Novel Mouse Models of Methylmalonic Aciduria
Recapitulate Phenotypic Traits with a Genetic Dosage Effect

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Running title: Phenotype and treatment study in MMAuria mouse model

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ABSTRACT

Methylmalonic aciduria (MMAuria), caused by deficiency of methylmalonyl-CoA mutase (MUT), usually presents in the newborn period with failure to thrive and metabolic crisis leading to coma or even death. Survivors remain at risk of metabolic decompensations and severe long-term complications, notably renal failure and neurological impairment. We generated clinically relevant mouse models of MMAuria using a constitutive Mut knock-in (KI) allele based on the p.Met700Lys patient mutation, used homozygously (KI/KI) or combined with a KO allele (KO/KI), to study biochemical and clinical MMAuria disease aspects. Transgenic Mut

ki/ki and Mut

ko/ki mice survive postweaning, show failure to thrive, increased methylmalonic acid, propionylcarnitine, odd-chain fatty acids and sphingoid bases - a new potential biomarker of MMAuria. Consistent with genetic dosage, Mut

ko/ki mice have lower Mut activity, are smaller and show higher metabolite levels than Mut

ki/ki mice. Further, Mut

ko/ki mice exhibit manifestations of kidney and brain damage, including increased plasma urea, impaired diuresis, elevated biomarkers and changes in brain weight. On a high protein diet, mice display disease exacerbation, including elevated blood ammonia, and catastrophic weight loss, which, in Mut

ki/ki mice, is rescued by hydroxocobalamin treatment. This study expands knowledge of MMAuria, introduces the discovery of new biomarkers and constitutes the first in vivo proof-of-principle of cobalamin treatment in mut-type MMAuria.

Methylmalonyl-CoA mutase (MUT) is a homodimeric enzyme that catalyzes the reversible isomerization of L-methylmalonyl-CoA to succinyl-CoA by using vitamin B12 (cobalamin; Cbl) in the cofactor form (5'-deoxyadenosylcobalamin; AdoCbl). In this anaplerotic reaction, several metabolic pathways, including the breakdown of branched chain amino acids, odd-chain fatty acids, and the side chain of cholesterol, converge on the propionate pathway. The vital importance of this enzyme is demonstrated by the severe inborn error of metabolism methylmalonic aciduria (MMAuria), which is caused by mutations in the MUT gene (mut-type MMAuria, OMIM #251000) or by defects of AdoCbl synthesis. Dysfunction of MUT leads to an accumulation of methylmalonic acid (MMA), 2-methylcitrate (2-MC), propionylcarnitine (C3) and other metabolites in body fluids and tissues (1,2). Although the clinical symptoms observed in mut-type MMAuria patients are variable, they often present in the newborn period with ketoacidosis, lethargy, failure to thrive, leading to coma or even death if untreated. In surviving patients, the chronic course involves organ-specific complications, the most common being neurological impairment and chronic kidney disease, manifesting as tubulo-interstitial nephritis and renal tubular acidosis. End-stage renal disease, occurring as early as the second decade of life in patients with severe MMAuria, is frequently seen (3-6). Besides symptomatic care, there are a few rational treatment approaches, including reduction of protein intake, carnitine and Cbl supplementation in order to decrease the flux through the propionate pathway and the formation of toxic metabolites (6). Although this treatment appears to partly improve metabolic control in some cases, most patients still suffer severe long-term complications.

The first effort to generate a mouse model of mut-type MMAuria was made in 2003, when a Mut knockout (KO) model on C57BL/6 background resulted in neonatal lethality of all homozygous (KO/KO) pups (7). Later, Peters et al. expressed varying copy numbers of the intact human MUT locus on the KO background and demonstrated a copy number dependent rescue of the KO lethality (8). They also developed another transgenic model containing an introduced stop codon on the human MUT locus crossed to the KO background resulting in a severe MMAuria phenotype (9). In 2009, Chandler et al. generated a Mut-KO mouse on the modified [(C57BL/6X129Sv/Ev) X FVB/N] background resulting in some KO mice surviving the neonatal period (10), although nearly all died within 25 days (11). Using this KO model, they applied adeno-associated virus-mediated gene therapy (11,12) and stable transgenic Mut expression restricted to the liver (13) to provide a long-term rescue of lethality. However, while these have proven to be useful models to study some aspects and rescue of disease, no model to date has been able to provide an accurate depiction of the patient situation. Neonatal or early lethality hamper long-term follow-up studies and the understanding of chronic disease manifestation, while transgenic rescue of Mut does not resemble the pathologic situation, where a patient carries the same genetic defect in all cells.
In order to devise a model of MMAuria reflective of the patient situation using a known human mutation and amenable to the study of the long-term disease course, we opted to knock-in (KI) a hypomorphlic allele based on a patient missense mutation. From our previous in depth MUT missense mutation characterization (14), we identified p.Met700Lys (c.2099T>A) human cDNA; p.Met698Lys in mouse) as an ideal candidate for this allele due to the residual enzymatic activity and in vitro response to hydroxocobalamin (OHCbl) found in MUT protein carrying this mutation. p.Met700Lys has been detected heterozygously in 4 MMAuria patients in conjunction with severe mutations (15,16), but at least 3 of these patients have had an intermediate disease course with a relatively later age of onset (between 14 days and 6 months) (16), supporting the potential of this mutation to cause moderate MMAuria. To assess the effect of genetic dosage on the phenotype and to generate a more severe disease model, the new KI mutation was combined with a KO allele (7). Here we present more severe disease model, the new KI mutation genetic dosage on the phenotype and to generate a cause moderate MMAuria. To assess the effect of (16), supporting the potential of this mutation to had an intermediate disease course with a relatively almost 100-fold increased

