organic acidemia has been screened in only a few major medical centers, where such instruments and expertise are available.

In conclusion, at present, solvent extraction with ethyl acetate, diethyl ether, or both, is widely used. This type of liquid extraction yields poor analytical recoveries of the more-polar compounds and is inconvenient for use with large numbers of samples. Also, GC-MS has become a wellestablished, easily automated, and reliable technology in the research field of metabolomics (23). Human urine contains many metabolites, and GC-MS analysis of urinary organic acids is an important technique for the diagnosis of inborn errors of lipid, amino acid, and carbohydrate metabolism (24). By means of urease pretreatment of urine samples and other methodological improvements, GC-MS has been applied to simultaneously analyze the numerous metabolic intermediates of multiple categories in urine, providing diagnostic evidence for more than 130 inborn errors of metabolism (IEM) (25-27).

## 3. Enzyme activity assay

MMA may be diagnosed by measuring MCM activity, with or without the addition of AdoCbl. This can be used to distinguish between two MMA variants (Cbl-responsive and Cbl-unresponsive) and distinguish between two MUT subtypes (mutand mut<sup>0</sup>). Thus, the in vitro measurement of MCM activity, with and without AdoCbl, is useful to investigate the Cbl pathway, diagnose MMA, identify MUT and cbl mutations, and gain insight into the biochemical changes accompanying vitamin B12 deficiencies. Methods described and employed to measure MCM activity include radiometric methods, in which [14C] succinyl-CoA is produced and separated from the substrate DL[CH3-14C]methylmalonyl-CoA by paper chromatography (28,29), thin layer

chromatography (30, 31), electrophoresis (32), potassium permanganate oxidation (33, 34), microwell filtration (35), extraction into ethyl acetate (36), high performance liquid chromatography (HPLC) (37) and GC (38). There are also nonradioactive assays based on the separation of methylmalonyl-CoA and succinyl-CoA by reverse-phase HPLC (39, 40), or on the direct spectrophotometric assay of succinyl-CoA (41-43). The first six methods are reputed to be laborious because they require many manipulations, and time-consuming because of the numerous incubations, and they have been criticized for their lack of sensitivity. The permanganate oxidation method is also criticized because the optimal conditions for oxidation vary depending on the permanganate concentration and heating time. The gas chromatographic radiometric assay method appears to be sensitive, but is also time-consuming. The lack of sensitivity and reproducibility, and the inconvenience of the radiometric assay for MCM activity make HPLC the method of choice. Therefore. the nonradioactive HPLC assay seems to be satisfactory for measuring the conversion of small fractions of methylmalonyl-CoA to succinyl-CoA, and is said to be simple, rapid, reliable, and highly reproducible. This method is sufficiently sensitive to measure low MCM activity, such as the holo-MCM activity in tissue extracts, or the total MCM activity of cells. We therefore believe this method is suitable for detecting abnormal MCM apoenzyme, whether to diagnose MMA or detect errors of cobalamin metabolism (44).

## 4. Mutation analysis *MUT*

MMA is an inborn error of metabolism due to the impaired isomerization of L-methylmalonyl-CoA to succinyl-CoA. This reaction is catalyzed by the mitochondrial protein MCM, an adenosylcobalamin-dependent enzyme (45). The human MUT gene, located on chromosome 6, is comprised of 13 exons spanning over 35 kb. The open reading frame consists of 2.7 kb, encoding 750 amino acids (46). Two classes of mutations in MUT are classically distinguished by studies of [14C]-propionate metabolism in primary fibroblasts from patients with MMA (47). Mut<sup>0</sup>

mutations result in no detectable MCM activity. Mut<sup>-</sup> mutations result in low residual enzyme activity. The human MUT gene was identified by Ledley et al., who screened an expression library with mutase antibodies to isolate the first human cDNA (48). Over the last 27 years a number of studies have described the spectrum of mutations observed at the MUT locus in human patients (45, 49-57). To date, 272 different mutations have been identified, including 187 missense/nonsense mutations, 24 splice-site mutations, 37 small deletions, 20 small insertions, three small indels, and one gross deletion.

