Functional Characterization and Categorization of Missense Mutations that Cause Methylmalonyl-CoA Mutase (MUT) Deficiency

Patrick Forny,1,2 D. Sean Froese,1,3 Terttu Suormala,1 Wyatt W. Yue,2,* and Matthias R. Baumgartner1,2,*

1Division for Metabolic Disorders and Children’s Research Center, University Children’s Hospital, Zurich, Switzerland; 2Zurich Center for Integrative Human Physiology, University of Zurich, Switzerland; 3Structural Genomics Consortium, University of Oxford, UK

Communicated by Elizabeth F. L Neufeld
Received 6 June 2014; accepted revised manuscript 27 July 2014.
Published online 14 August 2014 in Wiley Online Library (www.wiley.com/humanmutation). DOI: 10.1002/humu.22633

ABSTRACT: Methylmalonyl-CoA mutase (MUT) is an essential enzyme in propionate catabolism that requires adenosylcobalamin as a cofactor. Almost 250 inherited mutations in the MUT gene are known to cause the devastating disorder methylmalonic aciduria; however, the mechanism of dysfunction of these mutations, more than half of which are missense changes, has not been thoroughly investigated. Here, we examined 23 patient missense mutations covering a spectrum of exonic/structural regions, clinical phenotypes, and ethnic populations in order to determine their influence on protein stability, using two recombinant expression systems and a thermostability assay, and enzymatic function by measuring MUT activity and affinity for its cofactor and substrate. Our data stratify MUT missense mutations into categories of biochemical defects, including (1) reduced protein level due to misfolding, (2) increased thermolability, (3) impaired enzyme activity, and (4) reduced cofactor response in substrate turnover. We further demonstrate the stabilization of wild-type and thermolabile mutants by chemical chaperones in vitro and in bacterial cells. This in-depth mutation study illustrates the tools available for MUT enzyme characterization, guides future categorization of further missense mutations, and supports the development of alternative, chaperone-based therapy for patients not responding to current treatment.

Hum Mutat 35:1449–1458, 2014. Published 2014 Wiley Periodicals, Inc.*

KEY WORDS: methylmalonic aciduria; methylmalonyl-CoA mutase; MUT; cobalamin; thermolability

Introduction

Mitochondrial methylmalonyl-CoA mutase (MUT, EC 5.4.99.2) catalyzes the reversible isomerisation of t-methylmalonyl-CoA to succinyl-CoA, requiring vitamin B12 (cobalamin) in the form of adenosylcobalamin (AdoCbl) as a cofactor. In humans, this reaction represents an important step in propionate catabolism, funneling metabolites from the breakdown of amino acids (valine, isoleucine, methionine, and threonine), odd-chain fatty acids, and the side chain of cholesterol into the tricarboxylic acid cycle. The MUT enzyme is highly conserved from bacteria to human and is well studied at the enzymatic [Fenton et al., 1982; Banerjee and Ragsdale, 2003] and structural level [Mancia et al., 1996; Froese et al., 2010b]. The importance of the MUT-catalyzed reaction is further underlined by the metabolic disorder methylmalonic aciduria (MMA), which is caused by a genetic defect in the MUT enzyme itself (MIM# 251000, MMA mut type), or in one of several proteins (MMAA, MMAB, MMADHC) involved in the uptake, modification, and delivery of AdoCbl to the MUT enzyme for its activity (MMA cblA type, MIM# 251100; MMA cblB type, MIM# 251110; MMA cblD-variant 2 MIM# 277410) [Fowler et al., 2008].

The human MUT gene (MIM# 609058; chromosome location 6p12–21.2) [Ledley et al., 1988a, b] is the site of almost 250 deleterious mutations reported to cause MMA [Froese and Gravel, 2010] (human gene mutation database, HGMD Professional version as of December 2013, hgmd.org). Patient cell lines can be assigned to the mut-type MMA by complementation analysis of fibroblast heterokaryons [Gravel et al., 1975; Willard et al., 1978]. Further biochemical characterization, applying MUT activity assay and propionate fixation, allows mut classification into two subtypes. Mutants with residual mutant activity in cell homogenates under saturating concentrations of AdoCbl, and whose ability to incorporate [1–14C]propionate into acid precipitable material in intact skin fibroblasts is responsive to supplementation of the culture medium with hydroxocobalamin, are designated mut− mutants with no residual MUT activity and no response of propionate incorporation to hydroxocobalamin are designated mut+ [Willard and Rosenberg, 1980]. mut+ patients often present in the newborn period with ketoadidosis, lethargy, repeated vomiting, coma or even death, and suffer from severe long-term complications such as renal failure and neurological impairments. On the contrary, mut− patients have a lower occurrence of mortality, morbidity, and long-term complications [Horster et al., 2007]. Dietary interventions, carnitine supplementation, and symptom management are the mainstay of current treatment, although pharmacological doses of hydroxocobalamin are given to some MUT-deficient patients [Horster,...]
et al., 2007). Although some mut patients might respond to this treatment, others—including most if not all mut patients—do not [Horster et al., 2007]. Alternative therapeutic approaches should therefore be investigated, evaluating therapeutic potential on the basis of a systematic characterization of individual mut and mut alleles. Nevertheless, the sheer quantity of mutations in the MUT gene—most of which are private and rare [Acquaviva et al., 2005; Worga et al., 2006; Lempp et al., 2007; Fowler et al., 2008]—means that such a large-scale analysis has not been forthcoming, with previous reports of in vitro MUT characterization on only a handful of mutations [e.g., Crane and Ledley, 1994; Janata et al., 1997].

Common to many other metabolic disorders (e.g., phenylketonuria (PKU) [Mitchell et al., 2011]; ornithine transcarbamylase deficiency [Schelochkiv et al., 2009]), the largest proportion of MUT mutations are missense changes (131 of 243, 54%; hgmd.org) whose effects on the protein are difficult to predict a priori [Froese and Gravel, 2010; Yue et al., 2014]. MUT is therefore an attractive target to interrogate the genotype-specific biochemical penalties for missense mutations at the protein level, an approach adopted for other metabolic enzymes [Pey et al., 2007; Pekkala et al., 2010; Shi et al., 2012]. With this in mind, we have chosen seven mut and 16 mut missense mutations for an in-depth analysis. On the basis of recombinant expression (Escherichia coli and human fibroblasts), thermal denaturation, and enzyme assays, we catalogue each mutation as defective in stability, activity, or both, and further define subcategories in each. To the best of our knowledge, this work represents the first large-scale biochemical categorization of MUT mutations and sets the stage for exploring the potential of small molecule therapeutics, an approach that is gaining wide interest to augment mutant enzyme activity [Gomes, 2012].