man fibroblasts carrying a homozygous null allele 

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of genetic dosage on the phenotype and to generate
a more severe disease model, the new KI mutation
was combined with a KO allele (7). Here we present
a thorough characterization of these newly gener-
tated Mut

ki/ki and Mut

ko/ki mouse models, which
survive long-term and at the same time recapitulate
the key biochemical and clinical features of MMAuria.

RESULTS

In vitro characterization of the p.Met698Lys knock-in allele – To characterize the selected KI mutation, we tested the biochemical properties of the human mutation p.Met700Lys (hs-
p.Met700Lys) and the equivalent mutation in mouse (mm-p.Met698Lys). Separate constructs incorpora-
ing each allele were overexpressed in human fibroblasts carrying a homozygous null allele of MUT in order to determine enzyme activity and kinetics. In both, the residual activity was reduced to a few percent of wild-type (wt) (Table 1) with an almost 100-fold increased Km for AdoCbl (Table 1). These nearly identical biochemical results for the human and mouse equivalent mutations led us to conclude that the mm-p.Met698Lys mutation would be expected to produce an intermediate phenotype of MMAuria, similar to its hs-p.Met700Lys counterpart.

Generation of KI allele and initial charac-
terization – To generate the knock-in p.Met698Lys mutation, a targeting vector was designed to replace thymine with adenine at position c.2093 of Mut (based on NM_008650.3). The construct (Fig. 1A) was used for embryonic stem cell (C57BL/6-derived) targeting and a positive clone was confirmed by Southern hybridization (Fig. 1B) which was used for blastocyst injection. Screening for positive germ-line transmission was done by PCR genotyping (Fig. 1C, upper panel). Additionally, Mut

ko/ki mice were generated as a second disease model by crossing the previously established Mut

ko/ko mice (7) with our Mut

ki/ki mice, and followed by PCR genotyping of the KO allele (Fig. 1C, lower panel).

To determine the direct consequences of the missense change on Mut transcript, stability and function, we investigated Mut expression and activity in various tissues. In wt mice, we found Mut expression to be lower in brain than in liver or kidney (Fig. 1D), similar to previously published results (17). The KI allele did not result in an appreciable decrease of Mut transcript levels in all tissues tested, however, mice with one KO allele had ~50% of Mut transcript levels compared to wt (Fig. 1E), likely indicative of nonsense mediated decay. At the protein level, the amount of the homozygous Mut

ki/ki protein was significantly reduced compared to controls, suggesting an unstable mutant protein, while KO/KI Mut protein showed similar reduction (Fig. 1F). Mut activity from wt mice significantly varied among tissue types, with highest activities in kidney, liver and heart (Fig. 1G). Activities from KI/KI and KO/KI mice were greatly decreased in all tissues tested (Fig. 1H), comparable to the in vitro measurements (Table 1). Furthermore, a gene dosage-dependent decrease of enzyme activity was identified, with Mut

ko/ki activity slightly lower than Mut

ki/ki in most tissues (Fig. 1H). As expected, ~50% percent of residual activity was maintained in the Mut

ki/ko and Mut

ko/ko animals, reflecting the activity of the remaining wt allele (Fig. 1H).