## cblC

MMA, cobalamin deficiency type C (cblC) with homocystinuria (MMACHC gene) is the most common genetic defect in cobalamin metabolism (4, 58). The MMACHC gene responsible for cblC disorder is located on chromosome 1p34.1 and encodes a polypeptide of 282 amino acids. Exons 1-4 are coding and exon 5 is non-coding. The MMACHC protein may act as an intracellular cobalamin-trafficking chaperone and has been shown to act, in part, catalyzing the reductive decyanation of cyanocobalamin, generating cob(II)alamin, which is the substrate for assimilation into the active cofactor forms methylcobalamin (MeCbl) and adenosylcobalamin (AdoCbl) (59). To date, mutation analyses of MMACHC (60, 61) have shown 81 different mutations, which include 43 missense/nonsense mutations, five splice site mutations, 19 small deletions, eight small insertions, one small indel, four gross deletions, and one gross insertion.

## cblA

The cblA type is caused by mutations in the MMAA gene on 4q31. MMAA is involved in the synthesis of adenosylcobalamin (AdoCbl) (2). Two different enzymatic functions have been identified for the MMAA gene product: a role in vitamin B12 transport into the mitochondria, reduction of cobalamin II to cobalamin I, and the conservation or re-activation of MCM. Multiple mutations in various regions of the gene have been identified, which will help guide future structure and function studies (62, 63). The mutations includ

8

30 missense/nonsense mutations, four splice site mutations, five small deletions, five small insertions, and one gross deletion.

## cblB

In cblB type of MMA, the defective gene is MMAB. The MMAB gene, on chromosome 12q24.1, encodes the mitochondrial enzyme ATP: cobalamin adenosyl transferase (ATR), which catalyzes transfer of an adenosyl group from ATP to cobalamin (I) to form AdoCbl (3). Mutation analysis of the MMAB gene identified 20 missense/nonsense mutations, seven splice site mutations, two regulatory mutation, five small deletions, three small insertions, and one small indel (63, 64).

#### MCEE

D-methylmalonyl-CoA (D-MMCoA) is formed as a product of the propionyl-CoA carboxylase reaction. D-MMCoA requires racemization prior to becoming a substrate for the MCM reaction, and a deficiency of D-methylmalonyl-CoA racemase (MCR, EC 5.1.99.1) has long been postulated as a potential etiology of hereditary MMA (65). The epimerase gene (MCEE) on chromosome 2p13.3 was the first cobalamin-related gene to be identified on the basis of prokaryotic gene arrangements (66). То date. three missense/nonsense mutations have been identified.

## Other subtypes

At least four other genetic entities pathways can be associated with isolated MMA. cblD types are homozygous compound caused by or heterozygous mutations in the MMADHC gene, found on chromosome 2q23; these include eight missense/nonsense mutations, two small deletions, and three small insertions (67). Function of the product of this gene remains unknown; it shows homology to the putative ATPase component of a bacterial ABC transporter. Mutations in the Cterminal region were identified in patients with cblD variant 2, mutations in the N-terminal region were identified in patients with cblD variant 1, and truncating mutations were associated with the classic cbID phenotype. This supports suggestions that the MMADHC gene product plays a role in directing cobalamin to the 2 cobalamin-dependent enzymes of mammalian cells, Methylmalonyl CoA mutase and Methionine synthase (68).

cblF type is caused by homozygous or compound heterozygous mutations in the LMBRD1 gene, found on chromosome 6q13. This gene produces a lysosomal cobalamin transporter protein that facilitates lysosomal cobalamin export (69). To date, no mutations have been reported for this gene. The SUCLA2 gene encodes the betasubunit of the ADP-forming succinyl-CoA synthetase (SCS-A; EC 6.2.1.5). SCS is a mitochondrial matrix enzyme that catalyzes the reversible synthesis of succinyl-CoA from succinate and CoA. The reverse reaction occurs in the Krebs cycle, while the forward reaction may produce succinyl-CoA for activation of ketone bodies and heme synthesis. To date, ten missense/nonsense mutations, one splice site mutation, one small insertion, one gross deletion and one small indel have been identified (70). The cblX type is an X-linked (Xq28) recessive metabolic disorder characterized by severely delayed psychomotor development apparent in infancy and is caused by mutations in the HCFC1 gene. Mutation in HCFC1 gene inhibits its function in the transcriptional activation of MMACHC gene and showed that disorder of transcription can cause an inborn error of metabolism (71). To date, no mutations have been reported for this gene. Gene subtypes and mutations involved in MMA are shown in Table 3.

