# **Functional and Molecular Studies in Primary Carnitine Deficiency.**

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#### ABSTRACT

Primary carnitine deficiency is caused by a defect in the OCTN2 carnitine transporter encoded by the *SLC22A5* gene. It can cause hypoketotic hypoglycemia or cardiomyopathy in children, and sudden death in children and adults. Fibroblasts from affected patients have reduced carnitine transport. We evaluated carnitine transport in fibroblasts from 358 subjects referred for possible carnitine deficiency. Carnitine transport was reduced to 20% or less of normal in fibroblasts of 140/358 subjects. Sequencing of the 10 exons and flanking regions of the *SLC22A5* gene in 95/140 subjects identified causative variants in 84% of the alleles. The

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missense variants identified in our patients and others previously reported (n=92) were expressed in CHO cells. Carnitine transport was impaired by 73/92 variants expressed. Prediction algorithms (Polyphen-2, SIFT) correctly predicted the functional effects of expressed variants in about 80% of cases. These results indicate that mutations in the coding region of the *SLC22A5* gene cannot be identified in about 16% of the alleles causing primary carnitine deficiency. Prediction algorithms failed to determine the functional effects of amino acid substitutions in this transmembrane proteins in about 20% of cases. Therefore, functional studies in fibroblasts remain the best strategy to confirm or exclude a diagnosis of primary carnitine deficiency.

**KEY WORDS:** carnitine deficiency; carnitine uptake defect; fatty acid oxidation; carnitine transport; *SLC22A5*; OCTN2; mutations; newborn screening.

# **INTRODUCTION**

Primary carnitine deficiency (MIM# 212140) is an autosomal recessive disorder of the carnitine cycle resulting in defective fatty acid oxidation (Longo, et al., 2016). The incidence of primary carnitine deficiency varies with a frequency of approximately 1:40,000 newborns in Japan, 1:37,000–1:100,000 newborns in Australia, and 1:142,000 in the USA (Koizumi, et al., 1999; Longo, et al., 2016; Therrell, et al., 2014; Wilcken, et al., 2003). The highest incidence of primary carnitine deficiency (1:300) is in the Faroe Islands, an archipelago in the North Atlantic that remained geographically isolated for many centuries, where about 5% of the population is carrier for an abnormal allele (Longo, et al., 2016; Rasmussen, et al., 2014a; Rasmussen, et al., 2014b; Steuerwald, et al., 2017).

The gene for primary carnitine deficiency, *SLC22A5* (MIM# 603377), spans about 30 kb on chromosome 5q31 (chr5:132,369,752–132,395,614, hg38) and encodes for the organic cation transporter novel 2 (OCTN2) carnitine transporter (Longo, et al., 2016; Tamai, et al., 1998; Wu, et al., 1998). OCTN2 was identified in 1998 for its homology to the organic cation transport novel 1 (OCTN1) (Tamai, et al., 1998; Wu, et al., 1999; Wu, et al., 1998). OCTN1 and OCTN2 are part of the SLC22 family of membrane transporters, along with organic anion and cation transporters (Burckhardt and Wolff, 2000; Frigeni, et al., 2016; Longo, et al., 2016). OCTN2 operates as a Na<sup>+</sup>-independent organic cation transporter as well as a high affinity ( $K_m = 2.9 \pm 0.7 \mu M$ ) Na<sup>+</sup>-dependent carnitine transporter (Longo, et al., 2016; Tamai, et al., 1998; Wang, et al., 2000b; Wu, et al., 1999). Over 150 genetic variants are described in the *SLC22A5* mutation database

(http://www.arup.utah.edu/database/OCTN2/OCTN2\_display.php), of which 86 are pathogenic (including missense, nonsense, indels, frame-shifting, splice-site altering mutations and one large gene deletion).

Defective carnitine transport results in urinary carnitine wasting, low serum carnitine levels (< 9  $\mu$ M, normal 25-50  $\mu$ M), and decreased intracellular carnitine accumulation (Longo, et al., 2006; Longo, et al., 2016). Patients with primary carnitine deficiency lose most (10–95%) of the filtered carnitine in urine, and their heterozygous parents lose 2 to 3 times the normal amount, explaining their mildly reduced plasma carnitine levels (Longo, et al., 2016). Since carnitine is required for the transfer of long-chain fatty acids from the cytoplasm to the mitochondrial matrix for subsequent  $\beta$ -oxidation, the lack of carnitine impairs the ability to use fat as energy source during periods of metabolic stress (Longo, et al., 2016). If carnitine supplements are not promptly started, patients with primary carnitine deficiency can present with an acute metabolic decompensation early in life, or later in life with skeletal and cardiac myopathy or sudden death from arrhythmia (Longo, et al., 2016; Rose, et al., 2012; Wang, et al., 2000a). Siblings of affected children had only mild developmental delays or were asymptomatic (Longo, et al., 2016; Spiekerkoetter, et al., 2003; Wang, et al., 2001).

With the advent of expanded newborn screening, infants with primary carnitine deficiency or their affected mothers can be identified by very low levels of free carnitine (Longo, et al., 2016). The diagnosis of primary carnitine deficiency can be biochemically confirmed by demonstration of low free carnitine levels in plasma with reduced renal reabsorption (less than 90%), and normal renal function with no abnormalities in the urine organic acids (although a non-specific dicarboxylic aciduria has been reported during acute attacks) (Longo, et al., 2016; Scaglia, et al., 1998). Given the possibility of a maternal disorder causing primary or secondary carnitine deficiency, plasma and urine carnitine, plasma acylcarnitine profile, and urine organic acids should be evaluated in the mother as well when carnitine levels are low in the newborn screening (Longo, et al., 2016). The diagnosis can be definitively confirmed by verifying reduced carnitine transport activity (< 20% of normal controls) in skin fibroblasts from the patient or by mutational analysis of the *SLC22A5* gene (Longo, et al., 2016).

The measurement of carnitine transport in cultured fibroblasts is a very reliable method to confirm the diagnosis of primary carnitine deficiency, but is time demanding and requires a skin biopsy (Longo, et al., 2016). For this reason, molecular analysis (sequencing and deletion/duplication analysis) of the 10 exons of the *SLC22A5* gene and exon/intron boundaries in DNA obtained from blood is now the first-line test for diagnostic confirmation (Longo, et al., 2016).

Accepted Article

Here the reliability of DNA testing is compared to functional studies in fibroblasts to confirm the diagnosis of primary carnitine deficiency in 95 subjects. In addition, several novel missense mutations are expressed in heterologous cells to confirm their pathogenicity.

## **MATERIALS AND METHODS**

Patients and Membrane Transport. All studies were approved by the IRB of the University of Utah. Patients were referred for diagnostic confirmation either after symptomatic presentation or after identification by newborn screening programs. Some of the patients reported in this study were previously individually described (Amat di San Filippo and Longo, 2004; Amat di San Filippo, et al., 2006; Amat di San Filippo, et al., 2008; Amat di San Filippo, et al., 2003; De Biase, et al., 2012; Filippo, et al., 2011; Rose, et al., 2012; Schimmenti, et al., 2007; Wang, et al., 2000a; Wang, et al., 2001; Wang, et al., 2000b; Wang, et al., 2000c; Wang, et al., 1999). Fibroblasts from patients with primary carnitine deficiency were obtained by skin biopsy for diagnostic purposes. They were grown in Dulbecco's Modified Eagle Medium (Thermo Fisher Scientific Inc., Waltham, MA) supplemented with 12% fetal bovine serum (Sigma-Aldrich Corp., St. Louis, MO), 2 mM L-Glutamine (Thermo Fisher Scientific Inc., Waltham, MA), 100 U/ml Penicillin-Streptomycin (GE Healthcare Life Sciences HyClone Laboratories, South Logan, UT), 2.5 µg/mL Amphotericin B solution (GE Healthcare Life Sciences HyClone Laboratories, South Logan, UT). <sup>3</sup>H-Carnitine (0.5 µM) transport was measured for 4 hours at 37 °C in adherent cells using the cluster-tray method (Scaglia, et al., 1999; Wang, et al., 2000b). Non-saturable transport was measured in the presence of 2 mM cold carnitine and was subtracted from total transport to obtain saturable transport (Scaglia, et al., 1999; Wang, et al., 2000b). Values are reported as means  $\pm$  SE of 6 independent determinations obtained in two separate experiments.

**DNA Analysis and Molecular Techniques.** Genomic DNA was extracted from fibroblasts or peripheral blood by standard methods. GenBank sequence AB016625.1 was used as reference for the gene, NM\_003060.2 was used as the reference sequence for the cDNA, and NP\_001295051.1 for the protein. Nucleotide numbering uses the A of the ATG translation initiation start site as nucleotide +1. The 10 exons of the *SLC22A5* gene including exon/intron boundaries were amplified by PCR using a total of 10 primer pairs tailed with universal M-13 forward and reverse sequences. Amplification was verified by automated gel electrophoresis in the HDA-GT12<sup>TM</sup> Genetic Analyzer. PCR fragments were purified with ExoSap-IT (USB) and bi-directionally sequenced using BigDye-Terminator v.1.1 cycle sequencing kit and capillary electrophoresis in the AB13730. Generated sequences were aligned to reference sequence (GenBank AB016625.1) using Mutation Surveyor v.4.0 (Softgenetics) to detect mutations (Wang, et al., 2001). Multiplex ligation-dependent probe amplification (MLPA) analysis was performed to detect large deletions within the *SLC22A5* 

gene. Each DNA sample (400 ng) was analyzed using the SALSA MLPA P076 ACADVL-SLC22A5 Probemix for SLC22A5 (MRC-Holland, Amsterdam, The Netherlands) following the manufacturer's recommendations (http://mlpa.com/WebForms/WebFormMain.aspx).

Missense variants identified in patients with primary carnitine deficiency were recreated in the OCTN2-EGFP expression vector by site-directed mutagenesis using the Quick Change system (Agilent Technologies Inc., Santa Clara, CA) following the manufacturer's instructions (Amat di San Filippo, et al., 2003; Wang, et al., 2000b). The final plasmids were sequenced to confirm the presence of the mutation and the absence of PCR artifacts and stably transfected into Chinese Hamster Ovary (CHO) cells. The plasmid contained the neomycin-resistance gene and cells were selected for resistance to G418 (Geneticin, 0.8 mg/ml) for 3 weeks before testing. A mass culture including different clones was then used for functional studies. Confocal microscopy was used to confirm protein production, since in our expression vector the OCTN2 transporter is tagged with the green fluorescent protein (Wang, et al., 2000b). CHO cells were grown in Ham's F12 Nutrient Mixture (Thermo Fisher Scientific Inc., Waltham, MA) supplemented with 6% fetal bovine serum (Sigma-Aldrich Corp., St. Louis, MO), 2 mM L-Glutamine (Thermo Fisher Scientific Inc., Waltham, MA), 100 U/ml Penicillin-Streptomycin (GE Healthcare Life Sciences HyClone Laboratories, South Logan, UT), 2.5 µg/ml Amphotericin B solution (GE Healthcare Life Sciences HyClone Laboratories, South Logan, UT). Carnitine (0.5 µM) transport was measured at 37°C with the cluster-tray method as previously described (Amat di San Filippo, et al., 2003; Wang, et al., 2000b).