Gene dosage-dependent biochemical and clinical phenotype in Mut

ki/ki and Mut

ko/ko mice – Mice homozygous for the targeted point mutation (Mut

ki/ki) were viable and, when crossed with heterozygous Mut

ki/ko mice, reproduced at the expected Mendelian ratio of ~50% percent per genotype (51% Mut

ki/ko, 49% Mut

ki/ki litters = 33). Breeding of Mut

ko/ko also resulted in viable animals and displayed the expected Mendelian distribution of the genotypes (51% Mut

ko/ko, 49% Mut

ki/ko, litters = 36). Mice of both genotypes survived over one year, allowing us to monitor long-term development. At early stages of life, Mut

ki/ki and Mut

ko/ko mice were largely indistinguishable from their heterozygous littermates (Mut

ki/ko) and wt control mice. However,
after the age of about 100 days in females and about 150 days in males (Fig. 2A), mice of both genotypes showed signs of significant growth retardation. The difference in body weight between littermate control (Mut\(^{ki/wt}\)) and Mut\(^{ko/ki}\) female mice was 30% after approximately one year (Fig. 2A), while food intake remained constant (e.g., relative food intake at the age of 218 days in Mut\(^{ki/wt}\) was 0.079 grams of chow normalized to body weight in grams, S.D. ± 0.043 and in Mut\(^{ko/ki}\) 0.078 ± 0.048). In this cohort, C3 (normalized to acetylcarnitine, C2) in blood and MMA in urine were constantly elevated over the entire time span investigated (Fig. 2B and C). Survival was the same in all three groups (all mice still alive after one year except for one Mut\(^{ki/ki}\) and one Mut\(^{ko/ki}\), both of which died during blood collection).

To gain further insight into the consequences of the clearly reduced Mut enzymatic activity, we performed a cross-sectional analysis of hallmark metabolites of MMAuria (18) at 35 days of age. MMA was found elevated in a genetic dosage-dependent manner in tissues, urine and dried blood, with the highest levels found in Mut\(^{ko/ki}\) mice (Fig. 2D). Although Mut\(^{ki/ki}\) animals showed considerably lower levels, they were still significantly elevated compared to controls (Fig. 2D). C3 levels (normalized to C2) showed the same trend (Fig. 2E). Mut\(^{ki/wt}\) and Mut\(^{ko/wt}\) did not show any elevations, suggesting that one functional \(w\) allele is sufficient to cope with the demands of the propionate pathway (Fig. 2D and E). Ammonia levels, which are elevated during metabolic crisis in patients, were normal (Fig. 2F), suggesting these animals did not suffer from acute crises on normal chow. Similar to patients (19-21), both Mut\(^{ko/ki}\) and Mut\(^{ki/ki}\) showed elevated glycine levels (Fig. 2G), while other amino acid levels were mostly unchanged.

In addition to classic metabolites, we performed a comprehensive analysis of fatty acids and sphingoid bases. We found increased levels of odd-chain fatty acids, including the 17-carbon chain length fatty acid, which has recently been proposed as a biomarker of MMAuria (22,23), with normal levels of even-chain fatty acids (Fig. 2H). Further, we show here for the first time an elevation of odd-chain length sphingoid bases in plasma (Fig. 2I) and tissues (Fig. 2J), possibly as a consequence of the increased odd-chain fatty acids, while levels of even-chain sphingoid bases remained normal (Fig. 2J). Specific fatty acid and sphingoid base levels are freely available upon request.

**Initial characterization of the renal and neurological phenotypes** – We investigated whether Mut\(^{ko/ki}\) mice suffer from similar long-term complications as patients with MMAuria, e.g. renal dysfunction. When normalized to body weight, Mut\(^{ko/ki}\) mice consistently produced less urine during adulthood than littermate controls (Mut\(^{ki/wt}\)) (Fig. 3A), although water intake was not significantly different between the two groups (Fig. 3B). Consistent with renal dysfunction, these mice also had increased plasma urea throughout the time course measured (Fig. 3C), which has been observed in the context of a decreased glomerular filtration rate (24). Renal tubular dysfunction in these animals was evidenced by disturbed excretion of several electrolytes in urine (Fig. 3D). Kidney damage was further supported by increased levels of the biomarker Lipocalin-2 (Lcn2) in kidney tissue (Fig. 3E). Of note, light microscopy examination of hematoxylin-eosin stained sections of kidneys from Mut\(^{ki/wt}\) and Mut\(^{ko/ki}\) mice (female, 1-year old) did not reveal any structural abnormalities (Fig. 4A).

Neurological dysfunction is the other major long-term complication of MMAuria (25,26), often manifesting as movement disorder due to lesions in the basal ganglia and in acute metabolic crisis brain edema may be observed. In the Mut\(^{ko/ki}\) mice we observed an increase in brain weight at the age of one month as well as one year (Fig. 3F), which may be suggestive of brain edema. Although brain histology was normal in the basal ganglia (Fig. 4B), we also identified an upregulation in Lcn2 at the mRNA and protein level (Fig. 3G).