## Acknowledgments

This research was supported and funding by Immunobiochemistry Lab, Allergy Research Center, Medical School, Mashhad University of Medical Sciences, Mashhad, Iran. All authors declare they have no conflicts of interest.

Subtype	Gene	ions in genes involved in M Mutation	Number of mutation	Percentage
		Missense/nonsense	187	58.37
		Splice site mutation	24	
Unresponsive to		Small deletion	37	
vitamin B12	MUT	Small insertion	20	
		Small indel	3	
		Gross deletion	1	
		Missense/nonsense	30	9.65
		Splice site mutation	4	
cblA	MMAA	Small deletion	5	
		Small insertion	5	
		Gross deletion	1	
		Missense/nonsense	20	8.15
		Splice site mutation	7	
L ID		Regulatory	2	
cblB	MMAB	Small deletion	5	
		Small insertion	3	
		Small indel	1	
		Missense/nonsense	43	17.39
		Splice site mutation	5	
		Small deletion	19	
cblC	MMACHC	Small insertion	8	
		Small indel	1	
		Gross deletion	4	
		Gross insertion	1	
		Missense/nonsense	8	2.79
cblD	MMADHC	Small deletion	2	
		Small insertion	3	
		Missense/nonsense	10	3.01
		Splice site mutation	1	
SUCLA2	SUCLA2	Gross deletion	1	
		Small indel	1	
		Small insertion	1	
cblF	LMBRD1	Not reported	-	0
cblX	HCFC1	Not reported	-	0
MCEE	MCEE	Missense/nonsense	3	0.64

Table 3. Subtypes and mutations in genes involved in MMA (HGMD data base)

## References

1. Bell CG, Ledley FD, Lumetta MR, Zoghbi HY, VanTuinen P, Ledbetter SA, Ledbetter DH. Mapping of human methylmalonyl CoA mutase (MUT) locus on chromosome 6. American journal of human genetics. 1988;42(6):839-46.

2. Dobson CM, Wai T, Leclerc D, Wilson A, Wu X, Dore C, et al. Identification of the gene responsible for the cblA complementation group of vitamin B12responsive MMA based on analysis of prokaryotic gene arrangements. Proceedings of the National Academy of Sciences of the United States of America. 2002;99(24):15554-9.

3. Dobson CM, Wai T, Leclerc D, Kadir H, Narang M, Lerner-Ellis JP, et al. Identification of the gene responsible for the cblB complementation group of vitamin B12-dependent methylmalonicaciduria. Human molecular genetics. 2002;11(26):3361-9.

4. Lerner-Ellis JP, Tirone JC, Pawelek PD, Dore C, Atkinson JL, Watkins D, et al. Identification of the gene responsible for methylmalonicaciduria and homocystinuria, cblC type. Nature genetics. 2006;38(1):93-100.

5. Suormala T, Baumgartner MR, Coelho D, Zavadakova P, Kozich V, Koch HG, et al. The cblDdefect causes either isolated or combined deficiency of methylcobalamin and adenosylcobalamin synthesis. The Journal of biological chemistry. 2004;279(41):42742-9.

6. Seashore MR. The Organic Acidemias: An Overview. In: Pagon RA, Adam MP, Ardinger HH, Wallace SE, Amemiya A, Bean LJH, et al., editors. Gene Reviews. Seattle WA: University of Washington, Seattle; 1993.