Materials and Methods

Cloning

For expression in E. coli, we used a previously described construct of MUT (called hMUT in [Froese et al., 2010b]). For expression in fibroblasts, the MUT cDNA sequence was purchased from GeneCopoeia (Rockville, MD, USA) (Construct ID: E0205; RefSeq NM_000255.3) and subsequently subcloned into pTracer-CMV2 (Invitrogen, Carlsbad, CA, USA) following manufacturer’s instructions, using forward and reverse primers (Microsynth, Balgach, Switzerland) described in [Suormala et al., 2004]. pTracer-MUT wild-type (wt) and mutant constructs were transiently transfected into the immortalized fibroblasts by electroporation with a transfection efficiency of 6%–24%, as determined by fluorescence-associated cell sorting measuring the proportion of cells coexpressing the green fluorescent protein from the pTracer construct. This is comparable to previous observations [Coelho et al., 2008]. Cells were harvested 48 hr after electroporation by trypsinization, washed twice with phosphate buffered saline, and stored frozen at −80°C until assayed for MUT activity or used for Western blotting.

Western Blot

Fibroblast lysates were obtained by sonication (details under MUT activity assay). Crude cell lysates (25 μg of protein per sample) were mixed with 2× Laemmlli Sample Buffer (Bio-Rad, Hercules, CA, USA) and heated to 96°C for 6 min. Proteins were separated by 10% SDS-PAGE, transferred to a Protran BA85 nitrocellulose membrane (Whatman, GE Healthcare), blocked at room temperature for 1 hr with buffer A (5% skimmed milk, 1.2% w/v Tris-base, 9% w/v NaCl, 0.2% Tween 20, pH 7.6), incubated with polyclonal mouse anti-human MUT (Abcam, Cambridge, UK) (1:1,000 in buffer A), or β-actin (Sigma–Aldrich, Buchs SG, Switzerland) (1:4,000 in buffer A) overnight, detected by incubation with goat anti-mouse antibody coupled to horseradish peroxidase (Santa Cruz, Biotechnology, Dallas, TX, USA) (1:5,000 in buffer A) for 1 hr, and visualized with ECL Western Blotting Detection Reagents (GE Healthcare) on a Gel Logic 6000 Pro (Carestream, Gland, Switzerland).

MUT Activity Assay

MUT activity was assayed in crude cell lysates by measuring the production of [14C]succinate from [14C]methylmalonyl-CoA by a modification of earlier described methods [Baumgartner et al., 1983; Causey and Bartlett, 1984]. All operations were performed in a dark room illuminated with a 15 W red Safelight bulb (Dr. Fischer, Diez/Lahn, Germany). Cell lysates were prepared by pipetting 5 mM potassium phosphate buffer (pH 7.4) on frozen cells and disrupting the pellet by sonication twice for 15 sec using the microprobe of an XL-2000 sonicator (Miscoxon, Qsonica, Newtown, CT, USA) at amplitude 1.5. Specific activities were measured...
in 50 μl reaction mixture containing 0.1 M potassium phosphate buffer (pH 7.4), 1 mM DL-2-[methyl-1-14C]methylmalonyl-CoA (American Radiolabeled Chemicals (ARC), St. Louis, MO, USA; specific activity 7.03 MBq/mmol in assay), and 50–100 μg cell proteins without (holo-MUT activity) and with (total MUT activity) 50 μM AdoCbl. To estimate Kf for AdoCbl, its concentration was varied between 0.0025 and 50 μM. To estimate Kf for methylmalonyl-CoA, its concentration was varied between 0.01 and 1.0 mM in the presence of 50 μM AdoCbl. Reactions were initiated by the addition of cell lysate, allowed to proceed for 30 min at 37°C, and terminated by addition of 6 μl 5 N KOH (Merck, Whitehouse Station, NJ, USA). Samples were reincubated for 15 min at 37°C to hydrolyze CoA derivatives, neutralized by adding 5 μl 15 N perchloric acid (Merck), and spiked with 5 μl of 1% solution (w/v) of succinic acid (Merck) in order to visualize succinate during HPLC separation. Samples were centrifuged to remove precipitate, 50 μl of the supernatant was injected into an Animex HPX-87H Ion Exclusion column (300 × 7.8 mm2; H-form, 9 μm, Bio-Rad), and [14C]succinate was separated from [14C]methylmalonate by elution with 0.5 mM H2SO4 at 30°C using a flow rate of 0.4 ml/min. Succinate (retention time 17 min) and methylmalonate (11 min) peaks were visualized at 210 nm using a UV detector; 0.2 ml fractions covering the succinate peak were collected and [14C]succinate was quantitated in a Tri-Carb C1 900TR scintillation spectrometer (PerkinElmer, Waltham, MA, USA) with Optiphase HiSafe 2 counting cocktail (PerkinElmer). Protein in cell lysates was determined by the Lowry method. MUT activity was expressed as nanomole succinate formed per minute and milligram of protein. Kf values for AdoCbl were determined using Eadie–Hofstee plotting.

Differential Scanning Fluorimetry

Purified homodimeric MUT was assayed for shifts in melting temperature (Tm), which is the midpoint transition from a folded to an unfolded state, as previously described [Niesen et al., 2007; Froese et al., 2010a]. Each mutant and wt protein was assayed in the apo state, or with preincubation of 50 μM AdoCbl (Sigma–Aldrich), or a combination of 50 μM AdoCbl and 500 μM malonyl-CoA (MCoA; Sigma–Aldrich). MCoA was used as a substrate analogue because it has been confirmed to bind to the substrate pocket (PDF: 2XIQ) [Froese et al., 2010b]. For chemical cosolvent screening, various concentrations of proline, betaine, sorbitol, TMAO, sucrose (up to 1,200 mM), and glycerol (up to 20%) were preincubated with protein before analysis.