*In silico* analysis of missense mutations. The missense variants identified and expressed in CHO cells were evaluated using Sorting Intolerant From Tolerant (SIFT, <u>http://sift.jcvi.org/www/SIFT\_enst\_submit.html</u>) and Polymorphism Phenotyping Version v2.2.2r398 (POLYPHEN-2, http://genetics.bwh.harvard.edu/pph2/index.shtml) software programs for variant evaluation.

**Estimate of allele frequency.** The Exome Aggregation Consortium Browser Beta (ExAC, http://exac.broadinstitute.org/) and the Genome Aggregation Database (gnomAD, http://gnomad.broadinstitute.org/) were used to determine the allele frequency of the variants identified in the *SLC22A5* gene (Lek, et al., 2016).

# RESULTS

**Functional and molecular characterization of fibroblasts from patients with primary carnitine deficiency.** Carnitine transport was measured in skin fibroblasts from 358 subjects referred for possible primary carnitine deficiency (Figure 1). A wide range of carnitine transport activity was observed in these cells with a nearly continuous distribution of activity from 0% of control to values above that measured in matched controls. A

reduction of carnitine transport to 20% or less than paired normal control fibroblasts was used as cut-off for the diagnosis of primary carnitine deficiency, since all of our patients whose fibroblasts had carnitine transport below 20% of normal controls were not able to maintain free carnitine level  $\geq$ 9 µmol/L (normal 25-50 µM) without supplements (Longo, et al., 2016). Cells from 140/358 subjects (39%) had carnitine transport activity below 20% of paired normal controls.

Molecular studies (sequencing of the 10 exons and flanking regions of the SLC22A5 gene encoding the OCTN2 carnitine transporter and deletion/duplication analysis) were performed in 95/140 of the affected subjects. Table 1 shows carnitine transport activity in fibroblasts of the 95 patients diagnosed with primary carnitine deficiency along with the variants identified in their SLC22A5 gene. Among the 72 variants identified, 48 were missense, 9 caused a frameshift, 8 were nonsense, 2 were in-frame deletions, 1 was a synonymous variant, and 4 were localized in flanking regions of the SLC22A5 gene. Overall, the most frequent mutation was c.136C>T (p.Pro46Ser), which was identified in 13/190 alleles. This mutation has been identified in asymptomatic or minimally symptomatic (easy fatigability, muscle pain with exercise, fasting intolerance) adult women (Longo, et al., 2016; Schimmenti, et al., 2007) and in a mother (compound heterozygous p.Pro46Ser/p.Asn32Ser) who suffered repeated episodes of cardiac arrest requiring placement of a defibrillator, episodes that resolved after her diagnosis and initiation of carnitine therapy (De Biase, et al., 2012). This mutation (p.Pro46Ser) reduces, but does not abolish carnitine transport and affects glycosylation and maturation to the plasma membrane of the OCTN2 carnitine transporter (Filippo, et al., 2011; Longo, et al., 2016). This and other similar missense mutations seem protective against early clinical manifestations of primary carnitine deficiency, but can still be associated with fatal cardiac arrhythmia leading to sudden death (De Biase, et al., 2012; Longo, et al., 2016; Rose, et al., 2012). Molecular analysis identified causative variants in 84% of the affected alleles, but failed to identify significant variations in 6/95 patients, and a second mutation in 19/95 patients (Table 2), despite a functional diagnosis of primary carnitine deficiency.

**Expression of** *SLC22A5* variants in CHO cells. Molecular analysis is becoming the preferred method for diagnostic confirmation in patients with suspected primary carnitine deficiency since it does not require a skin biopsy and is relatively rapid. Among samples received for molecular analysis (including the 95 patients who had carnitine transport analyzed in fibroblasts), 133 different variants were identified, with 33 being novel (Table 3A). Of the 133 variants identified, 90 were missense, 16 caused a frameshift, 14 were nonsense, 8 affected splicing, 2 were in-frame deletions of single amino acids, 1 was a ~1.6 Mb deletion encompassing the entire OCTN2 gene (Li, et al., 2010), 1 was a 113 bp deletion encompassing the initiation codon of exon 1 (Nezu, et al., 1999), and 1 was a synonymous variant that created a splice site (c.453 G>A, p.Val151Val). An additional missense variant

(c.393G>C, p.Glu131Asp) in the last nucleotide abolished the donor site of exon 1 and is predicted to affect splicing. Twenty-six new missense variants and 1 new in-frame deletion detected in our patients, as well as 64 missense variants and 1 in-frame deletion reported in the literature (most never previously expressed in heterologous cells, see Figure 2 for their location on the OCTN2 transporter), were expressed in CHO cells to determine their effect on carnitine transport. Carnitine transport was expressed as percent of the value measured in CHO cells expressing the wild-type OCTN2 carnitine transporter, assumed to be 100% (Table 3A). Carnitine transport was reduced to less than 20% of control in 73/92 variants, indicating that these amino acid substitutions significantly impaired carnitine transport activity.

**Frequency of tested variants in the general population.** Of the 92 variants tested, 19 retained significant residual carnitine transport activity (>20% of normal), and several of them (p.Gly12Ser; p.Leu20His; p.Ala142Ser; p.Val151Met; p.Ala214Val; p.Arg254Gln; p.Thr264Met; p.Ile312Val; p.Glu317Lys; p.Ile348Thr; p.Arg488H; p.Pro549Ser) had frequency in the ExAC Browser Beta and in the gnomAD much higher than those impairing carnitine transport activity, with some individuals being homozygous for some of these milder variants (Table 3B). These variants affected different domains of the OCTN2 carnitine transporter (open circles in Figure 2), including transmembrane and hydrophilic regions in a manner similar to pathogenic missense variants (filled circles in Figure 2). While we cannot exclude that some of these changes might represent mutations (they could affect splicing which was not directly tested here, or they could be in combination with other variants impairing carnitine transport such when p.Ala142Ser is in *cis* with p.Arg488His (Amat di San Filippo, et al., 2006)), we can conclude that they do not affect the function of the protein if synthesized.

Functional studies allowed us to calculate the minimal estimated frequency of people carrying two abnormal alleles from the frequency in normal individuals reported in the ExAC Browser Beta and in the gnomAD (Table 3B). From these data, 1:221 (0.0045289) alleles contained either one of the missense mutations expressed in this study or a mutation causing the premature insertion of a stop codon or affecting splicing in the *SLC22A5* gene (Tables 3B and Supplemental Table 1). Note that one normal subject reported in the gnomAD browser was homozygous for the p.Tyr449Asp variant (Table 3B). This variant reduced carnitine transport to about 17% of normal OCTN2 and has only been identified in one heterozygous patient whose fibroblasts had about 50% of normal carnitine transport (carrier of primary carnitine deficiency) (Amat di San Filippo and Longo, 2004). We cannot exclude that the subject homozygous for this variant has a late-onset form of primary carnitine deficiency.

Given the frequency of pathogenic variants in the normal population (p), the predicted frequency of individuals with two abnormal alleles  $(p^2)$  in a general population would be  $1.6817*10^{-5}$  or 1:59,465.

**Variant analysis software.** Polyphen-2 and SIFT are software programs commonly used to predict the pathogenicity of missense variants. The 90 missense variants and 2 inframe deletions whose carnitine transport was tested in CHO cells were evaluated using these software programs and the results compared (Table 3C). Both programs were effective in separating benign from damaging variants (p=0.0005 for Polyphen-2 and p=0.0009 for SIFT using analysis of variance, Table 4). However, about 20% of the variants were incorrectly assigned by either program, with a more marked error in the prediction of benign variants (Table 4). When only variants in which the two programs provided a concordant interpretation (74/92) were analyzed, the correct prediction rate increased to 86% (Table 4).

#### DISCUSSION

Primary carnitine deficiency is a recessive condition caused by mutations in the OCTN2 carnitine transporter encoded by the *SLC22A5* gene (Longo, et al., 2016). The defective OCTN2 carnitine transporter is expressed in fibroblasts and these cells can be used for diagnostic confirmation (Longo, et al., 2016). By looking at a large number of diagnostic samples on which functional studies were performed, there was a continuum of distribution of carnitine transport activity, from 0% in some patients up to 1.5 times normal in certain subjects (Figure 1). While some of this variability might be in part related to variability of the assay, the measurements reported in individual cells were reproducible in repeated experiments, indicating the likely contribution of genetic factors. It is interesting that 218 of 358 fibroblast strains had no evidence of a carnitine uptake defect and that carnitine transport activity was above 60% of normal (the highest levels we have seen in a carrier) in about 70 patients. This suggests that in many cases confirmatory testing might have been requested immediately at the time of initial presentation, without following over time plasma carnitine levels to remove cases of secondary carnitine deficiency (in most cases nutritional).

Genetic analysis identified a number of missense variants, some of which decreased substantially transport activity, others that caused only a modest or no impairment (Table 3). The variability reported here for a limited number of subjects is only a portion of that observed in normal individuals, with about 354 missense variants reported in the ExAC Browser Beta (http://exac.broadinstitute.org/gene/ENSG00000197375) and in the gnomAD browser (http:// http://gnomad.broadinstitute.org/gene/ENSG00000197375). It is conceivable that missense variations, in addition to variations in regulatory sequences of the gene, are responsible for this continuous, rather that discontinuous distribution of carnitine transport activity in fibroblasts.

In our experience, patients whose fibroblasts had less than 20% carnitine transport activity were unable to maintain free carnitine level  $\geq 9 \ \mu mol/L$  and required carnitine

supplementation. For this reason, we established 20% of normal transport as the cut-off below which we diagnose primary carnitine deficiency and provide carnitine supplementation. Fibroblasts from 95 patients had carnitine transport activity <20% (Table 1). Molecular analysis identified causative mutations in 159/190 alleles, while no variants were identified in 31/190 alleles (Table 2), indicating that some mutations lie outside the coding region or flanking sequence of the *SLC22A5* gene. It must be noted that we still know very little of the promoter region of this gene and other regulatory sequences that are not routinely sequenced. For this reason, the diagnosis of primary carnitine deficiency can be missed in some patients if only DNA studies are performed. For example, 6 out of 95 of our patients would have been missed completely by molecular analysis (no significant sequence variations identified in either allele), if functional studies in fibroblasts had not been performed. This has profound implications since these patients could have escaped diagnosis and could have been left untreated without functional studies.