7. Manoli I, Venditti CP. MMA. In: Pagon RA, Adam MP, Ardinger HH, Wallace SE, Amemiya A, Bean LJH, et al., editors. Gene Reviews(R). Seattle WA: University of Washington, Seattle; 1993.

8. Van Gosen L. Organic acidemias: a methylmalonic and propionic focus. Journal of pediatric nursing. 2008;23(3):225-33.

9. Weisfeld-Adams JD, Morrissey MA, Kirmse BM, Salveson BR, Wasserstein MP, McGuire PJ, et al. Newborn screening and early biochemical followup in combined methylmalonicaciduria and homocystinuria, cblC type, and utility of methionine as a secondary screening analyte. Molecular genetics and metabolism. 2010;99(2):116-23. 10. Verhaeghe BJ, Lefevere MF, De Leenheer AP. Solid-phase extraction with strong anion-exchange columns for selective isolation and concentration of urinary organic acids. Clinical chemistry. 1988;34(6):1077-83.

11. Liebich H. Analysis of acidic metabolites by capillary column GC and GC/MS. Journal of High Resolution Chromatography 1983;6(12):640-50.

12. Holland JF, Leary JJ, Sweeley CC. Advanced instrumentation and strategies for metabolic profiling. Journal of chromatography. 1986;379:3-26.

13. Tuchman M, Bowers LD, Fregien KD, Crippin PJ, Krivit W. Capillary gas chromatographic separation of urinary organic acids. Retention indices of 101 urinary acids on a 5% phenylmethyl silicone capillary column. Journal of chromatographic science. 1984;22(5):198-202.

14. Cox EV, White AM. Methylmalonic acid excretion: an index of vitamin-B12 deficiency. Lancet. 1962;2(7261):853-6.

15. Giorgio AJ, Plaut GW.A method for the colorimetric determination of urinary methylmalonic acid in pernicious anemia. The Journal of laboratory and clinical medicine. 1965;66(4):667-76.

16. Gompertz D. The measurement of urinary methylmalonic acid by a combination of thin-layer and gas chromoatography. Clinica chimica acta; International journal of clinical chemistry. 1968;19(3):477-84.

17. Nakamura E, Rosenberg LE, Tanaka K. Microdetermination of methylmalonic acid and other short chain dicarboxylic acids by gas chromatography: use in prenatal diagnosis of MMA and in studies of isovaleric acidemia. Clinica chimica acta; International journal of clinical chemistry. 1976;68(2):127-40.

18. Millar KR, Lorentz PP. A gas chromatographic method for the determination of methylmalonic acid in urine. Journal of Chromatography A. 1974;101(1):177-81.

19. Tanaka K, West-Dull A, Hine DG, Lynn TB, Lowe T. Gas-chromatographic method of analysis for urinary organic acids. II. Description of the procedure, and its application to diagnosis of patients with organic acidurias. Clinical chemistry. 1980;26(13):1847-53. 20. Maties M, Shih VE, Evans J, Levy HL. Measurement of methylmalonic acid in urine filter paper specimens by gas chromatography. Clinica chimica acta; International journal of clinical chemistry. 1981;114(2-3):303-8.

21. Hyman DB, Saunders AM, Tanaka K. A rapid spot test for urinary methylmalonic acid collected on ion-exchange filter paper. Clinica Chimica Acta. 1983;132(3):219-27.

22. Hoffmann G, Aramaki S, Blum-Hoffmann E, Nyhan WL, Sweetman L. Quantitative analysis for organic acids in biological samples: batch isolation followed by gas chromatographic-mass spectrometric analysis. Clinical chemistry. 1989;35(4):587-95.

23. Nobeli I, Thornton JM. A bioinformatician's view of the metabolome. BioEssays: news and reviews in molecular, cellular and developmental biology. 2006;28(5):534-45.

24. Hori D, Hasegawa Y, Kimura M, Yang Y, Verma IC, Yamaguchi S. Clinical onset and prognosis of Asian children with organic acidemias, as detected by analysis of urinary organic acids using GC/MS, instead of mass screening. Brain & development. 2005;27(1):39-45.