Results

Rationale for Mutation Selection

From the repertoire of known MUT missense mutations, we selected 23 for this study (Table 1) with the rationale that they (1) cover both phenotypically severe mut+ (n = 7) and milder mut- (n = 16) subtypes, (2) are widely distributed across the gene (exons 2–13), and (3) include representative mutations prevalent in different ethnic populations, for example, African Americans (p.G717V and c.2150G>C), Caucasians (p.N219Y, c.655A>T; p.R369H, c.1106G>A; p.R694W, c.2080C>T) [Acquaviva et al., 2005; Lempp et al., 2007], and Turkish Asians (p.P615T, c.1843C>A) [Dundar et al., 2012]. These 23 mutations, reported in both homozygous and heterozygous states, are found in highly conserved residues, and their amino acid substitutions are predicted to be largely intolerated according to SIFT prediction (Table 1B). All selected mutations have been reported (Table 1A, references therein) with the exception of p.L736F, c.2206C>T, a novel mutation that exists in a compound heterozygous state with the splice site mutation c.753+2T>A [Worgan et al., 2006] in an MMA patient diagnosed in our laboratory. Propionate fixation and MUT activity assays on cells from this patient are consistent with the mut- subtype.

We mapped the missense mutations onto the human MUT crystal structure [Froese et al., 2010b] to understand their local structural environments (Supp. Fig. S1; an interactive version is available at www.thesgc.org/MUT). These mutations are located across the entire polypeptide, including the N-terminal substrate-binding domain (n = 13), C-terminal cofactor-binding domain (n = 9), and interdomain linker (n = 1) (Fig. 1A). A number of them contribute to the binding regions of the substrate (e.g., p.Y100C) and cofactor (e.g., p.M700K) (Fig. 1B), whereas four residues are buried at, or located near, the MUT dimer interface (p.P86L, p.Y316C, p.R369H, p.G426R) (Table 1B and Fig. 1B). Compatible with the large proportion of private mutations, no hot spot regions are found for the above, or other known MUT mutations.

Effects on Protein Integrity in Two Expression Systems

The diverse structural locations of the 23 missense mutations, many of which are distant from the catalytic center, suggest some mutations may impact on noncatalytic properties such as the stability of translated polypeptides. To investigate this, we reconstructed these alleles in two recombinant expression systems, E. coli and human fibroblasts (Fig. 2A and B), both of which have been used to produce MUT previously [Wil kemeyer et al., 1991; Janata et al., 1997; Froese et al., 2010b]. For E. coli expression, the mutant proteins (n = 21, excluding p.N219Y and p.Y316C that we were unable to clone) were assessed on their (1) expression level relative to wt protein, as judged by protein amount in total cell lysate, and (2) solubility, as judged by protein amount after affinity purification. We first tested several postinduction growth temperatures (37/26/18/12°C) for wt MUT and observed the highest solubility level with 18°C overnight growth (data not shown). Although all 21 mutants are expressed in E. coli at this temperature (Fig. 2A, lanes “L”), a number of them have poor solubility compared with wt, resulting in no (p.A191E and p.L328F) or reduced (p.S344F, p.F573S, p.P615T/L) protein recovery from the affinity eluant (lanes “E”). For chemical cosolvent screening, various concentrations of proline, betaine, sorbitol, TMAO, sucrose (up to 1,200 mM), and glycerol (up to 20%) were preincubated with protein before analysis.

Effects on Enzymatic Function

MUT catalyzes the isomerization of methylmalonyl-CoA to succinyl-CoA via reactive radical intermediates generated from a homolytic bond cleavage of AdoCbl [Banerjee and Ragsdale, 2003]. To investigate if and how each mutation affects MUT enzymatic function, we determined catalytic activity and ligand affinity for wt and mutant MUT expressed in MUT-deficient human fibroblasts. Under saturating AdoCbl levels (50 μM), wt MUT exhibited a mean substrate turnover of 20.2 nmol/min/mg protein (Supp. Table S2). All 23 mutants showed decreased enzyme activity,
Table 1. Selection of Mutations in the MUT Gene and Study Results