Together, the initial characterization of both the brain and kidney, in combination with the metabolites already identified in these organs, suggest an initial dysfunction in each, which has not yet manifested at the gross level.

**Diet-induced metabolic decompensation** – An increased throughput of the propionate pathway in patients with MMAuria, e.g. due to a catabolic state or an augmented protein intake, often results in metabolic decompensation (crisis). In order to replicate this state in our MMAuria mouse models and exacerbate their clinical phenotype, we used dietary challenge. Mice were fed a high protein (HP) diet, as used previously (13), or a precursor-enriched (PE) diet comprised of increased levels of precursor amino acids of propionate pathway metabolites, i.e. isoleucine, valine, and threonine. Upon initiation of both diets at the age of 2 months, both Mut\(^{ko/ki}\) and Mut\(^{ki/ki}\) mice experienced rapid
weight loss, which was so severe that the study had to be terminated after 3 days (Fig. 5A). The effect exerted by the PE diet appeared to be stronger than the HP diet, as shown by the more pronounced weight loss (Fig. 5A), likely reflecting the two-fold higher isoleucine, valine and threonine content in the PE diet (70 g/Kg, 84 g/Kg and 53 g/Kg compared to 35 g/Kg, 42 g/Kg and 27 g/Kg, respectively). Brain weight (normalized to body weight) was increased in Mut ko/ki mice under both diets, which was not the case on reference chow (RC: isoleucine, 10 g/Kg; valine, 12 g/Kg; threonine, 7.6 g/Kg) at this age (Fig. 5B). Investigation of the plasma before and after the diet change revealed, from both diets, further elevated levels of MMA, C3, and for the first time in these mice, increased ammonia levels in Mut koi/ki and Mut ko/ki mice (Fig. 5C, D and E), consistent with the induction of metabolic crisis. Further investigation of MMA and 2-MC concentrations in the brain, kidney and liver demonstrated a gene dosage-dependent increase in both metabolites for mice on either diet, compared to mice on the control diet (RC) (Fig. 5F, G and H).

Indeed, while these metabolites were already elevated in Mut ko/ki and Mut kiki/ki mice on RC (e.g. brain levels of MMA in Mut ko/ki were 4.4 times higher than in Mut kiki/ki, Fig. 5F), they were further increased up to 53-times when the diet was changed to HP or PE (Fig. 5F).

Cobalamin treatment partly rescues diet induced metabolic crisis – To attempt to mitigate the effects exerted by the HP diet, we treated the mice with daily intraperitoneal (i.p.) injections of hydroxocobalamin (OHCbl, 0.3 µg/g) for a week before and continuously during the dietary challenge. These injections resulted in nearly doubled levels of plasma Cbl (Fig. 6A), indicating efficient uptake of the vitamin into the blood stream. Mut kiki/ki mice appeared to be partially protected by this treatment, as, in contrast to mice on the HP diet alone (Fig. 5A), OHCbl treated Mut kiki/ki mice did not show significant weight loss (Fig. 6B). Further, elevations in C3 and ammonia were markedly delayed, although they increased considerably, along with MMA, by the study’s end (Fig. 6C-E). Mut ko/ki mice showed a less obvious, but still slight protective effect. Their weight loss was delayed compared to their untreated counterparts (compare Fig. 5A HP and Fig. 6B), and they did not have the significantly increased brain weight (Fig. 6F) seen previously (Fig. 5B). However, they did show immediate and striking increases in metabolite levels (Fig. 6C-E) suggesting they remained metabolically compromised.

DISCUSSION

Novel mouse models recapitulate clinical and biochemical phenotype of MMAuria – In this study, we aimed to generate novel mouse models of MMAuria which survive through adulthood but show clear biochemical and clinical signs of disease. To achieve these models, we knocked in a missense mutation to the mouse Mut gene. This constitutive KI allele causes Mut deficiency, which is further aggravated by combination with a KO allele. The most striking phenotypic sign in our mice was growth retardation, which is a correlate of failure to thrive described in human patients (3,6). This lack of weight gain is not readily explained by reduced energy intake, as we identified no difference in overnight food consumption between the mutant mice and controls, suggesting that other, possibly subcellular, processes are affected. Previous studies point towards inhibited tricarboxylic acid cycle enzymes and interference with oxidative phosphorylation in MMAuria (27-29). It will be important to assess these potential mechanisms in future studies.

Further clinical signs could be observed when the disease was accelerated by means of a high protein challenge. The success of this approach was demonstrated by the substantial elevation of metabolites and immediate loss of weight in both diets (HP and PE) for both mouse genotypes (Mut ko/ki and Mut kiki/ki), suggesting that the propionate pathway was unable to process the additional metabolic load contributed by the modified diets. As indicated by the significantly elevated ammonia levels, this induced situation is consistent with acute metabolic crisis, where aggressive treatment is required to prevent encephalopathy when ammonia levels rise above a threshold of 200 µmol/l (30).