25. Kuhara T. Noninvasive human metabolome analysis for differential diagnosis of inborn errors of metabolism.Journal of chromatography B, Analytical technologies in the biomedical and life sciences. 2007;855(1):42-50.

26. Kuhara T. Gas chromatographic-mass spectrometric urinary metabolome analysis to study mutations of inborn errors of metabolism. Mass spectrometry reviews. 2005;24(6):814-27.

27. Song Y-Z, Li B-X, Hao H, Xin R-L, Zhang T, Zhang C-H, et al. Selective screening for inborn errors of metabolism and secondary methylmalonicaciduria in pregnancy at high risk district of neural tube defects: A human metabolome study by GC-MS in China. Clinical Biochemistry. 2008;41(7–8):616-20.

28. Reed EB, Tarver H. Urinary methylmalonate and hepatic methylmalonyl coenzyme A mutase activity in the vitamin B12-deficient rat. The Journal of nutrition. 1970;100(8):935-47. 29. Whitaker TR, Giorgio AJ. A direct radioassay of methylmalonyl-coenzyme A mutase using enzymatically synthesized dl-[3-14C] methylmalonyl-CoA. Analytical Biochemistry. 1973;52(2):522-32.

30. Willard HF, Rosenberg LE. Inherited deficiencies of human methylmalonylCaAmutase activity: reduced affinity of mutant apoenzyme for adenosylcobalamin. Biochemical and biophysical research communications. 1977;78(3):927-34.

31. Scott JS, TrestonAM, Bowman EP, Owens JA, Cooksley WG. The regulatory roles of liver and kidney in cobalamin (vitamin B12) metabolism in the rat: the uptake and intracellular binding of cobalamin and the activity of the cobalamin-dependent enzymes in response to varying cobalamin supply. Clinical science (London, England: 1979). 1984;67(3):299-306.

32. Morrow G, Barness LA, Cardinale GJ, Abeles RH, Flaks JG. Congenital MMA: enzymatic evidence for two forms of the disease. Proceedings of the National Academy of Sciences of the United States of America. 1969;63(1):191-7. Kolhouse JF, Utley C, Allen RH. Isolation and characterization of MUT from human placenta. The Journal of biological chemistry. 1980;255(7):2708-12.

33. Kolhouse JF, Utley C, Allen RH. Isolation and characterization of MUT from human placenta. The Journal of biological chemistry. 1980;255(7):2708-12.

34. Kolhouse JF, Stabler SP, Allen RH. L-MUT from human placenta. Methods in enzymology. 1988;166:407-14.

35. Kakinuma H, Kobayashi A, Takahashi H. 14Cpropionate incorporation assay by rapid filtration in multiwell plates. Clinicachimicaacta; International journal of clinical chemistry. 2004;343(1-2):209-12.

36. Cannata JJ, Focesi A, Jr., Mazumder R, Warner RC, Ochoa S. metabolism of propionic acid in animal tissues. xii. Properties of mammalian methylmalonyl coenzyme a mutase. The Journal of biological chemistry. 1965;240:3249-57.

37. Causey AG, Bartlett K. A radio-HPLC assay for the measurement of MUT. Clinica Chimica Acta. 1984;139(2):179-86. 38. Goodey PA, Gompertz D. Methylmalonyl CoA mutase--a radiochromatographic assay. Clinicachimicaacta; International journal of clinical chemistry. 1972;42(1):119-23.

39. Kikuchi M, Hanamizu H, Narisawa K, Tada K. Assay of methylmalonyl CoA mutase with highperformance liquid chromatography. Clinicachimicaacta; International journal of clinical chemistry. 1989;184(3):307-13.

40. Riedel B, Ueland PM, Svardal AM. Fully automated assay for cobalamin-dependent methylmalonyl CoA mutase. Clinical chemistry. 1995;41(8 Pt 1):1164-70.

41. Wood HG, Kellermeyer RW, Stjernholm R, Allen SHG. metabolism of methylmalonyl-coa and the role of biotin and B12 coenzymes. Annals of the New York Academy of Sciences. 1964;112(2):661-79.