| Predicted amino acid change | Nucleotide change | Genomic location | mut subtype | References | Category of biochemical defect | Conservation score | Buried surface at dimer (%) | SIFT tolerance |
|-----------------------------|------------------|------------------|-------------|------------|-------------------------------|-------------------|-----------------------------|---------------|
| p.P66L                      | c.257C>T         | Exon 2           | mut<sup>a</sup> | [Worgan et al., 2006] | Thermolabile, K<sub>M</sub>   | 0.705             | 100                         | 0.00           |
| p.Y100C                     | c.299A>G         | Exon 2           | mut<sup>a</sup> | [Lempp et al., 2007] | Folding, catalytic            | 0.953             | 0                           | 0.00           |
| p.A191E                     | c.572C>A         | Exon 3           | mut<sup>a</sup> | [Acquaviva et al., 2005; Worgan et al., 2006] | Catalytic          | 0.936             | 0                           | 0.00           |
| p.Q211H                     | c.654A>C         | Exon 3           | mut<sup>a</sup> | [Worgan et al., 2006] | Catalytic                    | 1.000             | 0                           | 0.00           |
| p.N219F                     | c.655A>T         | Exon 3           | mut<sup>a</sup> | [Acquaviva et al., 2005; Worgan et al., 2006; Lempp et al., 2007] | Catalytic          | 1.000             | 0                           | 0.00           |
| p.Y231N                     | c.691T>A         | Exon 3           | mut<sup>a</sup> | [Worgan et al., 2006; Lempp et al., 2007] | K<sub>M</sub>       | 0.951              | 0                           | 0.00           |
| p.Y316C                     | c.947A>G         | Exon 5           | mut<sup>a</sup> | [Worgan et al., 2006] | Unclear                      | 0.817             | 97                          | 0.00           |
| p.L528F                     | c.982C>T         | Exon 5           | mut<sup>a</sup> | [Acquaviva et al., 2005; Lempp et al., 2007] | Folding, catalytic | 0.901             | 0                           | 0.01           |
| p.S344F                     | c.1031C>T        | Exon 5           | mut<sup>a</sup> | [Lempp et al., 2007] | Folding, catalytic            | 0.944             | 0                           | 0.00           |
| p.N366S                     | c.1097A>G        | Exon 6           | mut<sup>a</sup> | [Lempp et al., 2007] | Catalytic                    | 1.000             | 0                           | 0.00           |
| p.R369H                     | c.1106G>C        | Exon 6           | mut<sup>a</sup> | [Worgan et al., 2006; Lempp et al., 2007] | Catalytic          | 1.000             | 99                          | 0.00           |
| p.T387I                     | c.1160C>T        | Exon 6           | mut<sup>a</sup> | [between 2007] | Unclear                      | 1.000             | 0                           | 0.00           |
| p.G426R                     | c.1276G>A        | Exon 6           | mut<sup>a</sup> | [Worgan et al., 2006] | Thermolabile, K<sub>M</sub>   | 0.702             | 98                          | 0.00           |
| p.F573S                     | c.1718T>C        | Exon 10          | mut<sup>a</sup> | [Worgan et al., 2006] | Folding                      | 0.586             | 0                           | 0.03           |
| p.P615T                     | c.1843C>A        | Exon 11          | mut<sup>a</sup> | [Acquaviva et al., 2005; Lempp et al., 2007] | Folding            | 0.862             | 0                           | 0.00           |
| p.P615L                     | c.1844C>T        | Exon 11          | mut<sup>a</sup> | [between 2007] | Folding                      | 0.862             | 0                           | 0.00           |
| p.V633G                     | c.1898T>G        | Exon 11          | mut<sup>a</sup> | [between 2007] | K<sub>M</sub>                 | 0.794             | 0                           | 0.00           |
| p.G648D                     | c.2150G>T        | Exon 12          | mut<sup>a</sup> | [between 2007] | Thermolabile, K<sub>M</sub>   | 0.729             | 0                           | 0.00           |
| p.R694G                     | c.2080C>T        | Exon 12          | mut<sup>a</sup> | [Acquaviva et al., 2005; Lempp et al., 2007] | Catalytic          | 0.549             | 0                           | 0.00           |
| p.M700K                     | c.2081G>T        | Exon 12          | mut<sup>a</sup> | [Acquaviva et al., 2005; Lempp et al., 2007] | Thermolabile, K<sub>M</sub> | 0.539             | 0                           | 0.08           |
| p.G717V                     | c.2150G>T        | Exon 13          | mut<sup>a</sup> | [between 2007] | Mutant                        | 0.840             | 0                           | 0.00           |
| p.L736F                     | c.2206C>T        | Exon 13          | mut<sup>a</sup> | [between 2007] | Unclear                      | 0.625             | 0                           | 0.02           |

<sup>a</sup>Nucleotide numbering uses +1 as the A of the ATG translation initiation codon in the reference sequence (NM_000255.3), with the initiation codon as codon 1.

<sup>b</sup>The conservation score (0 = unconserved, 1 = strictly conserved) was calculated using Scorecons server (http://www.ebi.ac.uk/thornton-srv/databases/cgi-bin/valdar/scorecons_server, as of June 2014) [Vaalder, 2002], based on a multiple sequence alignment of 146 MUT homologues from different phyla, generated from the CONSURF server (http://consurf.tau.ac.il/, as of June 2014) [Ashkenazy et al., 2010].

<sup>c</sup>Percentage of the amino acid total surface that is buried in the human MUT dimer, calculated from the PISA server v1.48 (http://www.ebi.ac.uk/pdbe/pisat) [Krissinel and Henrick, 2007].

<sup>d</sup>SIFT tolerance score (0–0.05 corresponds to amino acid changes that likely affect protein function) was calculated using the SIFT server (http://sift.jcvi.org/, as of June 2014) [Kumar et al., 2009].

<sup>e</sup>The new variant has been submitted to dbSNP (http://www.ncbi.nlm.nih.gov/SNP/).

Effects on Protein Thermolability

To test whether the catalytic impairment of certain MUT mutants relates to changes in their structural integrity, we expressed and purified these mutants in large scale to write sufficiently soluble in E. coli (Fig. 2A, n = 15). They had similar retention volumes for AdoCbl and methylmalonyl-CoA. wt MUT had a mean K<sub>M</sub> for AdoCbl of 4.7 nM (Supp. Table S2), similar to published values [Morrow et al., 1978]. All characterized mutants exhibited higher K<sub>M</sub> values than wt, to varying degrees (Fig. 2D). Seven proteins had a K<sub>M</sub> of >200 times wt levels, of which four (p.Y231N, p.G426R, p.G648D, and p.G717V) are at least three orders of magnitude higher, indicating a strong K<sub>M</sub> defect. On the contrary, five proteins had near-wt K<sub>M</sub> (e.g., p.Y316C and p.F573S), suggesting their catalytic impairment is not primarily due to reduced cofactor affinity. No mutants showed large increases in K<sub>M</sub> for methylmalonyl-CoA (data not shown), suggesting substrate binding was not defective in these mutants, as has been shown previously [Willard and Rosenberg, 1980].
of wt MUT, with the rationale that binding of the cognate ligands should stabilize MUT thereby increasing its $T_m$. Consistent with this, wt MUT (at 1 μM) exhibits a $T_m$ of 58.2°C in the apo form, which increases in a dose-dependent manner upon addition of the native-like ligands (Supp. Fig. S3A and B), reaching $T_m$ values of 60.8°C ($\Delta T_m = 2.6°C$) with AdoCbl (50 μM), and 73.7°C ($\Delta T_m = 15.5°C$) with both AdoCbl (50 μM) and MCoA (500 μM) (Fig. 3, inset). The extent of these “$T_m$ shifts” corroborates previous structural observations [Froese et al., 2010b], in which the smaller $\Delta T_m$ by binding AdoCbl alone is consistent with a modest rearrangement in the MUT C-terminal domain (Supp. Fig. S4A), whereas the larger $\Delta T_m$ with AdoCbl + MCoA is associated with a substantial conformational change in the N-terminal domain (Supp. Fig. S4B). Notably, MCoA alone (500 μM) did not increase the $T_m$ of MUT beyond apo levels (data not shown), suggesting an ordered binding mechanism of AdoCbl cofactor first, followed by substrate, in the ternary complex.