Along with these general organism-wide symptoms, we were able to identify organ-specific disturbances in the kidney and the brain. Under normal diet conditions, adult Mut ko/ki mice showed manifestations of kidney dysfunction, as evidenced by increased plasma urea, impaired diuresis and changes in the urinary excretion of electrolytes. These changes mirror the organ manifestations of MMAuria patients, in whom chronic kidney disease occurs in almost half of all cases in an age-dependent manner (3-6). In our mice, these changes were...
accompanied by increased renal levels of Lcn2, a protein that has been suggested to be an early biomarker of chronic kidney damage (31), including in MMAuria (13). Further, we found increased brain weight in the Mut<sup>ko/ki</sup> mice at one month and one year of age, possibly indicative of cytotoxic edema as a sign of neurotoxicity. Interestingly, we also found increased Lcn2 mRNA and protein levels in the brain of these mice. In contrast to its potential role as a kidney specific marker, upregulation of Lcn2 in the brain has been suggested to be a biomarker in multiple sclerosis (32), and was found to be upregulated in animal models of brain-specific diseases (33,34). Hence, the specific role of Lcn2 as an organ-specific biomarker remains unclear; however, the finding of upregulation in affected tissues suggests at least a general role in dysfunctional cells.

Biochemically, the phenotype of human patients was accurately emulated, as illustrated by the types of metabolites which were increased and the extent of their elevation. For example, MMA levels in the urine were detected in a range comparable to human MMAuria patients (35), while glycine, thought to be increased by inhibition of the intra-mitochondrial glycine cleavage enzyme due to accumulated organic acids or their CoA esters (36), was found at high levels in our mice, similar to the description in the first MMAuria patients (37). Further, we found elevated levels of odd-chain fatty acids, most likely as a consequence of increased C3 which is used as a primer in fatty acid synthesis (6,23,38,39). In addition, we present for the first time elevations of 14- or 18-chain length fatty acids. The pathophysiological significance of this newly discovered phenomenon remains unknown.

Overall, these novel models of MMAuria recapitulate many of the biochemical and clinical hallmarks of MMAuria in patients and thus are excellent tools to study the pathogenesis of disease. Application of a long-term dietary challenge with less potent diets will potentially allow monitoring of chronic disease progression, leading to more pronounced impacts on brain and kidney tissue.

**Gene dose-dependence allows titration of disease severity** – Patients carrying the p.Met700Lys mutation show an intermediate phenotype with relatively late onset and residual enzyme and pathway activity in fibroblast cell homogenates (15,16,40). Based on these observations, the measurable residual enzyme activity in vitro and the missense type of the variant used for the generation of the KI allele, we expected that the KI allele would result in a milder phenotype than the KO allele, which is in fact a truncating mutation. Indeed, a gene dose-dependent effect was corroborated by all experimental data: Mut<sup>ko/ki</sup> mice displayed higher concentrations of MMA, 2-MC, and C3, more pronounced growth retardation, and a stronger response to dietary challenge than homozygous Mut<sup>ki/ki</sup> animals. Therefore, the gene dose effect detected in our study is in analogy to the phenotypic differences observed in patients. This circumstance is a true benefit of this study since the novel animal models allow us to modulate the phenotypic severity of MMAuria and to systematically assess milder and more severe forms of MMAuria.

**Response to cobalamin treatment** – It is recommended to evaluate every mut-type MMAuria patient for responsiveness to cobalamin treatment (6,35). However, no systematic study has thus far investigated the effectiveness of cofactor treatment in mut-type MMAuria. If effective, the cofactor would be expected to ameliorate disease by supporting the enzyme’s activity and/or stability, similar to the cofactor response displayed by phenylalanine hydroxylase (41). The human mutant p.Met700Lys is an excellent candidate to test cobalamin responsiveness, as patient fibroblasts compound heterozygous for this and a premature stop mutation showed 4.5 times increased incorporation of 14C-propionate into acid precipitable material when supplemented with OHCbl (40), and biochemically this mutation was found to affect the K<sub>M</sub> of MUT for AdoCbl (14). Indeed, in our HP diet study, Mut<sup>ki/ki</sup> mice appear to have been at least partly protected by pre-administration of OHCbl. These mice did not show the immediate weight loss associated with the initiation of the HP diet, and showed delayed accumulation of metabolites. The reduced degree of protection for Mut<sup>ko/ki</sup> mice suggests that with the level of Cbl achieved by these injections, both mutant alleles are required to be potentially Cbl-responsive. An even better rescue may be facilitated by the application of higher Cbl doses in these mice, potentially reaching levels suggested for treatment in human patients (35). Nevertheless, our results set the stage for further exploration of cofactor response –
a concept which may be applicable to many other missense mutations, since about one quarter of all mut-type mutations show an in vitro response to cobalamin (3).