42. Frenkel EP, Kitchens RL, Hersh LB, Frenkel R. Effect of vitamin B12 deprivation on the in vivo levels of coenzyme A intermediates associated with propionate metabolism. The Journal of biological chemistry. 1974;249(21):6984-91.

43. Watanabe F, Tamura Y, Saido H, Nakano Y. Enzymatic Assay for Adenosylcobalamin-dependent Methylmalonyl Coenzyme A Mutase. Bioscience, Biotechnology and Biochemistry. 2014;57(9):1593-4.

44. Gaire D, Sponne I, Droesch S, Charlier A, Nicolas J-P, Lambert D. Comparison of two methods for the measurement of rat liver methylmalonyl-coenzyme A mutase activity: HPLC and radioisotopic assays. The Journal of Nutritional Biochemistry. 1999;10(1):56-62.

45. Crane AM, Ledley FD. Clustering of mutations in methylmalonyl CoA mutase associated with mut-MMA. American journal of human genetics. 1994;55(1):42-50.

46. Nham SU, Wilkemeyer MF, Ledley FD.Structure of the human MUT (MUT) locus. Genomics. 1990;8(4):710-6.

47. Rosenberg LE, Fenton WA. Disorders of propionate metabolism. In: Scriver CR, Beaudet AL, Sly WS, Valle D (eds) The metabolic basis of inherited disease, 6th ed. McGraw-Hill, New York, 1989; pp 822-844.

48. Ledley FD, Lumetta M, Nguyen PN, Kolhouse JF, Allen RH. Molecular cloning of L-MUT: gene transfer and analysis of mut cell lines. Proceedings of the National Academy of Sciences of the United States of America. 1988;85(10):3518-21.

49. Ogasawara M, Matsubara Y, Mikami H, Narisawa K. Identification of two novel mutations in the MUT gene with decreased levels of mutant mRNA in MMA. Human molecular genetics. 1994;3(6):867-72.

50. Fuchshuber A, Mucha B, Baumgartner ER, Vollmer M, Hildebrandt F. mut0 MMA: eleven novel mutations of the methylmalonyl CoA mutase including a deletion-insertion mutation. Human mutation. 2000;16(2):179.

51. Benoist JF, Acquaviva C, Callebaut I, Guffon N, Ogier de Baulny H, Mornon JP, et al. Molecular and structural analysis of two novel mutations in a patient with mut(-) methylmalonyl-CoA deficiency. Molecular genetics and metabolism. 2001;72(2):181-4.

52. Peters HL, Nefedov M, Lee LW, Abdenur JE, Chamoles NA, Kahler SG, et al. Molecular studies in mutase-deficient (MUT) methylmalonicaciduria: identification of five novel mutations. Human mutation. 2002;20(5):406.

53. Acquaviva C, Benoist JF, Pereira S, Callebaut I, Koskas T, Porquet D, et al. Molecular basis of MUT apoenzyme defect in 40 European patients affected by mut(o) and mut- forms of MMA: identification of 29 novel mutations in the MUT gene. Human mutation. 2005;25(2):167-76.

54. Chandler RJ, Venditti CP. Genetic and Genomic Systems to Study MMA. Molecular genetics and metabolism. 2005;86(1-2):34-43.

55. Worgan LC, Niles K, Tirone JC, Hofmann A, Verner A, Sammak A, et al. Spectrum of mutations in mut MMA and identification of a common Hispanic mutation and haplotype. Human mutation. 2006;27(1):31-43.

56. Keeratichamroen S, Cairns JR, Sawangareetrakul P, Liammongkolkul S, Champattanachai V, Srisomsap C, et al. Novel mutations found in two genes of thai patients with isolated MMA. Biochemical genetics. 2007;45(5-6):421-30. 57. Sakamoto O, Ohura T, Matsubara Y, Takayanagi M, Tsuchiya S. Mutation and haplotype analyses of the MUT gene in Japanese patients with MMA. Journal of human genetics. 2007;52(1):48-55.