We subsequently measured the $T_m$ values of purified mutant proteins in the apo, AdoCbl bound, and AdoCbl + MCoA bound states (Fig. 3), to determine if they had a lower $T_m$ than wt, an indicator of protein destabilization [Leandro et al., 2011]. For clarity, the $\Delta T_m$ of apo mutants compared to apo-wt is given in Figure 2E. Six of 15 proteins exhibited near-wt $T_m$ in their apo forms, indicating intact protein integrity. Of these, two (p.F573S, p.V633G) responded to AdoCbl + MCoA stabilization with similar $T_m$ shifts as wt, whereas four (p.Y100C, p.N366S, p.R369H, p.T387I) did not, suggesting that the latter mutants lost the capacity for native ligand stabilization. In particular, p.T387I showed a concentration-dependent decrease in $T_m$ when adding AdoCbl alone (Supp. Fig. S3C). The other nine proteins, however, were more thermolabile than wt in their apo forms. Although five (p.Q218H, p.Y231N, p.G648D, p.R694W, and p.M700K) had slightly lower $T_m$ than wt ($\Delta T_m$ range -2.7 to -1.3) and responded to AdoCbl + MCoA stabilization, four (p.P86L, p.G426R, p.R694L, and p.G717V) were substantially more thermolabile than wt (maximum destabilization: p.P86L, $\Delta T_m$ = -8.7°C) and did not respond to ligand supplementation.

### Stabilization of MUT Mutants by Osmolytes

The insolubility and thermolability of a number of MUT mutants, compared with wt, implicate protein destabilization as a biochemical defect and prompted us to investigate the potential for posttranslational stabilization of mutant proteins by six different osmolytes using DSC and bacterial expression (Fig. 4; Supp. Figs. S5 and S6). When applied in high millimolar concentrations, these uncharged, low-molecular-weight compounds function as “chemical chaperones” and stabilize a number of disease-associated proteins prone to misfolding, an effect likely mediated by osmotic changes in the milieu [Nascimento et al., 2008; Majtan et al., 2010]. The two most thermolabile MUT mutants in our study, p.P86L and p.G426R, showed that $T_m$ increases at high osmolyte concentrations (10% glycerol, 600 mM others), with the efficacy of stabilization dependent on the osmolyte used (Fig. 4A and B). These osmolytes confer a nonsaturating dose response, and the effect appears nonsselective, as wt and all mutant proteins tested demonstrated stabilization in their apo forms to similar degrees (Supp. Fig. S5). To determine if osmolytes could also stabilize proteins cotranslationally, we expressed several mutants with varying levels of solubility in *E. coli* and supplemented their growth (Supp. Fig. S6). Compared with no supplement control, soluble protein yield was improved with glycerol (at 1% concentration: p.A191E and p.G426R; 5%: p.R694W), TMAO (100 mM: p.P615T and p.R694W), and to a lesser extent betaine (100 mM: p.R694W). No improvement for p.L328F was seen,
Figure 2. MUT missense mutations confer different effects on stability and activity. A: Coomassie staining of SDS-PAGE following small-scale bacterial expression and affinity purification. For each mutation, lanes for total cell lysate (“L,” left) and eluant after purification (“E,” right) are shown. B: Western blot following expression of each mutation in a MUT-deficient patient cell line. Vector, empty vector. C: Total MUT activity (assay with 50 μM AdoCbl) of each mutation expressed as percent of mean wt activity. D: $K_M$ for AdoCbl, expressed as times the mean wt value. E: $\Delta T_m$ of apo mutant compared to apo-wt protein (see Fig. 3). –, not applicable.

whereas wt protein was very soluble under all conditions, suggesting the rescue could be mutant specific. Together, our data demonstrate the amenability of small exogenous molecules to modulate MUT stability, and support the notion to develop target-specific “pharmacological chaperones” (PCs).

**Discussion**

Our study aimed to establish a genotype–phenotype correlation by characterizing the biochemistry associated with a set of MUT missense mutations. Earlier small-scale studies have tested the activity and expression of MUT misense mutants, for example, using the lower eukaryote Saccharomyces cerevisiae (five mutations) (Crane and Ledley, 1994) and E. coli (four mutations) [Janata et al., 1997] as recombinant hosts. Our selected set of 23 mutants, five of which overlap with those previous studies (p.Y231N, p.R369H, p.G648D, p.R694W, p.G717V), constitutes to our knowledge the largest biochemical analysis of MUT mutations to date.

**A Catalogue of Biochemical Defects in MUT Mutants**

Traditionally, MMA phenotypes are broadly classified into $\text{mut}^0$ and $\text{mut}^-$ subtypes based on the response of a patient cell line to hydroxocobalamin in propionate incorporation [Lempp et al., 2007]. In order to understand the molecular mechanism(s) governing the $\text{mut}^0/\text{mut}^-$ subtypes and catalogue the associated biochemical defects, we have further assessed MUT mutations based on stability and catalytic properties as summarized in Table 1B.

We evaluated mutants on two stability criteria, folding and thermolability. Folding defects refer to the aggregation or degradation of translated polypeptides, thereby reducing the steady-state protein. These mutants displayed drastically reduced soluble
Figure 3. Thermolability of MUT mutations. The change in $T_m$ values ($\Delta T_m$, compared with apo wt) for each mutant MUT in the apo state, or following the addition of AdoCbl alone or with AdoCbl and MCoA, is shown. Black, apo protein; red, with 50 $\mu$M AdoCbl; blue, with 50 $\mu$M AdoCbl and 500 $\mu$M MCoA. Error bars depict SEM from at least three measurements. (Inset) Representative DSF melting curves of wt MUT in the apo form (black), with 50 $\mu$M AdoCbl (red) and with 50 $\mu$M AdoCbl and 500 $\mu$M MCoA (blue).

Figure 4. Stabilization of mutant MUT by osmolytes. Representative DSF melting curves for p.P86L (A) and p.G426R (B) mutants in the apo form (gray) and in the presence of osmolytes (colors) are shown along with wt MUT (black).

protein yield, as exemplified by p.P615T/L. Four other mutants, two from each mut subtype (mut$: p.A191E, p.L328F; mut$: p.S344F, p.F573S), also fit this category having reduced detectable protein in one of the two expression systems. A few mutants displayed only slightly decreased soluble E. coli expression, such as p.G717V and p.R694W, consistent with a previous report [Crane and Ledley, 1994] and were not included in the folding category. Thermolabile mutants had near-wt levels of recombinant expression and solubility, reflecting near-native folding, but increased temperature sensitivity. Four mutations in this study fell into the thermolabile category (p.P86L, p.G426R, p.R694L, p.G717V).