**EXPERIMENTAL PROCEDURES**

**In vitro cell culture and enzyme activity assay** – Constructs of human (hs) MUT wild-type and mouse (mm) Mut wild-type and the mutants hs-p.Met700Lys and mm-p.Met698Lys were made by site-directed mutagenesis in the pTracer-CMV2 vector (Thermo Fisher Scientific) using the QuickChange site-directed mutagenesis kit (Stratagene) following manufacturer’s instructions. Constructs were transiently transfected into immortalized fibroblasts carrying the homozygous MUT mutation p.Gln30* by electroporation. Cell lysates were obtained by sonication in 5 mM potassium phosphate buffer (pH 7.4) and enzymatic activity and $K_M$ determined as previously described (14).

**Mouse tissue preparation** – Liver, kidney, brain, heart, and muscle as well as blood samples were removed from euthanized mice and stored at -80°C. Lysates of homogenized (TissueLyser II, Qiangen) tissues were resuspended in 300-600 µl of a lysis buffer dependent on the subsequent usage of the sample, either Buffer A (TritionX-100 0.5% (v/v), 10 mM HEPES at pH 7.4, 2 mM DTT, 1 tablet of Complete Protease Inhibitor Cocktail Tablets, Roche) for Western blot and activity assay or Buffer B (250 mM sucrose, 50 mM KCl, 5 mM MgCl2, 20 mM Tris base, adjusted to pH 7.4) for determination of MMA and 2-MC levels. The homogenate was then centrifuged at 14'000 rpm for 10 min at 4°C and the supernatant was used for MUT activity assay (14), Western blotting and qRT-PCR (see below).

**Housing of mice and generation of knock-in allele** – Animal experiments were performed in accordance with policies of the Veterinary Office of the State of Zurich and Swiss law on animal protection. Animal studies were approved by the Cantonal Veterinary Office Zurich under the license number 202/2014. Animals were kept in single-ventilated cages and under controlled humidity and temperature (21-23°C). Whenever possible littermate controls were used to compare experimental groups.

The generation of mice carrying the Mut-p.Met698Lys mutation was performed by Polygene (Rümlang, Switzerland) using embryonic stem cell targeting. To generate Mut$^{kox}$ mice, female Mut$^{kowt}$ (7) were crossed to Mut$^{koki}$ males. Mouse genotyping was performed on genomic DNA from ear punch biopsies using the primers 5’-GTGGGTGTCAACACACTTG-3’ (forward) and 5’-CGTATGACTGGGATGCCT-3’ (reverse) for the ki-allele and 5’-ACAACTCTTGTG-TAGTGTC-3’ (forward) and 5’-CCITTAGGATGTCCATCTCG-3’ (reverse) for the ko-allele.

**Long-term studies and metabolic cage studies** – Long-term monitoring of mice entailed weekly weight measurements and regular blood collections as well as urine collections. Animals were single caged overnight to collect urine and measure individual chow and water intake. Urine was collected in the morning, the sediment removed and supernatant frozen at -80°C.

**Quantitative real time PCR analysis and Western blot** – Total RNA (QIAmp RNA Blood Mini Kit, Qiagen) was extracted from frozen mouse tissue lysates and analyzed for expression using specific probes for Mut (Mm00485312_m1) and Lcn2 (Mm01324470_m1) via the TaqMan Gene Expression Master Mix on a 7900HT Fast Real-Time PCR System, with normalization to beta-actin (Mm00607939_s1) (all Thermo Fisher Scientific). Experiments were performed in triplicates. For Western blotting analysis, antibodies against MUT (ab67869, 1:500, Abcam), Lcn2 (ab63929, 1:1000, Abcam) and beta-actin (A2228, Sigma-Aldrich) were detected by horseradish peroxidase (HRP)-labelled goat anti-mouse (sc-2302, Santa Cruz) or goat anti-rabbit (sc-2301, Santa Cruz) IgG at a dilution of 1:5000. Bands were quantified by normalization to internal beta-actin control bands.