It is crucial to correctly identify all the patients affected by primary carnitine deficiency and promptly start carnitine supplements. If left untreated, primary carnitine deficiency can lead to hypoketotic hypoglycemia early in life, or manifest with skeletal and cardiac myopathy, and in some cases sudden death at any age (Longo, et al., 2016). At the same time, a subject wrongly diagnosed with the disease will have to face an unnecessary lifelong therapy with carnitine supplements (Longo, et al., 2016).

Unlike mutations causing the premature insertion of a stop codon or affecting splicing, the interpretation of missense variants found by molecular analysis can be difficult and requires additional studies. Here we expressed 90 missense variants and 2 in-frame deletions identified in our patients in CHO cells to determine their carnitine transport activity (Table 3A). A carnitine transport activity of 20% or less than CHO expressing the wild-type OCTN2 carnitine transporter (transport activity of 100%) was considered pathogenic. Of the variants tested, 73/92 were pathogenic, and 19/92 were benign (Table 3A).

Polyphen-2 and SIFT are software programs commonly used to predict the effect of missense variants on protein function. We compared the predictions of Polyphen-2 and SIFT against functional studies of carnitine transport in CHO cells (Table 3C). Our results indicate that while Polyphen-2 and SIFT were effective in separating benign from damaging variants, they incorrectly identified the effect of variants in about 20% of the cases (Table 4). For variants in which Polyphen-2 and SIFT provided a concordant interpretation, the effect of benign variants was wrongly predicted in 53% of the cases, while the error was reduced to 3% when predicting the effect of damaging variants (Table 4). For these reasons, these programs seem skewed in predicting more damaging effects of missense mutations than experimentally observed (false positive results) in the OCTN2 carnitine transporter. We do not know whether this could extend to the interpretation of functional effects in other membrane transporters.

The incidence of primary carnitine deficiency is approximately 1:142,000 in the USA as ascertained by newborn screening (Therrell, et al., 2014). From the frequency of pathogenic alleles reported in the ExAC Browser Beta and in the gnomAD Browser in about 120,000 healthy (heterozygotes) individuals (Table 3B and supplemental table 1) (Lek, et al., 2016), the calculated frequency of mutant alleles is 1:221, with a minimal estimated frequency of people carrying two abnormal alleles in the general population of 1.6817\*10<sup>-5</sup> or 1:59,465 individuals. This frequency is significantly higher than the one reported by newborn screening, indicating that some affected individuals might be missed by the current neonatal screening protocols. We do not know whether this is due to specific mutations (Chen, et al., 2013) and/or to problems related to the timing of collection of the newborn screening sample, in which the blood samples are collected at 1-3 days of age and carnitine levels might reflect those of the mother (Pasquali and Longo, 2013; Rasmussen, et al., 2017; Schimmenti, et al., 2007). Carnitine is transferred from the mother to the child via the placenta, and immediately after birth levels of free carnitine (C0) are usually lower in infants of mothers with primary carnitine deficiency as compared to infants with the disease themselves (Pasquali and Longo, 2013). With time, levels decrease in infants with primary carnitine deficiency, but remain stable or slightly increase in infants of mothers with primary carnitine deficiency (Pasquali and Longo, 2013). The diagnosis can therefore be missed if screening is performed too close to birth and no second screening is obtained. In the Faroe Islands, post-neonatal screening (age >2 months) was able to identify additional patients who had unrevealing newborn screening, confirming that patients with primary carnitine deficiency are missed by current newborn screening methods (Steuerwald, et al., 2017). For these reasons, primary carnitine deficiency should be considered in children and adults with a suitable clinical presentation (hypoglycemia, cardiomyopathy, arrhythmia, sudden death) even in the presence of a normal newborn screening test.

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The Authors have no conflict of interest to declare for this manuscript.

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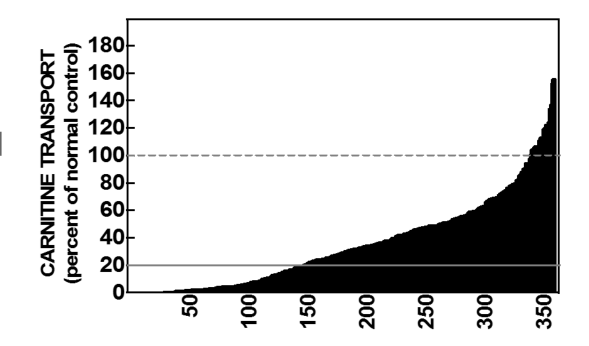
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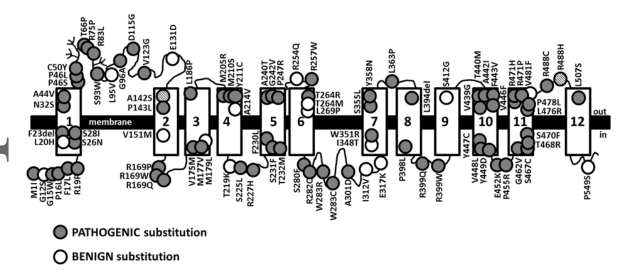
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#### **LEGENDS TO FIGURES.**

Figure 1. Carnitine transport by fibroblasts obtained from 358 individuals referred for possible primary carnitine deficiency. <sup>3</sup>H-Carnitine (0.5  $\mu$ M) transport was measured for 4 hours at 37°C. Nonsaturable transport, measured in the presence of 2 mM cold carnitine, was subtracted from total transport to obtain saturable carnitine transport. Transport activity was normalized to the one of simultaneous normal controls (100% activity). Each bar represents the average of 6 observations in two separate experiments with the sample numbered from 1 (lowest activity) to 358 (highest activity) indicated on the X-axis. Carnitine transport activity of 20% or less of normal controls was indicative of primary carnitine deficiency (continuous line). Fibroblasts from some of the subjects had carnitine transport higher than their paired normal controls (100% activity, dashed line).



**Figure 2. Schematic of the OCTN2 carnitine transporter with location of the 92 variants expressed in CHO cells.** The OCTN2 carnitine transporter is composed of 557 amino acids. Hydropathy analysis and conservation with other organic cation transporters indicates that the transporter forms 12 transmembrane spanning domains (rectangles) with both the N- and C-termini facing the cytoplasm. Glycosylation sites are indicated by branching (Filippo, et al., 2011). The 92 OCTN2 variants identified and expressed in CHO cells are indicated by circles. Mutations reducing carnitine transport to 20% or less than wild-type when expressed in CHO cells are indicated by full circles; variants retaining more that 20% residual activity are indicated by empty circles. The A142S and R488H mutations (indicated by black stripes circles) impair carnitine transport when in *cis* on the same allele, but not alone (Amat di San Filippo, et al., 2006).



#### **TABLES**

Table 1. Functional and molecular analysis of cells from 95 patients with primary carnitine deficiency. Carnitine transport activity was measured in fibroblasts from subjects referred for possible primary carnitine deficiency (Figure 1) and expressed as percent of normal activity. The 10 exons and flanking regions of the *SLC22A5* gene encoding the OCTN2 carnitine transporter were sequenced and subjected to deletion/duplication analysis in patients whose fibroblasts had <20% of normal transport activity (transport activity is reported in the second column). Variations from the standard sequence (NM\_003060.2) are reported in columns 3-10. No causative variants were identified in 16% of the alleles (*NOT FOUND*).

	Pt#	Transport activity % of control	ALLELE 1 cDNA	PROTEIN	ALLELE 2 cDNA	PROTEIN	ADDITIONAL VA	ARIANT	
	1	0.0	c.12C>G	p.Tyr4*	c.12C>G	p.Tyr4*			
	2	0.1	c.865C>T	p.Arg289*	c.865C>T	p.Arg289*			
	3	0.1	c.458_459del	p.Val153Alafs*41	c.844C>T	p.Arg282*			
	4	0.2	c.1202_1203insA	p.Tyr401*	c.1302del	p.Gly435fs*24			
	5	0.2	c.844C>T	p.Arg282*	c.844C>T	p.Arg282*			
	6	0.3	c.740C>G	p.Pro247Arg	c.740C>G	p.Pro247Arg			
	7	0.3	c.64_66del	p.Phe22del	c.64_66del	p.Phe22del			
	8	0.3	c.254_264dup	p.lle89Glyfs*45	c.254_264dup	p.lle89Glyfs*45			
	9	0.3	c.1319C>T	p.Thr440Met	c.1319C>T	p.Thr440Met			
	10	0.4	c.248G>T	p.Arg83Leu	c.248G>T	p.Arg83Leu			
J	11	0.5	c.760C>T	p.Arg254*	c.760C>T	p.Arg254*			
٦	12	0.5	c.844C>T	p.Arg282*	c.3G>T	p.Met1Leu			
	13	0.5	c.12C>G	p.Tyr4*	c.12C>G	p.Tyr4*			
	14	0.5	c.839C>T	p.Ser280Phe	c.1463G>A	p.Arg488His	c.424G>T	p.Ala142Ser	
	15	0.6	c.506G>A	p.Arg169GIn	c.506G>A	p.Arg169Gln			
	16	0.7	c.1392_1409delinsCA	p.Val465Thrfs*29	c.1392_1409delinsCA	p.Val465Thrfs*29			
	17	0.9	c.458_459del	p.Val153Alafs*41	c.865C>T	p.Arg289*			
	18	0.9	c.760C>T	p.Arg254*	c.760C>T	p.Arg254*			
	19	0.9	c.760C>T	p.Arg254*	c.760C>T	p.Arg254*			
	20	1.0	c.565_568del	p.Phe189Argfs*14	c.1316T>G	p.Val439Gly			
	21	1.0	c.1340A>G	p.Tyr447C	c.1340A>G	p.Tyr447Cys			
	22	1.0	c.768G>A	p.Trp256*	c.1403C>G	p.Thr468Arg			
	23	1.0	c.825G>A	p.Trp275*	c.1267+3_1267+23del				
	24	1.1	c.505C>T	p.Arg169Trp	c.1051T>C	p.Trp351Arg			
1					I		I		; <b>I</b>