Inspection of the MUT crystal structure shows that both folding and thermolabile mutants involve amino acid changes that (1) remove a conformationally restricted proline residue (e.g., p.P86L and p.P615L/T), (2) cause steric clashes with a bulky side chain (e.g., p.L328F and p.S344F), or (3) disrupt the natively charged/polar (e.g., p.R694L) or nonpolar (e.g., p.A191E) environment. Furthermore, stability mutations are distributed across the entire polypeptide, indicating that all domains of the protein contribute to its intrinsic stability.

In addition to stability mutants, we also identified direct kinetic impairment, which we termed catalytic and $K_M$ mutants. Catalytic mutants had low residual activity (<5% of wt) despite detectable eukaryotic expression and constituted the largest category of mutants (10 in this study, Table 1B). Of particular note, two catalytic mutants, p.Q218H and p.A219E, involve structurally adjacent residues lining the substrate-binding channel (Fig. 1A; Supp. Fig. S1D and E), where p.Q218 has been implicated in transition state stabilization [Loferer et al., 2003]. $K_M$ mutants exhibited >200-fold increased $K_M$ for AdoCbl compared with wt. Among the seven $K_M$ mutants
identified (Table 1B), only three (p.V633G, p.G648D, and p.G717V) are buried within the C-terminal cobalamin-binding domain (Fig. 1A; Supp. Fig. S1,F, Q, and T). The other four are toward the N-terminal end: p.Y231N is situated at the interdomain interface, within a loop that contacts the cobalamin corrin ring (Supp. Fig. S1F); p.G426R is situated at the dimeric interface; and p.P86L and p.Y100C are located near the N-terminal substrate-binding channel, distant from any obvious involvement in cobalamin binding. Further investigation is therefore warranted to understand how the Kd effect of these mutations is mediated.

It is of note that our classification scheme is not aimed at pigeonholing mutant alleles into disparate, mutually exclusive categories; on the contrary, there is a high degree of interdependence between various stability and catalytic properties that together contribute to proper enzyme function. This is clearly exemplified with seven mutants fitting multiple criteria. Three of them had a significantly increased Kd but were also thermolabile (p.P86L, p.G426R, and p.G717V), one was thermolabile with a catalytic defect (p.R694L), and three met the criteria of folding and catalytic defects (p.A191E, p.L528F, and p.S344F). In addition, three mutations were termed “unclear” (p.Y316C, p.T387I, and p.L736F) because in our assays they lack definitive aberration in stability and enzyme kinetics, showed detectable protein by Western blot, intermediate enzyme activity, and only moderate Km change.

Future studies should aim to address the lesser characterized biochemical aspects of MUT and possible mutational effects, which are beyond the scope of this study. These include (1) possible allosteries between substrate and cofactor binding, which could potentially explain the Tm shift from MCoA + AdoCbl but not MCoA alone; (2) potential formation of a supramolecular complex with the GT-Pase MMAA and adenosyltransferase MMAB [Froese and Gravel, 2010], required for the proper assembly of AdoCbl onto the MUT enzyme; and (3) interallelic rescue of certain MUT disease alleles (e.g., p.R93H and p.G717V) [Worgan et al., 2006].

Clinical Utility and Implications of In Vitro Characterization

A clinical application for our biochemical categorization of MUT genotypes is to help prognosticate the course of the disease. As expected, the characterized mut+ mutants harbor folding or catalytic defects, resulting in very low or undetectable residual enzyme activities consistent with their predominance in neonatal onset patients with severe long-term complications. On the contrary, the characterized mut- mutants can present in any one, or combinations, of the four biochemical defects defined here. We postulate that mutations that do not interfere with protein stability and have less of a catalytic penalty would result in milder disease (e.g., p.P86L or p.G426R). Similar to other metabolic diseases where an activity threshold of ~5%–10% of wt is sufficient to ameliorate disease [Leinekugel et al., 1992; Suzuki, 2013], mut+ patients often show a later onset and less long-term complications, presumably due to the intermediate level of residual MUT activity.

Whether any MUT-deficient patient, including those of the mut- subtype, benefit from treatment with high doses of hydroxocobalamin is controversial. Early publications documented no response [Wilcken et al., 1977; Matsui et al., 1983] and a recent documentation of a possible response has relied on retrospective reviews [Horster et al., 2007], whereas up to now, no controlled studies clearly demonstrating response to hydroxocobalamin have been published. This picture is very different from PKU, where >40% patients respond clinically to the cognate cofactor tetrahydropterin (BH4) of phenylalanine hydroxylase (PAH) [Zurfluh et al., 2008]. For mut-type MMA, a clinical response to treatment is usually defined as reduced plasma or urinary concentration of methylmalonic acid. Although difficult to measure because of its broad variation in body fluids [Thompson and Chalmers, 1990; Horster et al., 2007; Fowler et al., 2008], and clinical improvement even more difficult to determine reliably, lower methylmalonic acid levels are thought to reflect a higher residual enzyme activity protecting patients from severe long-term complications (e.g., chronic renal failure and neurological complications) [Thompson and Chalmers, 1990; Horster et al., 2007; Cosson et al., 2009]. Therefore, careful evaluation of potential responsiveness to hydroxocobalamin treatment in all MMA patients is warranted, and a protocol has been proposed [Fowler et al., 2008]. Our thermal denaturation data offer a molecular explanation to a possible mechanism of cobalamin treatment by revealing AdoCbl-induced stability improvement in certain mutants in a ligand-specific manner similar to that of BH4 on mutant PAH. Our results also predict that patients harboring a Kd defect (e.g., p.G648D) are more likely to benefit from cobalamin treatment based on the assumption that increased hydroxocobalamin plasma levels result in increased cellular AdoCbl production because, in our enzyme assay, these mutants exhibited high activity levels in the presence of high AdoCbl concentration. Nevertheless, hydroxocobalamin treatment may only benefit a small subset of patients with intermediate to high residual activity, if any, outlining an imperative to search for alternative therapies.