**Metabolite measurements** – MMA in mice urine was analysed by liquid chromatography mass spectrometry (LC-MS/MS) on a Thermo Scientific UltiMate 3000 Rapid Separation LC coupled to an AB Sciex 5500 TripleQuad mass spectrometer using a commercial Kit (Recipe ClinMass® advanced). For determination of the concentrations of amino acids, acylcarnitines (C2 and C3) as well as MMA in dried blood spots, blood was collected from tail vein onto a filter card and dried. Punches from filter cards were analyzed by tandem mass spectrometry similar to (42). Total fatty acid profiles from plasma/serum were determined by direct methylation and gas chromatography, as described (43). Sphingoid bases were analyzed in tissue homogenates, as published (44). Ammonia levels in whole blood were measured by the PocketChem
blood ammonia meter (PA-4140, Arkray). Electrolytes in urine were measured by a timed endpoint method (SYNCHRON System kit, Beckman Coulter). For urea determination in plasma the SYNCHRON CX3 Delta System was used (Beckman Coulter, Nyon), kit no. 443350.

Diet and cobalamin therapy – We applied a high protein (HP, 60% protein, TD.140830, Harlan) and precursor-enriched (PE, 700% isoleucine, valine and threonine compared to reference, TD.140829, Harlan) diet to 60-day-old mice. For the PE diet, leucine (19 g/kg, 119%) was enriched since its uptake might compete with the uptake of the other amino acids which are increased in the diet and cystine was increased (3.5 g/kg, 700%) to elevate the overall sulfur content. For cobalamin rescue, mice were injected with 0.3 µg hydroxocobalamin (Streuli Pharma AG) i.p. 7 days before and throughout diet treatment.
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CONFLICT OF INTEREST STATEMENT
The authors declare no conflict of interest.

AUTHOR CONTRIBUTIONS
PF designed the research together with MRB, DSF, OD and performed most of the experiments. Experimental work was also carried out by AS, MM, NN, AZ. New reagents or analytic tools were contributed by DM, CDL, JH, LS, RF, HLP, TH, BT, SK, OD. Data was analyzed by PF, MRB, DSF, OD, PB, TH. The manuscript was written by PF, MRB, DSF, OD.

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**FIGURE/TABLE LEGENDS**

**Table 1.** Human and mouse mutant proteins show similar enzyme parameters when expressed in a mut\(^0\) patient cell line. Mean enzyme activities were measured in fibroblast homogenates after overexpression of the respective constructs. Mutant values were calculated in relation to their wt counterparts and normalized to percent residual activity of the respective wt (set to 100%). \(K_M\) values for AdoCbl were determined by measuring MUT activity with increasing concentrations of AdoCbl ranging from 0.0025 to 50 \(\mu\)M in the same homogenates as above. Displayed numbers represent mean values ± S.D.

**Fig. 1.** Generation of KI allele and initial characterization. (A) Based on the wt mouse Mut gene, a targeting construct was generated to replace the Mut gene region spanning from intron 8 to intron 12 (shaded in grey), consisting of a long and short arm of homology as well as an insert with the missense mutation (p.Met698Lys) and a neomycin cassette (neo, green arrow) flanked by FRT sites (vertical green triangles). Homologous recombination and Flp-deletor breeding resulted in the final targeted KI allele with the mutation and one remaining FRT site. Exons are represented by red triangles or boxes, arrows “fwd” and “rev” indicate forward and reverse genotyping primers, respectively. (B) Southern blotting of targeted clone 12H5. (C) Upper panel: genotyping PCR of ear biopsy DNA for the KI allele with fwd and rev primers depicted in panel A. Asterisk indicates an additional heterodimer band as confirmed by Sanger sequencing. Lower panel: genotyping PCR of KO allele. (D) Mut expression among wt tissues (error bars are SEM; \(n = 4\); *: \(p < 0.01\)) and (E) KO allele dependent loss of Mut expression (bars are means normalized to wt in each tissue; error bars depict SEM; \(n ≥ 4\); *: \(p < 0.05\)). (F) Western blot analysis of Mut protein levels using \(β\)-actin as loading control. The lower band in the top panel of each organ represents an unspecific band. (G) Mut activity varies by tissue and (H) by genotype (bars are mean values and in each tissue normalized to the wt value; error bars are SEM; \(n ≥ 4\); *: \(p < 0.0001\)). Significance levels are the same in all five tissues (not depicted for clarity).
Fig. 2. Clinical and biochemical phenotype of Mut<sup>ki/ki</sup> and Mut<sup>ko/ki</sup> mice (red, females; blue, males, genotypes see legend). (A) Monitoring of body weight over time. Decreased body weight in female Mut<sup>ki/ki</sup> mice is significant compared to Mut<sup>ki/wt</sup> from day 162, and in Mut<sup>ko/ki</sup> significant compared to Mut<sup>ki/wt</sup> from day 114. For both: p < 0.05. (B) C3 levels in dried blood spots normalized to acetylcarnitine (C2). (C) MMA levels measured by LC-MS/MS in urine collected overnight in metabolic cages. In panels A-C points represent mean values, error bars depict SEM and n = 5 per group. (D) MMA levels in tissues (left panel), urine (middle) and dried blood (right), determined by tandem mass spectrometry. (E) C3 levels normalized to C2 in dried blood. (F) Ammonia levels measured in whole blood. (G) Glycine concentration determined in dried blood by tandem mass spectrometry. (H) Fatty acid levels determined in plasma and expressed as percent of total fatty acids. (I) Plasma levels of sphingoid bases (expressed as pmol per µl plasma). (J) 17-carbon chain sphingoid base levels in different tissues expressed as pmol per µg protein. Panels D-J: bars represent mean values from 35 days old mice (error bars depict S.D.; n ≥ 4; *: p < 0.05).