	25	1.1	c.1193C>T	p.Pro398Leu	c.839C>T	p.Ser280Phe				
	26	1.2	c.1319C>T	p.Thr440Met	c.1463G>A	p.Arg488H	c.424G>T	p.Ala142Ser		
	27	1.3	c.344A>G	p.Asp115Gly	c.344A>G	p.Asp115Gly	c.283C>G	p.Leu95Val	c.283C>G	p.Leu95Val
	28	1.4	c.844C>T	p.Arg282*	c.1463G>A	p.Arg488His	c.424G>T	p.Ala142Ser		
	29	1.5	c.1193C>T	p.Pro398Leu	c.557T>C	p.Leu186Pro	c.1645C>T	p.Pro549Ser		
	30	1.5	c.506G>C	p.Arg169Pro	c.1088T>C	p.Leu363Pro				
	31	2.0	c.902C>A	p.Ala301Asp	c.902C>A	p.Ala301Asp				
	32	2.0	c.760C>T	p.Arg254*	c.1400C>G	p.Ser467Cys				
	33	2.1	c.248G>T	p.Arg83Leu	c.248G>T	p.Arg83Leu				
	34	2.2	c.629A>G	p.Asn210Ser	c.1463G>A	p.Arg488His	c.424G>T	p.Ala142Ser		
	35	2.2	c.56G>C	p.Arg19Pro	c.254_264dup	p.lle89Glyfs*45				
	36	2.5	c.136C>T	p.Pro46Ser	c.760C>T	p.Arg254*				
	37	2.8	c.1196G>A	p.Arg399GIn	c.1463G>A	p.Arg488His	c.424G>T	p.Ala142Ser		
	38	2.8	c.725G>T	p.Gly242Val	c.725G>T	p.Gly242Val				
	39	2.8	c.844C>T	p.Arg282*	c.136C>G	p.Pro46Ser				
	40	2.9	c.847T>A	p.Trp283Arg	c. 847T>A	p.Trp283Arg				
λ	41	2.9	c.43G>T	p.Gly15Trp	c.248G>T	p.Arg83Leu				
	42	3.0	c.760C>T	p.Arg254*	c.760C>T	p.Arg254*				
	43	3.2	c.505C>T	p.Arg169Trp	c.505C>T	p.Arg169Trp				
	44	3.3	c.1196G>A	p.Arg399Gin	c.1196G>A	p.Arg399GIn				
	45	3.3	c.695C>T	p.Thr232Met	c.1403C>G	p.Thr468Arg				
	46	3.4	c.136C>T	p.Pro46Ser	c.1556_1559dup	p.lle521Hisfs*3				
	47	3.4	c.232delC	p.His79Thrfs*51	c.232delC	p.His79Thrfs*51				
	48	3.5	c.248G>T	p.Arg83Leu	c.641C>T	p.Ala214Val				
	49	3.8	c.254_264dup	p.lle89Glyfs*45	c.1385G>T	p.Gly462Val				
	50	3.8	c.845G>A	p.Arg282GIn	c.845G>A	p.Arg282GIn				
	51	4.0	c.695C>T	p.Thr232Met	NOT FOUND					
	52	4.0	c.136C>G	p.Pro46Ser	c.1463G>A	p.Arg488His	c.424G>T	p.Ala142Ser		
	53	4.2	c.136C>G	p.Pro46Ser	c.1556_1559dup	p.lle521Hisfs*3				
	54	4.2	c.1354G>A	p.Glu452Lys	c.1354G>A	p.Glu452Lys				
	55	4.3	c.77G>A	p.Ser26Asn	c.845G>A	p.Arg282GIn				
	56	4.3	c.232delC	p.His79Thrfs*51	c.1342G>T	p.Val448Leu	c.641C>T	p.Ala214Val		
Acc	57	4.8	c.95A>G	p.Asn32Ser	c.136C>G	p.Pro46Ser				
	58	4.9	c.1324_1325GC>AT	p.Ala442lle	c.1324_1325GC>AT	p.Ala442lle				
	59	5.0	c.1462C>T	p.Arg488Cys	c.1462C>T	p.Arg488Cys				
	60	5.3	Exon 3 deletion	p.Phe167Aspfs*61	NOT FOUND		c.394-16T>A			
	61	5.5	c.695C>T	p.Thr232Met	c.695C>T	p.Thr232Met				
									1	

	62	5.6	c.136C>T	p.Pro46Ser	c.695C>T	p.Thr232Met			
	63	5.6	c.136C>T	p.Pro46Ser	c.1463G>A	p.Arg488His	c.424G>T	p.Ala142Ser	
	64	5.7	c.844C>T	p.Arg282*	NOT FOUND		c.824+13T>C		
	65	5.7	c.136C>T	p.Pro46Ser	c.1193C>T	p.Pro398Leu			
	66	5.8	c.1319C>T	p.Thr440Met	c.453G>A	p.Val151Val			
	67	5.9	c.1319C>T	p.Thr440Met	NOT FOUND				
$\overline{\mathbf{O}}$	68	6.3	c.136C>T	p.Pro46Ser	c.523G>A	p.Val175Met			
	69	7.0	c.458_459del	p.Val153Alafs*41	NOT FOUND				
	70	7.2	c.137C>T	p.Pro46Leu	c.1175_1177del	p.Leu394del			
	71	7.4	c.1195C>T	p.Arg399Trp	c.653-2A>C		c.653-2A>C		
	72	7.5	c.865C>T	p.Arg289*	NOT FOUND				
Itti	73	8.5	c.248G>T	p.Arg83Leu	c.641C>T	p.Ala214Val			
	74	8.9	c.43G>T	p.Gly15Trp	NOT FOUND				
	75	9.0	c.453G>A	p.Val151Val	Exon 3 deletion	p.Phe167Aspfs*61			
	76	9.1	c.136C>T	p.Pro46Ser	NOT FOUND				
	77	9.6	NOT FOUND		NOT FOUND		c.1441G>T	p.Val481Phe	
	78	11.1	c.136C>T	p.Pro46Ser	NOT FOUND				
	79	11.1	NOT FOUND		NOT FOUND		c.1451G>T	p.Gly484Val	
	80	11.7	NOT FOUND		NOT FOUND				
ote	81	12.4	c.806delT	p.Leu269Argfs*26	c.1345T>G	p.Tyr449Asp			
	82	13.7	c.839C>T	p.Ser280Phe	NOT FOUND				
	83	13.9	NOT FOUND		NOT FOUND				
	84	14.0	c.505C>T	p.Arg169Trp	NOT FOUND				
	85	14.0	c.419G>A	p.Trp140*	NOT FOUND				
	86	14.3	c.51C>G	p.Phe17Leu	NOT FOUND				
	87	15.0	NOT FOUND		NOT FOUND				
	88	16.2	c.641C>T	p.Ala214Val	c.641C>T	p.Ala214Val			
	89	16.8	c.393G>C	p.Glu131Asp	NOT FOUND				
	90	16.9	NOT FOUND		NOT FOUND				
	91	17.7	c.1202_1203insA	p.Tyr401*	NOT FOUND				
	92	17.8	c.136C>T	p.Pro46Ser	NOT FOUND				
Acc	93	17.9	c.1193C>T	p.Pro398Leu	NOT FOUND				
	94	18.7	c.458_459del	p.Val153Alafs*41	NOT FOUND				
	95	19.4	c.1193C>T	p.Pro398Leu	NOT FOUND				

**Table 2. Molecular studies in patients with primary carnitine deficiency.** Sequencing and deletion/duplication analysis of all 10 exons of the *SLC22A5* gene and flanking regions was performed in 95 patients diagnosed with primary carnitine deficiency because of a low level of carnitine transport in fibroblasts (activity <20% of matching normal controls). No variants were identified in 31/190 (16%) of the alleles sequenced. For 6/95 (6%) affected patients no variants were identified in either alleles, while for 19/95 (20%) affected patients no variant was identified in one allele.

95	NUMBER of PATIENTS
190	NUMBER of ALLELES
31/190 (16%)	Alleles with unknown mutations
19/95 (20%)	Patients with no identified variant in one allele
6/95 (6%)	Patients with no identified variant in either alleles

**Table 3. Variants in the** *SLC22A5* gene in patients referred for possible primary carnitine. (A) Molecular analysis of the *SLC22A5* gene in our patients identified 133 variants, with 33 being novel (\*). Of the 133 variants identified, 90 missense variants and 2 in-frame deletions were expressed in CHO cells to evaluate carnitine transport (functional activity). Carnitine transport was expressed as percent of the activity in CHO cells expressing the wild-type OCTN2 carnitine transporter (100% activity). Carnitine transport was reduced to less than 20% of wild-type OCTN2 by 73/92 variants. (B) Table 3B shows the frequency in normal individuals in the ExAC Browser Beta and in the gnomAD of the alleles found in our patients along with the number of normal individual homozygous for a specific allele. 1:221 alleles contained either one of the missense mutations expressed in this study or a mutation causing the premature insertion of a stop codon or affecting splicing in the *SLC22A5* gene. (C) The 90 missense variants and 2 in-frame deletions whose carnitine transport was tested in CHO cells were evaluated for pathogenicity using the PolyPhen-2 v2.2.2r398 (http://genetics.bwh.harvard.edu/pph2/index.shtml, using default parameters, defining damaging any variant with a score >0.15) and SIFT

(<u>http://sift.jcvi.org/www/SIFT\_enst\_submit.html</u>, using default parameters and assuming as damaging any change with a score of <0.05) software with the predicted effects indicated. Actual values for each amino acid substitution are reported in Supplemental Table S2. Variants were submitted to ClinVar

https://www.ncbi.nlm.nih.gov/clinvar/?term=SLC22A5%5Bgene%5D.

		Α	
Exon	cDNA	PROTEIN	Transpo rt activity % wild- type OCTN2
	OCTN2 deletion		
1	c91_22del		
1	c.3G>T	p.Met1Leu	0.16
1	c.4_5insC	p.Arg2Profs*136	
1	c.12C>G	p.Tyr4*	
1	c.34G>A	p.Gly12Ser	51.7
1	c.43G>T	p.Gly15Trp	11.05
1	c.47C>T	p.Pro16Leu	0.00
1	c.51C>G	p.Phe17Leu	10.74
1	c.56G>C	p.Arg19Pro	2.03
1	c.59T>A	p.Leu20H	46.77
1	c.64_66del	p.Phe23del	0.46
1	c.77G>A	p.Ser26Asn	5.51
1	c.83G>T	p.Ser28lle	1.35
1	c.95A>G	p.Asn32Ser	0.88
1	c.131C>T	p.Ala44Val	8.22
1	c.136C>T	p.Pro46Ser	2.47
1	c.137C>T	p.Pro46Leu	4.10
1	c.149G>A	p.Cys50Tyr	0.00
1	c.196A>C	p.Thr66Pro	1.80
1	c.224G>C	p.Arg75Pro	1.93
1	c.232delC	p.His79Thrfs*51	
1	c.248G>T	p.Arg83Leu	0.14
1	c.254_264dup	p.lle89Glyfs*45	
1	c.278C>G	p.Ser93Trp	0.00
1	c.283C>G	p.Leu95Val	28.92

1

1

c.287G>C

c.344A>G

p.Gly96Ala

p.Asp115Gly

l	В	C	;
Allele Frequency (ExAC)	Homozygous (ExAC)	Polyphen-2	SIFT
3.23E-05	0	damaging	damaging
0.0000173	0		
0.0007152	1	damaging	damaging
0.0001028	0	damaging	damaging
	-		
		benign	benign
0.0000601	0	damaging	damaging
		damaging	benign
0.0013610	1	damaging	benign
0.0000259	0	/	damaging
0.0000262	0	damaging	benign
		damaging	benign
0.0000268	0	damaging	benign
0.0000197	0	damaging	damaging
0.0005515	0	damaging	damaging
0.0000202	0	damaging	damaging
		damaging	damaging
		damaging	damaging
0.0000177	0	benign	benign
0.0000181	0		
0.0004323	0	damaging	damaging
0.0000385	0		
		damaging	damaging
		benign	benign
0.0003927	0	damaging	damaging
		damaging	damaging