Small Molecule Chaperones as Alternative Therapy?

Our stability data add MUT to the growing list of metabolic enzymes in which mutation-induced protein destabilization is widely believed to play a causative role in disease pathogenesis [Gregersen et al., 2006]. In this model, mutant polypeptides tend to misfold, aggregate, and be degraded by protein quality control in the cell. Chemical or pharmacologic approaches to partially correct misfolding, divert the mutant polypeptide from degradation/aggregation pathways, and deliver it to the native subcellular destination may therefore allow a sufficient recovery of physiological function. This small molecule “chaperoning” approach has been proposed as a viable treatment for a number of protein misfolding diseases [Pey et al., 2008; Gomes, 2012]. We provide proof of principle for MUT chaperone therapy by showing in vitro and in cell stabilization of several mutant proteins using chemical osmolytes. These small organic solutes enhance protein stability via a nonselective, nontarget-specific mode of preferential exclusion from the protein’s surface [Arakawa and Timasheff, 1985], although necessitating high effective concentrations (in millimolar range) that may be toxic to the cell—restricting their therapeutic utility.

Alternatively, target-specific PCs could be developed for MUT that, unlike chemical chaperones, exert a direct stabilization effect by binding to the affected protein, potentially allowing lower, clinically tolerable dosage concentrations. PCs have shown promise in several lysosomal storage diseases, entering early-stage clinical trials for Gaucher, Tay-Sachs, and Fabry diseases [Boyd et al., 2013]. In the case of MUT, we anticipate that an effective PC molecule need only to stabilize the mutant protein “long enough” to be delivered to mitochondria, where it can be further stabilized by its native ligands and protein interaction partners (MMAA and MMAB). Such a PC molecule may increase the residual enzyme activity of the mutant protein to a sufficient level that copes with the cellular demands of propionate catabolism. Therefore, PC therapy is likely to rescue mild stability defects (i.e., alleles that give rise to destabilized protein and partial loss of function; e.g., p.G426R, p.M700K, and p.G717V),
either to be administered alone or as a complement to the existing diet/hydroxocobalamin treatment. Other catalytic, Km mutations, without stability defects, may also benefit from PCs that potentially increase residual activity, acting as "enzymatic activators." Altogether, the stage is set for a systematic, unbiased screening regime in the search for PC molecules targeting MUT, and our established DSF assay may provide a good starting point for this screening.

Acknowledgments

The Structural Genomics Consortium is a registered charity (number 1097737) that receives funds from AbbVie, Boehringer Ingelheim, the Canada Foundation for Innovation, the Canadian Institutes for Health Research, Genome Canada, GlaxoSmithKline, Janssen, Lilly Canada, the Novartis Research Foundation, the Ontario Ministry of Economic Development and Innovation, Pfizer, Takeda, and the Welcome Trust [092809/Z/10/Z].

References

Acquaviva C, Benoist JP, Pereira S, Cellebaut I, Koskas T, Poquet D, Elion J. 2005. Molecular basis of methylmalonyl-CoA mutase phenotype diversity in 40 European patients affected by mut(o) and mut-forms of methylmalonic acidemia: identification of 29 novel mutations in the MUT gene. Hum Mutat 25:167–76.

Arakawa T, Timashoff SN. 1985. The stabilization of proteins by osmolytes. Biophys J 47:411–414.

Ashkenazy H, Erez E, Martz E, Pupko T, Ben-Tal N. 2010. ConSurf 2010: calculating evolutionary conservation in sequence and structure of proteins and nucleic acids. Nucleic Acids Res 38:W529–W533.

Banerjee R, Ragsdale SW. 2003. The many faces of vitamin B12: catalysis by cobalamin-dependent enzymes. Annu Rev Biochem 72:209–247.

Baumgartner R. 1983. Activity of the cobalamin-dependent methylmalonyl-CoA mutase. In: Hall CA, editor. The cobalamins – volume 10 of methods in hematology. Churchill Livingstone, London, UK, p 181–193.

Baumgartner MR, Almanzah S, Suormala T, Obie C, Cole RN, Packman S, Baumgartner ER, Valle D. 2001. The molecular basis of human 3-methylcrotonyl-CoA carboxylase deficiency. J Clin Invest 107:495–504.

Boyd RE, Lee G, Rybczynski P, Benjamin ER, Khanna R, Wustman BA, Valenzano KJ. 2011. Understanding carbamoyl-phosphate synthetase I (CPS1) deficiency by using expression studies and structure-based analysis. Hum Mutat 31:801–810.

Horster F, Baumgartner MR, Viardot C, Suormala T, Burgard P, Fowler B, Hofmann GF, Garbade SF, Kolker S, Baumgartner ER. 2007. Long-term outcome in methylmalonic acidurias is influenced by the underlying defect (mut0, mut-, cblA, cblB), variants on protein function using the SIFT algorithm. Nat Protoc 3:1073–1082.

Janata J, Kogekar N, Fenton WA. 1997. Expression and kinetic characterization of methylmalonyl-CoA mutase from patients with the mut- phenotype: evidence for naturally occurring interallelic complementation. Hum Mol Genet 6:1457–1464.

Krisnissen E, Henrick K. 2007. Inference of macromolecular assemblies from crystalline state. J Mol Biol 372:774–797.

Kumar P, Henikoff S, Ng PC. 2009. Predicting the effects of coding non-synonymous variants on protein function using the SIFT algorithm. Nat Protoc 4:1073–1082.

Ledley FD, Muneta M, Nguyen PN, Kolhouse JF, Allen RH. 1988A. Molecular cloning of L-methylmalonyl-CoA mutase: gene transfer and analysis of mut cell lines. Proc Natl Acad Sci USA 85:3518–3521.

Ledley FD, Muneta MR, Zoghbi HY, Vantsunen P, Ledbetter SA, Ledbetter DH. 1988B. Mapping of human methylmalonyl-CoA mutase (Mut) locus on chromosome 6. Am J Hum Genet 42:839–846.

Ledley FD, Rosenblatt DS. 1997. Mutations in mut methylmalonic acidemia: clinical and enzymatic correlations. Hum Mutat 9:1–6.