58. Rosenblatt DS, Wayne AF. Inherited disorders of folate and cobalamin transport and metabolism. In: Scriver CR, Beaudet AL, Sly WS, Valle D, editors. The metabolic and molecular basis of inherited diseases. New York: McGraw-Hill. 2001; p 3897–3933.

59. Kim J, Gherasim C, Banerjee R. Decyanation of vitamin B12 by a trafficking chaperone. Proceedings of the National Academy of Sciences of the United States of America. 2008;105(38):14551-4.

60. Richard E, Jorge-Finnigan A, Garcia-Villoria J, Merinero B, Desviat LR, Gort L, et al. Genetic and cellular studies of oxidative stress in methylmalonicaciduria (MMA) cobalamin deficiency type C (cblC) with homocystinuria (MMACHC). Human mutation. 2009;30(11):1558-66.

61. Carrillo-Carrasco N, Chandler RJ, Venditti CP. Combined MMA and homocystinuria, cblC type. I. Clinical presentations, diagnosis and management. Journal of inherited metabolic disease. 2012;35(1):91-102.

62. Morel CF, Watkins D, Scott P, Rinaldo P, Rosenblatt DS. Prenatal diagnosis for MMA and inborn errors of vitamin B12 metabolismand transport. Molecular genetics and metabolism. 2005;86(1-2):160-71.

63. Yang X, Sakamoto O, Matsubara Y, Kure S, Suzuki Y, Aoki Y, et al. Mutation analysis of the MMAA and MMAB genes in Japanese patients with vitamin B(12)-responsive MMA: identification of a prevalent MMAA mutation. Molecular genetics and metabolism. 2004;82(4):329-33.

64. Lemer-Ellis JP, Gradinger AB, Watkins D, Tirone JC, Villeneuve A, Dobson CM, et al. Mutation and biochemical analysis of patients belonging to the cblB complementation class of vitamin B12-dependent methylmalonic aciduria. Molecular genetics and metabolism. 2006;87(3):219-25.

65. Gradinger AB, Bélair C, Worgan LC, Li CD, Lavallée J, Roquis D, et al. Atypical methylmalonicaciduria: frequency of mutations in the methylmalonyl CoA epimerase gene (MCEE). Human mutation. 2007;28(10):1045.

66. Bobik TA, Rasche ME. Identification of the human methylmalonyl-CoA racemase gene based on the analysis of prokaryotic gene arrangements. Implications for decoding the human genome. The Journal of biological chemistry. 2001;276(40):37194-8.

67. Plesa M, Kim J, Paquette SG, Gagnon H, Ng-Thow-Hing C, Gibbs BF, et al. Interaction between MMACHC and MMADHC, two human proteins participating in intracellular vitamin B(1)(2) metabolism. Molecular genetics and metabolism. 2011;102(2):139-48.

68. Miousse IR, Watkins D, Coelho D, Rupar T, Crombez EA, Vilain E, et al. Clinical and molecular heterogeneity in patients with the cbID inborn error of cobalamin metabolism. The Journal of pediatrics. 2009; 154(4):551-6.

69. Gailus S, Suormala T, Malerczyk-Aktas AG, Toliat MR, Wittkampf T, Stucki M, et al. A novel mutation in LMBRD1 causes the cblF defect of vitamin B(12) metabolism in a Turkish patient. Journal of inherited metabolic disease. 2010;33(1):17-24.

70. Carrozzo R, Dionisi-Vici C, Steuerwald U, Lucioli S, Deodato F, Di Giandomenico S, et al. SUCLA2 mutations are associated with mild methylmalonic aciduria, Leigh-like encephalomyopathy, dystonia and deafness. Brain : a journal of neurology. 2007;130(Pt 3):862-74.

71. Yu HC, Sloan JL, Scharer G, Brebner A, Quintana AM, Achilly NP, et al. An X-linked cobalamin disorder caused by mutations in transcriptional coregulator HCFC1. American journal of human genetics. 2013;93(3):506-14.