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(Li, et al., 2010)

19.65

4.04

Reference(s)

(Li, et al., 2010) (Nezu, et al., 1999)

(Dobrowolski, et al., 2005)

(Nezu, et al., 1999) (Wang, et al., 2001) (Li, et al., 2010)

(El-Hattab, et al., 2010)

(Wang, et al., 2001)

(Rose, et al., 2012)

(Rahbeeni, et al., 2002)

(Lamhonwah, et al., 2002; Rasmussen, et al., 2014b)

(Schimmenti, et al., 2007)

(Li, et al., 2010)

(Li, et al., 2010)

(Amat di San Filippo, et al., 2006)

(Makhseed, et al., 2004)

(Lamhonwah, et al., 2002; Wang, et al., 2001)

(Lamhonwah, et al., 2002)

(Lee, et al., 2010; Urban, et al., 2006)

	1	c.350G>A	p.Trp117*	
	1	c.368T>G	p.Val123Gly	13.96
	1	c.393G>C	p.Glu131Asp	32.84
	IVS1	c.393+5G>A		
	IVS1	c.394-16T>A		
	2	c.396G>A	p.Trp132*	
$\mathbf{O}$	2	c.419G>A	p.Trp140*	
•	2	c.424G>T	p.Ala142Ser	25.68
	2	c.428C>T	p.Pro143Leu	1.18
	2	c.433dupA	p. Thr145Asnfs*50	
	2	c.451G>A	p.Val151Met	65.93
		c.453G>A	p.Val151Val	
	2	c.458_459del	p.Val153fs*41	
	IVS2	c.497+1G>T		
		Exon 3 deletion	p.Phe167Aspfs*61	
	3	c.505C>T	p.Arg169Trp	0.01
	3	c.506G>C	p.Arg169Pro	0.24
	3	c.506G>A	p.Arg169GIn	0.22
	3	c.517delC	p.Leu173Cysfs*3	
	3	c.523G>A	p.Val175Met	7.03
	3	c.529A>G	p.Met177Val	14.40
	3	c.535A>T	p.Met179Leu	48.13
	3	c.557T>C	p.Leu186Pro	0.00
	3	c.565_568del	p.Phe189Argfs*13	
O	3	c.573delG	p.Asn192llefs*12	
()	3	c.614T>G	p.Met205Arg	0.00
	3	c.629A>G	p.Asn210Ser	0.13
	3	c.632A>G	p.Tyr211Cys	0.00
	3	c.641C>T	p.Ala214Val	32.56
	IVS3	c.652+1G>A		
	IVS3	c.653-2A>C		

c.656C>A

c.674C>T

4

4

p.Thr219Lys

p.Ser225Leu

0.0000042	0	
		damaging
		damaging
0.0000082	0	
0.0000494	0	benign
0.0000081	0	damaging
0.0000330	0	damaging
0.0000433	0	
0.0000082	0	damaging
0.0000577	0	damaging
0.0000447	0	damaging
0.0000330	0	damaging
0.0001071	0	damaging
		benign
0.0000041	0	damaging
0.0000494	0	
		damaging
0.0000203	0	damaging
0.0000081	0	damaging
0.0007989	2	damaging
0.0000122	0	
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28.25

11.34

(Li, et al., 2010)

(Li, et al., 2010)

(Han, et al., 2014) (Rose, et al., 2012)

(Koizumi, et al., 1999; Nezu, et al., 1999; Tang, et al., 1999)

(Amat di San Filippo, et al., 2006)

(Dobrowolski, et al., 2005)

(Lamhonwah, et al., 2002;

Wang, et al., 2000c)

(Burwinkel, et al., 1999)

(Han, et al., 2014)

(Li, et al., 2010)

(Li, et al., 2010)

(Li, et al., 2010)

(Vaz, et al., 1999)

(Lamhonwah, et al., 2002)

(Koizumi, et al., 1999)

(Han, et al., 2014)

(Li, et al., 2010)

(Han, et al., 2014)