Majtan T, Liu L, Carpenter JP, Kraus JP. 2010. Rescue of cystathionine beta-synthase (CBS) mutants with chemical chaperones: purification and characterization of eight CBS mutant protein enzymes. J Biol Chem 285:15866–15873.

Niesen FH, Campilo I, Ainscow PM, Sandhoff K. 1992. Quantitative correlation between the residual activity of beta-lysosomal aminidase-a and a-lysosomal a-aminidase-a and the severity of the resulting lysosomal storage disease. Hum Genet 88:513–523.

Pekkala S, Martinez AI, Barcelona B, Yefimenko I, Finckh U, Rubio V, Cervera J. 2010. Identification of pharmacological chaperones as potential therapy for phenylalanine hydroxylase deficiency. Hum Mutat 31:2212–2221.

Pendse S, Martinez Al, Barcelona B, Yefimenko I, Finckh U, Rubio V, Cervera J. 2010. Phenylalanine hydroxylase deficiency. Gene Med 31:808–817.

Pey AL, Stricher F, Serrano L, Martinez A. 2007. Predicted effects of missense mutations on native-state stability accounted for by free energy of binding in the protein environment. J Mol Biol 372:562–572.

Rohrer ME, Remmers JE, Krysztofinska E, Wang X, Kroon E, Ugochukwu E, Gravel RA. 2009. Predicting the effects of coding non-synonymous variants on protein function using the SIFT algorithm. Nat Protoc 4:1073–1082.

Rohrer ME, Remmers JE, Krysztofinska E, Wang X, Kroon E, Ugochukwu E, Gravel RA. 2009. Predicting the effects of coding non-synonymous variants on protein function using the SIFT algorithm. Nat Protoc 4:1073–1082.

Rohrer ME, Remmers JE, Krysztofinska E, Wang X, Kroon E, Ugochukwu E, Gravel RA. 2009. Predicting the effects of coding non-synonymous variants on protein function using the SIFT algorithm. Nat Protoc 4:1073–1082.

Rohrer ME, Remmers JE, Krysztofinska E, Wang X, Kroon E, Ugochukwu E, Gravel RA. 2009. Predicting the effects of coding non-synonymous variants on protein function using the SIFT algorithm. Nat Protoc 4:1073–1082.

Rohrer ME, Remmers JE, Krysztofinska E, Wang X, Kroon E, Ugochukwu E, Gravel RA. 2009. Predicting the effects of coding non-synonymous variants on protein function using the SIFT algorithm. Nat Protoc 4:1073–1082.

Rohrer ME, Remmers JE, Krysztofinska E, Wang X, Kroon E, Ugochukwu E, Gravel RA. 2009. Predicting the effects of coding non-synonymous variants on protein function using the SIFT algorithm. Nat Protoc 4:1073–1082.

Rohrer ME, Remmers JE, Krysztofinska E, Wang X, Kroon E, Ugochukwu E, Gravel RA. 2009. Predicting the effects of coding non-synonymous variants on protein function using the SIFT algorithm. Nat Protoc 4:1073–1082.

Rohrer ME, Remmers JE, Krysztofinska E, Wang X, Kroon E, Ugochukwu E, Gravel RA. 2009. Predicting the effects of coding non-synonymous variants on protein function using the SIFT algorithm. Nat Protoc 4:1073–1082.

Rohrer ME, Remmers JE, Krysztofinska E, Wang X, Kroon E, Ugochukwu E, Gravel RA. 2009. Predicting the effects of coding non-synonymous variants on protein function using the SIFT algorithm. Nat Protoc 4:1073–1082.
Suzuki Y. 2013. Chaperone therapy update: Fabry disease, G(M1)-gangliosidosis and Gaucher disease. Brain Dev-Jpn 35:515–523.

Thompson GN, Chalmers RA. 1990. Increased urinary metabolite excretion during fasting in disorders of propionate metabolism. Pediatr Res 27:413–416.

Valdar WS. 2002. Scoring residue conservation. Proteins 48:227–241.

Vedadi M, Niesen FH, Allali-Hassani A, Fedorov OY, Finerty PJ, Jr., Wawney GA, Yeung B, Arrowsmith C, Ball LI, Berglund H, Hui R, Maresden BD et al. 2006. Chemical screening methods to identify ligands that promote protein stability, protein crystallization, and structure determination. Proc Natl Acad Sci USA 103:15835–15840.

Wilcken B, Kilham HA, Faull K. 1977. Methylmalonic aciduria: a variant form of methylmalonyl coenzyme A apomutase deficiency. J Pediatr 91:428–430.

Wilkenmeyer MF, Crane AM, Ledley FD. 1991. Differential diagnosis of mut and cbl methylmalonic acidemia by DNA-mediated gene transfer in primary fibroblasts. J Clin Invest 87:915–918.

Wilkemeyer MF, Crane AM, Ledley FD. 1991. Differential diagnosis of mut and cbl methylmalonic aciduria by DNA-mediated gene transfer in primary fibroblasts. J Clin Invest 87:915–918.

Willard HF, Mellman IS, Rosenberg LE. 1978. Genetic complementation among inherited deficiencies of methylmalonyl-CoA mutase activity – evidence for a new class of human cobalamin mutant. Am J Hum Genet 30:1–13.

Willard HF, Rosenberg LE. 1980. Inherited methylmalonyl CoA mutase apoenzyme deficiency in human fibroblasts: evidence for allelic heterogeneity, genetic compounds, and codominant expression. J Clin Invest 65:690–698.

Worgan LC, Niles K, Tirone JC, Hofmann A, Verner A, Sammak A, Kucic T, Lepage P, Rosenblatt DS. 2006. Spectrum of mutations in mut methylmalonic acidemia and identification of a common Hispanic mutation and haplotype. Hum Mutat 27:31–43.

Yue WW, Froese DS, Brennan PE. 2014. The role of protein structural analysis in the next generation sequencing era. Top Curr Chem 336:67–98.

Zurfluh MR, Zschocke J, Lindor M, Feilet F, Chery C, Burzina A, Stevens RC, Thony B, Blau N. 2008. Molecular genetics of tetrahydrobiopterin-responsive phenylalanine hydroxylase deficiency. Hum Mutat 29:167–175.