Fig. 3. Renal and neurological phenotype. (A) Urine production measured in metabolic cages overnight and expressed as microliters of urine per hour normalized to body weight. (B) Overnight water intake measured in grams of water normalized to body weight. (C) Urea in plasma expressed in mg per deciliter. (D) Electrolytes measured in urine and normalized to grams of creatinine. (E) Western blot of Mut and Lcn2 in kidney at two different time points. (F) Brain weight normalized to body weight at three different time points. (G) Expression of Lcn2 in brain tissue on mRNA level normalized to β-actin and on protein level (β-actin as loading control). In panels A-D and F-G, points and bars represent mean values; error bars depict SEM; *: p < 0.05.

Fig. 4. Histological analysis by hematoxylin-eosin staining of (A) kidney and (B) brain sections from Mut<sup>ki/wt</sup> and Mut<sup>ko/ki</sup> female mice at the age of 1 year. Scale bars are 10 µm in the kidney pictures, 1 mm on brain overview pictures and 100 µm in the insets of the basal ganglia.

Fig. 5. Modified diet leads to disease acceleration. (A) Start of both modified diets (upper panel: PE, precursor-enriched; lower panel: HP, high protein) at 60 days of age is indicated by the black arrows. Monitoring of body weight showed significant differences among the three groups on day 63 for both diets. (B) Weight of brain normalized to body weight after diet study. (C) MMA in dried blood. (D) C3 (normalized to C2) in dried blood. (E) Whole blood ammonia before and after diet modification. (F-H) MMA and 2-MC levels in homogenates of (F) brain, (G) kidney, and (H) liver tissue on reference chow (RC), high protein (HP) and precursor-enriched (PE) as determined by GC-MS. For all panels, from 5 animals for each genotype bars and points represent means; error bars are SEM. For panels A-E, *: p < 0.01; n.s., not significant; statistics were not calculated for panels F-H.

Fig. 6. Treatment of HP diet-induced phenotype by OHCbl. (A) Cobalamin levels in plasma of untreated mice and OHCbl treated mice; *: p < 0.0001. (B) Time course of body weight. Black arrow indicates start of HP diet on day 60. Differences of body weight on day 64 were significant between Mut<sup>ko/ki</sup> and the other two genotypes (Mut<sup>ki/ki</sup> and wt), which did not differ significantly from each other. (C) C3 (normalized to C2), (D) MMA levels in dried blood and (E) whole blood ammonia levels were measured on the day the diet was started (day 60) and on day 62 and day 64 following diet commencement. (F) Weights of brain whole organs in sacrificed animals at the end of the study. In panels B-E duration of HP diet was from day 60 to day 64 and OHCbl treatment from day 53 to day 64 as indicated by capped horizontal lines. Points and bars represent mean values; error bars in all panels depict SEM.
<table>
<thead>
<tr>
<th>Activity</th>
<th>Human</th>
<th>Mouse</th>
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<tbody>
<tr>
<td></td>
<td>100.0 ± 22.4</td>
<td>3.46 ± 0.38</td>
</tr>
<tr>
<td>$K_M$</td>
<td>2.59 ± 0.87</td>
<td>575.6 ± 46.0</td>
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</tbody>
</table>
A Mouse wild-type Mut gene

Targeting construct

B wt 12H5

10.3 kb: targeted
8.4 kb: wild-type

C

ES cell targeting and Flp breeding

Targeted allele

D Mut-wt expression

E Mut expression in tissues

F

G Mut-wt activity

H Mut activity in tissues

Figure 1.
Figure 2.
Figure 3.
Figure 4.
Figure 5.
Figure 6.
Novel Mouse Models of Methylmalonic Aciduria Recapitulate Phenotypic Traits with a Genetic Dosage Effect

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