4	c.680G>A	p.Arg227His	7.25	(Li, et al., 2010)	0.0000577
4	c.688T>C	p.Phe230Leu	0.80	(Li, et al., 2010)	0.0000108
4	c.692C>T	p.Ser231Phe	0.00	*	
4	c.695C>T	p.Thr232Met	19.14	(Dobrowolski, et al., 2005)	0.0001318
4	c.718G>A	p.Ala240Thr	2.06	(Li, et al., 2010)	
4	c.725G>T	p.Gly242Val	0.00	(Wang, et al., 2000c)	
4	c.740C>G	p.Pro247Arg	0.07	*	
4	c.745_748del	p.Phe249Leufs*13		(Han, et al., 2014)	
4	c.760C>T	p.Arg254*		(Tang, et al., 2002)	0.0001071
4	c.761G>A	p.Arg254Gin	34.24	*	0.0001236
4	c.768G>A	p.Trp256*		(Amat di San Filippo, et al., 2006)	
4	c.769C>T	p.Arg257Trp	7.65	(Li, et al., 2010)	0.0000906
4	c.791C>T	p.Thr264Met	46.00	(Amat di San Filippo, et al., 2008)	0.0000330
4	c.791C>G	p.Thr264Arg	2.48	(Li, et al., 2010)	0.0000406
4	c.806delT	p.Leu269Argfs*27		(Cederbaum, et al., 2002)	
4	c.806T>C	p.Leu269Pro	44.86	*	
IVS4	c.824+13T>C			(Mutlu-Albayrak, et al., 2015)	0.4346000
5	c.825G>A	p.Trp275*		(Dobrowolski, et al., 2005)	0.0000082
5	c.839delC	p.Arg282Aspfs*14		(Lamhonwah, et al., 2002)	0.0000082
5	c.839C>T	p.Ser280Phe	0.53	(Amat di San Filippo, et al., 2006)	0.0000081
5	c.844C>T	p.Arg282*		(Burwinkel, et al., 1999; Vaz, et al., 1999) (Wang, et al., 1999)	0.0000494
5	c.844delC	p.Val295*		(Komlosi, et al., 2009)	
5	c.845G>A	p.Arg282GIn	5.60	(Amat di San Filippo, et al., 2006)	
5	c.847T>C; T>A	p.Trp283Arg	0.45	(Mayatepek, et al., 2000)	
5	c.849G>T	p.Trp283Cys	0.00	(Koizumi, et al., 1999)	
5	c.865C>T	p.Arg289*		(Dobrowolski, et al., 2005)	0.0000082
5	c.902C>A	p.Ala301Asp	0.33	(Wang, et al., 2000c)	
5	c.934A>G	p.lle312Val	67.14	(Amat di San Filippo, et al., 2008; Li, et al., 2010)	0.0008072
5	c.949G>A	p.Glu317Lys	100.00	(Amat di San Filippo, et al., 2008)	0.0000082
5	c.955C>T	p.Gln319*		(Li, et al., 2010)	0.0000041
6	c.1008delA	p.Thr337Profs*12		(Lamhonwah, et al., 2002)	
6	c.1043T>C	p.lle348Thr	57.69	*	0.0000496
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7    c.1064C>T    p.Ser358Leu    1.55    (Li, et al., 2010)    0.0000081      7    c.1072T>A    p.Tyr35Xan    0.08    (Li, et al., 2010)    0.0000082      7    c.108ET>C    p.Leu36SPro    0.00    (Akpinar, et al., 2010)    0.0000082      7    c.1195CT    p.Leu36SPro    0.46    (Arrard IS an Filippo, et al., 2010)    0.0000082      7    c.1195C>T    p.Arg339Try    4.54    (El-Heittab, et al., 2010)    0.000082      7    c.108C>T    p.Arg339Gin    0.45    (Wang, et al., 1089)    0.000082      7    c.108C>A    p.Arg339Gin    0.45    (Wang, et al., 2010)    0.000082      7    c.1202_1203insA    p.Tyr401*    (Lamforwah, et al., 2001)    0.000082      7    c.1362To    p.Ser412Giy    107.72    *       8    c.1362Colin    p.Alg439Gily    0.52    *       8    c.1382T>G    p.Tyr440Met    0.00    (Lamforwah, et al., 2010)    0.00000162      8    c.1382CoT    p.Vat439Gily    0.	6	c.1051T>C	p.Trp351Arg	0.00	(Wang, et al., 2000c)	
7    c.1088T>C    p.Leu383Pro    0.00    (Akpinar, et al., 2010)    0.0000041      7    c.1161T>G    p.Tyr387*    (Tang, et al., 2002)    0.0000323      7    c.1175_1177del    p.Leu384del    5.29    *    0.0000323      7    c.1195C>T    p.Pr0398Leu    0.46    (Amat di San Filippo, et al., 2010)    0.000082      7    c.1195C>T    p.Arg399Fry    4.54    (El-Hattab, et al., 2010)    0.000082      7    c.1196C>A    p.Arg399Gin    0.45    (Wang, et al., 2010)    0.000082      7    c.1202_1203insA    p.Tyr401*    (Larrhonwah, et al., 2002)    0.000082      7    c.1267+3_1267+23del    (Dobrowolak, et al., 2001)    0.000082      8    c.1316T>G    p.Val439Giy    0.52    *      8    c.1316T>G    p.Val439Giy    0.52    *      8    c.1316T>G    p.Val439Giy    0.52    *    0.00000165      8    c.1316T>G    p.Val4421le    14.83    (El-Hattab, et al., 2010)    0.0000001      8    c.1	7	c.1064C>T	p.Ser355Leu	1.55	(Li, et al., 2010)	0.0000081
7    c.1161T-G    p.Tyr387*    (Tang, et al., 2002)    0.0000323      7    c.1175_[1177del    p.Leu394del    5.29    *    0.000082      7    c.1193C>T    p.Pro398Lou    0.46    (Amat di San Filippo, et al., 2010)    0.000082      7    c.1195C>T    p.Arg399Try    4.54    (El-Hattab, et al., 2010)    0.000082      7    c.1196C>A    p.Arg399Gin    0.45    (Wang, et al., 2010)    0.000082      7    c.1202_1203insA    p.Tyr401*    (Larrhorivah, et al., 2002)    0.000082      7    c.12127_232del    (Dobrowolski, et al., 2005)    0.000082    0.000082      7    c.12274>3    p.Ser412Gly    107.72    *    0.000082      8    c.1316T>G    p.Val439Gly    0.52    *    0.0000165      8    c.1316T>G    p.Val439Gly    0.52    *    0.000006165      8    c.1324_1325delGCin    p.Ala442le    14.83    (El-Hattab, et al., 2010)    0.0000041      8    c.13462>T    p.Val448Phe    0.58    (Mayatep6k, et al., 20	7	c.1072T>A	p.Tyr358Asn	0.08	(Li, et al., 2010)	0.0000082
7    c.1175_1177del    p.Leu394del    5.29    ·    0.0000882      7    c.1195C>T    p.Pro398Leu    0.46    (Amat di San Filippo, et al., 2010)    0.0000882      7    c.1195C>T    p.Arg399Try    4.54    (El-Hattab, et al., 2010)    0.0000882      7    c.1195C>T    p.Arg399Gin    0.45    (Wang, et al., 2001)    0.000082      7    c.1202_1203InsA    p.Tyr401*    (Lamforoweh, et al., 2002)    0.000082      7    c.1264A>G    p.Ser412Gly    107.72    *       8    c.1316T>G    p.Val439Gly    0.52    *       8    c.1316T>G    p.Val430H    0.34    (Li, et al., 2010)        8    c.1324_1325delGCin    p.Aie42lie    14.83    (El-Hattab, et al., 2002)	7	c.1088T>C	p.Leu363Pro	0.00	(Akpinar, et al., 2010)	0.0000041
T. H.	7	c.1161T>G	p.Tyr387*		(Tang, et al., 2002)	0.0000323
7  c.1193C-1  p.Pr0382.01  0.48  2006)  0.0000741    7  c.1196G>A  p.Arg399Try  4.54  (EI-Hattab, et al., 2010)  0.0000741    7  c.1202_1203insA  p.Tyr401*  (Lamhonwah, et al., 2002)  0.000082    7  c.1202_1203insA  p.Tyr401*  (Uanhonwah, et al., 2002)  0.000082    7  c.1267+3_1267+23del  (Dobrowolski, et al., 2005)  0.000082    8  c.1302delG  p.Siy435Alafs*24  (Wang, et al., 1999)  0.0000185    8  c.1316T>G  p.Val439Giy  0.52  *    8  c.1316CT  p.Thr440Met  0.00  (Lamhonwah, et al., 2002)  0.0000185    8  c.1324_1325delGCIn sAT  p.Nat4201e  14.83  (EI-Hattab, et al., 2010)  0.0000041    8  c.132GST  p.Val446Phe  0.58  (Mayatepek, et al., 2000)  0.000022    8  c.134GST  p.Val446Phe  0.58  (Mayatepek, et al., 2000)  0.00003247    8  c.134GS-T  p.Val446Phe  0.58  (Mayatepek, et al., 2000)  0.00003247    8  c.1346D>G  p.Tyr449Asp  16.56	7	c.1175_1177del	p.Leu394del	5.29	*	0.0000082
7    c.1196G>A    p.Arg399Gin    0.45    (Wang, et al., 2001)    0.0000182      7    c.1202_1203insA    p.Tyr401*    (Lamhonwah, et al., 2002; Wang, et al., 1999)    0.0000082      7    c.1234A>G    p.Ser412Giy    107.72    *      NS7    c.1267+3_1267+23del    (Dobrowolski, et al., 2005)    0.0000082      8    c.1302delG    p.Gly435Alafs'24    (Wang, et al., 1999)    0.0000165      8    c.1316T>G    p.Val439Gly    0.52    *    0.0000165      8    c.1316T>G    p.Val439Gly    0.52    *    0.0000165      8    c.1324_1325delGCin    p.Ala442lle    14.83    (EI-Hattab, et al., 2010)    0.00000165      8    c.1326ST    p.Val446Phe    0.58    (Mayatepek, et al., 2000)    0.0000041      8    c.134GA>G    p.Tyr447Cys    0.00    (Arnat di San Filippo and Longo, 2004; Rahbeeni, et al., 2002)    0.00003247      8    c.134GSA    p.Glu452Lys    2.14    (Wang, et al., 2004)    0.0000041      8    c.134GA>A    p.Glu452Lys    2.14	7	c.1193C>T	p.Pro398Leu	0.46		0.0000989
7  c.1202_1203insA  p.Tyr401*  (Lamhonwah, et al., 2002; Wang, et al., 1999)  0.000082    7  c.1267+3_1267+23del  (Dabrowolski, et al., 2005)  0.000082    8  c.1302delG  p.Gly435Alafs*24  (Wang, et al., 1999)  0.000082    8  c.1302delG  p.Val439Gly  0.52  *  0.0000165    8  c.1316T>G  p.Val439Gly  0.52  *  0.0000165    8  c.1316T>G  p.Val439Gly  0.52  *  0.0000165    8  c.1324_1325delGCin  p.Ala442lle  14.83  (El-Hattab, et al., 2010)  0.0000041    8  c.1320A>G  p.Tyr447Cys  0.00  (Amat di San Filippo and Longo, 2004; Rahbeeni, et al., 2002)  0.0000041    8  c.134GST  p.Val448Leu  10.62  *  *  *    8  c.134GSA  p.Gly462Val  2.80  *  *  *    8  c.134GSA  p.Gly462Val  2.80  *  *  *    8  c.134GSA  p.Gly462Val  2.80  *  *  *    8  c.1346CSG  p.Val486Thrfs*29	7	c.1195C>T	p.Arg399Try	4.54	(El-Hattab, et al., 2010)	0.0000741
r  c. 122_1233ISA  p.1yH01  Wang. et al., 1999)  0.0000002    7  c.1267+3_1267+23del  (Dobrowolski, et al., 2005)    8  c.1302delG  p.Gly435Alafs*24  (Wang, et al., 1999)    8  c.1316T>G  p.Val439Gly  0.52    8  c.1319C>T  p.Thr440Met  0.00  (Lamhonwah, et al., 2002)  0.0000165    8  c.1324_1325deliGCin  p.Ala442lle  14.83  (El-Hattab, et al., 2010)  0.0000165    8  c.1336G>T  p.Val446Phe  0.58  (Mayatepek, et al., 2000)  0.0000041    8  c.1340A>G  p.Tyr447Cys  0.00  2004; Rahbeeni, et al., 2000)  0.0000041    8  c.1342G>T  p.Val446Phe  0.58  (Mayatepek, et al., 2000)  0.000022    8  c.1342G>T  p.Val448Leu  10.62  *  -    8  c.1342G>A  p.Gly452Y  2.14  (Wang, et al., 2010)  0.00003247    8  c.1342G>A  p.Gly452YS  2.14  (Wang, et al., 2010)  0.00003247    8  c.1364C>G  p.Pro455Arg  0.18  (L. et al., 2010)  0.000072	7	c.1196G>A	p.Arg399GIn	0.45	(Wang, et al., 2001)	0.0000162
IVS7    c.1267+3_1267+23del    (Dobrowolski, et al., 2005)      8    c.1302delG    p.Gly435Alafs*24    (Wang, et al., 1999)      8    c.1316T>G    p.Val439Gly    0.52    *      8    c.1319C>T    p.Thr440Met    0.00    (Lamhonwah, et al., 2002)    0.0000165      8    c.1324_1325delGCIn    p.Ala442lle    14.83    (El-Hattab, et al., 2010)    0.0000165      8    c.1324_1325delGCIn    p.Ala442lle    14.83    (El-Hattab, et al., 2010)    0.00000000000000000000000000000000000	7	c.1202_1203insA	p.Tyr401*			0.0000082
8  c.1302delG  p.Gly435Alafs*24  (Wang, et al., 1999)    8  c.1316T>G  p.Val439Giy  0.52  *    8  c.1319C>T  p.Thr440Met  0.00  (Lamhonwah, et al., 2002)  0.0000165    8  c.1324_1325delGCin  p.Ala442lie  14.83  (El-Hattab, et al., 2010)  0.0000041    8  c.1326DFT  p.Val446Phe  0.58  (Mayatepek, et al., 2000)  0.0000041    8  c.1326DFT  p.Val446Phe  0.58  (Mayatepek, et al., 2002)  0.0000041    8  c.1340A>G  p.Tyr447Cys  0.00  2004; Rahbeeni, et al., 2002)  0.0000247    8  c.1342G>T  p.Val448Leu  10.62  *  -    8  c.1342G>T  p.Val448Leu  10.62  *  -    8  c.1342G>T  p.Val448Leu  10.62  *  -    8  c.1342G>A  p.Glu452Lys  2.14  (Wang, et al., 2002)  0.00003247    8  c.1364C>G  p.Pro455Arg  0.18  (Li, et al., 2010)  0.000072    8  c.1372delG  p.Val465Thrfs*29  *  -  -<	7	c.1234A>G	p.Ser412Gly	107.72	*	
8    c.1316T>G    p.Val439Gly    0.52    *      8    c.1316T>G    p.Thr440Met    0.00    (Lamhonwah, et al., 2002)    0.0000165      8    c.1324_1325delGCin sAT    p.Ala442lle    14.83    (El-Hattab, et al., 2010)    0.0000165      8    c.13267>G    p.Phe443Val    0.34    (Li, et al., 2010)    0.0000041      8    c.1336G>T    p.Val446Phe    0.58    (Mayatepek, et al., 2000)    0.0000041      8    c.1340A>G    p.Tyr447Cys    0.00    (Amat di San Filippo and Longo, 2004; Rahbeeni, et al., 2002)    0.00003247      8    c.1342G>T    p.Val448Leu    10.62    *    0.00003247      8    c.1354G>A    p.Glu452Lys    2.14    (Wang, et al., 2002)    0.0000424      8    c.1384G>A    p.Glu452Lys    2.14    (Wang, et al., 2010)    0.000072      8    c.1385G>T    p.Glu452Val    2.80    *       8    c.1385G>T    p.Glu452Thrfs*29    *        8    c.1382_14093delinsCA    p.Val465Thrfs*29	IVS7	c.1267+3_1267+23del			(Dobrowolski, et al., 2005)	
8    c.1319C>T    p.Thr440Met    0.00    (Lamhonwah, et al., 2002)    0.0000165      8    c.1324_1325delGCin sAT    p.Ala442lle    14.83    (EI-Hattab, et al., 2010)    0.0000165      8    c.1326G>T    p.Nal442Ple    14.83    (EI-Hattab, et al., 2010)    0.0000041      8    c.1336G>T    p.Val446Phe    0.58    (Mayatepek, et al., 2000)    0.0000041      8    c.1340A>G    p.Tyr447Cys    0.00    (Amat di San Filippo and Longo, 2004; Rahbeeni, et al., 2002)    0.0003247      8    c.1342G>T    p.Val448Leu    10.62    *    0.0000494      8    c.1384G>A    p.Glu452Lys    2.14    (Wang, et al., 2000a)    0.00003247      8    c.1384G>A    p.Glu452Lys    2.14    (Wang, et al., 2010)    0.000072      8    c.1385G>T    p.Gly462Val    2.80    *    *      8    c.1385G>T    p.Gly462Val    2.80    *      8    c.1382_1409delinsCA    p.Val465Thrfs*29    *    *      8    c.1400C>G    p.Ser467Cys    16.56	8	c.1302delG	p.Gly435Alafs*24		(Wang, et al., 1999)	
8  c.1324_1325delGCin sAT  p.Ala442lle  14.83  (El-Hattab, et al., 2010)    8  c.1327T>G  p.Phe443Val  0.34  (Li, et al., 2010)    8  c.1326G>T  p.Val446Phe  0.58  (Mayatepek, et al., 2000)    8  c.1340A>G  p.Tyr447Cys  0.00  (Amat di San Filippo and Longo, 2004; Rahbeeni, et al., 2002)    8  c.1342G>T  p.Val448Leu  10.62  *    8  c.1345G>A  p.Glu452Lys  2.14  (Wang, et al., 2000a)  0.000041    8  c.1364G>A  p.Glu452Lys  2.14  (Wang, et al., 2000a)  0.0000494    8  c.1364G>G  p.Val458*  (Han, et al., 2010)  0.000072    8  c.1385G>T  p.Glu452Lys  2.80  *    8  c.1385G>T  p.Gly462Val  2.80  *    8  c.1382_1409delinsCA  p.Val465Thrfs*29  *     8  c.1400C>G  p.Ser467Cys  16.56  (Koizumi, et al., 1999)  0.0000165    8  c.1409C>T  p.Ser470Phe  0.00  (Lamhonwah, et al., 2002)  0.0000165    8  c.1412G>C	8	c.1316T>G	p.Val439Gly	0.52	*	
sAT    p.Aid442iie    14.63    (El-Hattab, et al., 2010)      8    c.1327T>G    p.Phe443Val    0.34    (Li, et al., 2010)      8    c.1326C>T    p.Val446Phe    0.58    (Mayatepek, et al., 2000)      8    c.1340A>G    p.Tyr447Cys    0.00    (Amat di San Filippo and Longo, 2004; Rahbeeni, et al., 2002)      8    c.1342G>T    p.Val448Leu    10.62    *      8    c.1342G>T    p.Val448Leu    10.62    *      8    c.1345T>G    p.Tyr447Cys    2.14    (Wang, et al., 2000a)    0.0000247      8    c.1354G>A    p.Glu452Lys    2.14    (Wang, et al., 2000a)    0.000072      8    c.1364C>G    p.Pro455Arg    0.18    (Li, et al., 2010)    0.0000072      8    c.1372delG    p.Val465Thrfs*29    *         8    c.1392_1409dellinsCA    p.Val465Thrfs*29    *         8    c.1400C>G    p.Ser467Cys    16.56    (Koizumi, et al., 1999)	8	c.1319C>T	p.Thr440Met	0.00	(Lamhonwah, et al., 2002)	0.0000165
8  c.1336G>T  p.Val446Phe  0.58  (Mayatepek, et al., 2000)    8  c.1340A>G  p.Tyr447Cys  0.00  (Arnat di San Filippo and Longo, 2004; Rahbeeni, et al., 2002)    8  c.1342G>T  p.Val448Leu  10.62  *    8  c.1342G>T  p.Val448Leu  10.62  *    8  c.1345T>G  p.Tyr449Asp  16.56  (Arnat di San Filippo and Longo, 2004)  0.0003247    8  c.1354G>A  p.Glu452Lys  2.14  (Wang, et al., 2000a)  0.0000494    8  c.1364C>G  p.Pro455Arg  0.18  (Li, et al., 2010)  0.0000072    8  c.1372delG  p. Val458*  (Han, et al., 2014)  0.0000072    8  c.1392_1409delinsCA  p.Val465Thrfs*29  *  *    8  c.1403C>G  p.Ser467Cys  16.56  (Koizumi, et al., 1999)  0.0000081    8  c.1403C>G  p.Thr468Arg  0.64  (Lamhonwah, et al., 2002)  0.0000081    8  c.1403C>C  p.Arg471Pro  0.00  (Filippo, et al., 2011)  0.0000165    8  c.1412G>C  p.Arg471His  1.40  (Spiekerko	8		p.Ala442lle	14.83	(El-Hattab, et al., 2010)	
8  c.1340A>G  p.Tyr447Cys  0.00  (Amat di San Filippo and Longo, 2004; Rahbeeni, et al., 2002)    8  c.1342G>T  p.Val448Leu  10.62  *    8  c.1342G>T  p.Val448Leu  10.62  *    8  c.1345T>G  p.Tyr447Cys  2.00  (Amat di San Filippo and Longo, 2004)  0.0003247    8  c.1345T>G  p.Tyr449Asp  16.56  (Amat di San Filippo and Longo, 2004)  0.00003247    8  c.1354G>A  p.Glu452Lys  2.14  (Wang, et al., 2000a)  0.0000494    8  c.1364C>G  p.Pro455Arg  0.18  (Li, et al., 2010)  0.0000072    8  c.1372delG  p.Val485*  (Han, et al., 2014)  0.0000072    8  c.1385G>T  p.Gly462Val  2.80  *    8  c.1302_1409delinsCA  p.Val465Thrfs*29  *    8  c.1400C>G  p.Ser467Cys  16.56  (Koizumi, et al., 1999)  0.00001565    8  c.1403C>G  p.Thr468Arg  0.64  (Lamhonwah, et al., 2002)  0.0000165    8  c.1412G>C  p.Arg471Pro  0.00  (Filippo, et al., 2011) <td< td=""><td>8</td><td>c.1327T&gt;G</td><td>p.Phe443Val</td><td>0.34</td><td>(Li, et al., 2010)</td><td></td></td<>	8	c.1327T>G	p.Phe443Val	0.34	(Li, et al., 2010)	
8  c.1340A-3  p.1yr4476ys  0.00  2004; Rahbeeni, et al., 2002)    8  c.1342G>T  p.Val448Leu  10.62  *    8  c.1345T>G  p.Tyr449Asp  16.56  (Amat di San Filippo and Longo, 2004)  0.0003247    8  c.1354G>A  p.Glu452Lys  2.14  (Wang, et al., 2000a)  0.0000494    8  c.1354G>A  p.Glu452Lys  2.14  (Wang, et al., 2010)  0.0000072    8  c.1372delG  p. Val458*  (Han, et al., 2014)  0.0000072    8  c.1385G>T  p.Gly462Val  2.80  *    8  c.1302_1409delinsCA  p.Val465Thrfs*29  *    8  c.1400C>G  p.Ser467Cys  16.56  (Koizumi, et al., 1999)  0.00001565    8  c.1403C>G  p.Thr468Arg  0.64  (Lamhonwah, et al., 2002)  0.0000165    8  c.1412G>C  p.Arg471Pro  0.00  (Filippo, et al., 2011)  0.0000165    8  c.1412G>A  p.Arg471His  1.40  (Spiekerkoetter, et al., 2003)  0.0000165	8	c.1336G>T	p.Val446Phe	0.58	(Mayatepek, et al., 2000)	0.0000041
8    c.1345T>G    p.Tyr449Asp    16.56    (Amat di San Filippo and Longo, 2004)    0.0003247      8    c.1354G>A    p.Glu452Lys    2.14    (Wang, et al., 2000a)    0.0000494      8    c.1364C>G    p.Pro455Arg    0.18    (Li, et al., 2010)    0.0000072      8    c.1372delG    p. Val458*    (Han, et al., 2014)    0.0000072      8    c.1385G>T    p.Gly462Val    2.80    *      8    c.1392_1409delinsCA    p.Val465Thrfs*29    *      8    c.1392_1409delinsCA    p.Val465Thrfs*29    *      8    c.1400C>G    p.Ser467Cys    16.56    (Koizumi, et al., 1999)    0.00001565      8    c.1403C>G    p.Thr468Arg    0.64    (Lamhonwah, et al., 2002)    0.0000165      8    c.1412G>C    p.Arg471Pro    0.00    (Filippo, et al., 2011)    0.0000165      8    c.1412G>A    p.Arg471His    1.40    (Spiekerkoetter, et al., 2003)    0.0000165	8	c.1340A>G	p.Tyr447Cys	0.00	(Amat di San Filippo and Longo, 2004; Rahbeeni, et al., 2002)	
6  C.13431 > G  p.1y1443Asp  16.56  2004)  0.0003247    8  c.1354G>A  p.Glu452Lys  2.14  (Wang, et al., 2000a)  0.0000494    8  c.1364C>G  p.Pro455Arg  0.18  (Li, et al., 2010)  0.0000072    8  c.1372delG  p. Val458*  (Han, et al., 2014)  0.0000072    8  c.1385G>T  p.Gly462Val  2.80  *    8  c.1392_1409delinsCA  p.Val465Thrfs*29  *    8  c.1400C>G  p.Ser467Cys  16.56  (Koizumi, et al., 1999)  0.00001565    8  c.1403C>G  p.Thr468Arg  0.64  (Lamhonwah, et al., 2002)  0.0000165    8  c.1409C>T  p.Ser470Phe  0.00  (Lamhonwah, et al., 2002)  0.0000165    8  c.1412G>C  p.Arg471Pro  0.00  (Filippo, et al., 2011)  0.0000165    8  c.1412G>A  p.Arg471His  1.40  (Spiekerkoetter, et al., 2003)  0.0000165	8	c.1342G>T	p.Val448Leu	10.62	*	
8  c.1364C>G  p.Pro455Arg  0.18  (Li, et al., 2010)  0.0000072    8  c.1372delG  p. Val458*  (Han, et al., 2014)	8	c.1345T>G	p.Tyr449Asp	16.56		0.0003247
8  c.1372delG  p. Val458*  (Han, et al., 2014)    8  c.1385G>T  p.Gly462Val  2.80  *    8  c.1392_1409delinsCA  p.Val465Thrfs*29  *    8  c.1392_1409delinsCA  p.Val465Thrfs*29  *    8  c.1400C>G  p.Ser467Cys  16.56  (Koizumi, et al., 1999)  0.0001565    8  c.1403C>G  p.Thr468Arg  0.64  (Lamhonwah, et al., 2002)  0.0000081    8  c.1409C>T  p.Ser470Phe  0.00  (Lamhonwah, et al., 2002)  0.0000165    8  c.1412G>C  p.Arg471Pro  0.00  (Filippo, et al., 2011)  0.0000165    8  c.1412G>A  p.Arg471His  1.40  (Spiekerkoetter, et al., 2003)  0.0000165	8	c.1354G>A	p.Glu452Lys	2.14	(Wang, et al., 2000a)	0.0000494
8  c.1385G>T  p.Giy462Val  2.80  *    8  c.1392_1409delinsCA  p.Val465Thrfs*29  *    8  c.1400C>G  p.Ser467Cys  16.56  (Koizumi, et al., 1999)  0.0001565    8  c.1403C>G  p.Thr468Arg  0.64  (Lamhonwah, et al., 2002)  0.0000081    8  c.1409C>T  p.Ser470Phe  0.00  (Lamhonwah, et al., 2002)  0.0000165    8  c.1412G>C  p.Arg471Pro  0.00  (Filippo, et al., 2011)  0.0000165    8  c.1412G>A  p.Arg471His  1.40  (Spiekerkoetter, et al., 2003)  0.0000165	8	c.1364C>G	p.Pro455Arg	0.18	(Li, et al., 2010)	0.0000072
8  c.1392_1409delinsCA  p.Val465Thrfs*29  *    8  c.1400C>G  p.Ser467Cys  16.56  (Koizumi, et al., 1999)  0.0001565    8  c.1403C>G  p.Thr468Arg  0.64  (Lamhonwah, et al., 2002)  0.0000081    8  c.1409C>T  p.Ser470Phe  0.00  (Lamhonwah, et al., 2002)  0.0000165    8  c.1412G>C  p.Arg471Pro  0.00  (Filippo, et al., 2011)  0.0000165    8  c.1412G>A  p.Arg471His  1.40  (Spiekerkoetter, et al., 2003)  0.000165	8	c.1372delG	p. Val458*		(Han, et al., 2014)	
8    c.1400C>G    p.Ser467Cys    16.56    (Koizumi, et al., 1999)    0.0001565      8    c.1403C>G    p.Thr468Arg    0.64    (Lamhonwah, et al., 2002)    0.0000081      8    c.1409C>T    p.Ser470Phe    0.00    (Lamhonwah, et al., 2002)    0.0000165      8    c.1412G>C    p.Arg471Pro    0.00    (Filippo, et al., 2011)    0.0000165      8    c.1412G>A    p.Arg471His    1.40    (Spiekerkoetter, et al., 2003)    0.000165	8	c.1385G>T	p.Gly462Val	2.80	*	
8    c.1403C>G    p.Thr468Arg    0.64    (Lamhonwah, et al., 2002)    0.0000081      8    c.1409C>T    p.Ser470Phe    0.00    (Lamhonwah, et al., 2002)    0.0000165      8    c.1412G>C    p.Arg471Pro    0.00    (Filippo, et al., 2011)    0.0000165      8    c.1412G>A    p.Arg471His    1.40    (Spiekerkoetter, et al., 2003)    0.0000165	8	c.1392_1409delinsCA	p.Val465Thrfs*29		*	
8    c.1409C>T    p.Ser470Phe    0.00    (Lamhonwah, et al., 2002)    0.0000165      8    c.1412G>C    p.Arg471Pro    0.00    (Filippo, et al., 2011)    0.0000165      8    c.1412G>A    p.Arg471His    1.40    (Spiekerkoetter, et al., 2003)    0.0000165	8	c.1400C>G	p.Ser467Cys	16.56	(Koizumi, et al., 1999)	0.0001565
8    c.1412G>C    p.Arg471Pro    0.00    (Filippo, et al., 2011)      8    c.1412G>A    p.Arg471His    1.40    (Spiekerkoetter, et al., 2003)	8	c.1403C>G	p.Thr468Arg	0.64	(Lamhonwah, et al., 2002)	0.0000081
8 c.1412G>A p.Arg471His 1.40 (Spiekerkoetter, et al., 2003)	8	c.1409C>T	p.Ser470Phe	0.00	(Lamhonwah, et al., 2002)	0.0000165
	8	c.1412G>C	p.Arg471Pro	0.00	(Filippo, et al., 2011)	
8 c.1427T>G p.Leu476Arg 0.00 (Mutlu-Albayrak, et al., 2015)	8	c.1412G>A	p.Arg471His	1.40	(Spiekerkoetter, et al., 2003)	
	8	c.1427T>G	p.Leu476Arg	0.00	(Mutlu-Albayrak, et al., 2015)	

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c.1433C>T	p.Pro478Leu	0.28	(Tang, et al., 1999)					damaging	damaging
c.1441G>T	p.Val481Phe	63.00	(Amat di San Filippo, et al., 2008)	-				benign	benign
c.1451-1G>A			(Nezu, et al., 1999)	-	0.0004777	0	_		
c.1462C>T	p.Arg488Cys	9.50	(Schimmenti, et al., 2007)		0.0000412	0	_	damaging	damaging
c.1463G>A	p.Arg488His	40.19	(Amat di San Filippo, et al., 2006)		0.0032620	4	_	damaging	damaging
c.1520T>C	p.Leu507Ser	4.85	(Li, et al., 2010)		0.0000323	0	_	damaging	damaging
c.1556_1559dup	p.lle521Hisfs*3		(Schimmenti, et al., 2007)	+			_		
c.1645C>T	p.Pro549Ser	23.79	(Amat di San Filippo, et al., 2008; Urban, et al., 2006)	-	0.0064990	28		benign	benign
	c.1441G>T c.1451-1G>A c.1462C>T c.1463G>A c.1520T>C c.1556_1559dup	c.1441G>T  p.Val481Phe    c.1451-1G>A    c.1462C>T  p.Arg488Cys    c.1463G>A  p.Arg488His    c.1520T>C  p.Leu507Ser    c.1556_1559dup  p.lle521Hisfs*3	c.1441G>T  p.Val481Phe  63.00    c.1451-1G>A	c.1441G>T  p.Val481Phe  63.00  (Amat di San Filippo, et al., 2008)    c.1451-1G>A  (Nezu, et al., 1999)    c.1452C>T  p.Arg488Cys  9.50  (Schimmenti, et al., 2007)    c.1463G>A  p.Arg488His  40.19  (Amat di San Filippo, et al., 2007)    c.1463G>A  p.Arg488His  40.19  (Amat di San Filippo, et al., 2006)    c.1520T>C  p.Leu507Ser  4.85  (Li, et al., 2010)    c.1556_1559dup  p.Ile521Hisfs*3  (Schimmenti, et al., 2007)    c.1645C>T  p.Bro£49Sor  23.79  (Amat di San Filippo, et al., 2007)	c.1441G>T  p.Val481Phe  63.00  (Amat di San Filippo, et al., 2008)    c.1451-1G>A  (Nezu, et al., 1999)    c.1462C>T  p.Arg488Cys  9.50  (Schimmenti, et al., 2007)    c.1463G>A  p.Arg488His  40.19  (Amat di San Filippo, et al., 2006)    c.1520T>C  p.Leu507Ser  4.85  (Li, et al., 2010)    c.1556_1559dup  p.lle521Hisfs*3  (Schimmenti, et al., 2007)	c.1441G>T  p.Val481Phe  63.00  (Amat di San Filippo, et al., 2008)    c.1451-1G>A  (Nezu, et al., 1999)  0.0004777    c.1452C>T  p.Arg488Cys  9.50  (Schimmenti, et al., 2007)    c.1463G>A  p.Arg488His  40.19  (Amat di San Filippo, et al., 2007)    c.1463G>A  p.Arg488His  40.19  (Amat di San Filippo, et al., 2006)    c.1520T>C  p.Leu507Ser  4.85  (Li, et al., 2010)  0.0000323    c.1556_1559dup  p.lle521Hisfs*3  (Schimmenti, et al., 2007)  0.0064990	c.1441G>T  p.Val481Phe  63.00  (Amat di San Filippo, et al., 2008)    c.1451-1G>A  (Nezu, et al., 1999)  0.0004777  0    c.1462C>T  p.Arg488Cys  9.50  (Schimmenti, et al., 2007)  0.0000412  0    c.1463G>A  p.Arg488His  40.19  (Amat di San Filippo, et al., 2006)  0.0032620  4    c.1520T>C  p.Leu507Ser  4.85  (Li, et al., 2010)  0.0000323  0    c.1556_1559dup  p.lle521Hisfs*3  (Schimmenti, et al., 2007)  0.0064090  28	c.1441G>T  p.Val481Phe  63.00  (Amat di San Filippo, et al., 2008)    c.1451-1G>A  (Nezu, et al., 1999)  0.0004777  0    c.1462C>T  p.Arg488Cys  9.50  (Schimmenti, et al., 2007)  0.0000412  0    c.1463G>A  p.Arg488His  40.19  (Amat di San Filippo, et al., 2006)  0.0032620  4    c.1520T>C  p.Leu507Ser  4.85  (Li, et al., 2010)  0.0000323  0    c.1556_1559dup  p.lle521Hisfs*3  (Schimmenti, et al., 2007)  0.0064090  28	c.1441G>T  p.Val481Phe  63.00  (Amat di San Filippo, et al., 2008)  benign    c.1451-1G>A  (Nezu, et al., 1999)  0.0004777  0    c.1462C>T  p.Arg488Cys  9.50  (Schimmenti, et al., 2007)  0.0000412  0    c.1463G>A  p.Arg488His  40.19  (Amat di San Filippo, et al., 2006)  0.0032620  4  damaging    c.1520T>C  p.Leu507Ser  4.85  (Li, et al., 2010)  0.0000323  0  damaging    c.1556_1559dup  p.lle521Hisfs*3  (Schimmenti, et al., 2007)  0.0064990  28  benign

**Table 4. Correlation between functional studies and predictions of variant analysis software programs.** Polyphen-2 and SIFT are software programs commonly used to predict the pathogenicity of missense variants. Functional studies of carnitine transport in CHO cells expressing 90 missense variants and 2 in-frame deletions identified in our patients were compared to the predictions of Polyphen-2 and SIFT. Carnitine transport activity of 20% or less that the wild-type transporter was considered pathogenic. Both programs were effective in separating benign from damaging variants (p=0.0005 for Polyphen-2 and p=0.0009 for SIFT using Fisher exact text). However, about 20% of the variants were incorrectly predicted by either program, with a more marked error in the prediction of benign variants. Polyphen-2 and SIFT provided a concordant prediction of the effect of a variant in 80% of the cases (74 of 92 variants). In 86% of the cases the results provided by the two programs were concordant with the results obtained analyzing carnitine transport in CHO cells, with 57/59 (97%) variants correctly identified as pathogenic, but only 7/15 (47%) correctly identified as benign.

POLYPHEN-2 VARIANT ANALYSIS SOFTWARE					
90 variants analyzed	benign	damaging			
transport <20% of WT-OCTN2	3	68			
transport >20% of WT-OCTN2	7	12			
75/90 (83%) correctly identified. Two-tailed p value 0.0005					

SIFT VARIANT ANALYSIS SOFTWARE						
92 variants analyzed	benign	damaging				
transport <20% of WT-OCTN2	13	60				
transport >20% of WT-OCTN2	11	8				
71/92 (77%) correctly identified. Two-tailed p value 0.0009						

# POLYPHEN-2 AND SIFT CONCORDANT PREDICTIONS

	benign	damaging
transport <20% of WT-OCTN2	2	57
transport >20% of WT-OCTN2	7	8
64/74 (86%) correctly identified.	Two-tailed	p value 0.0001