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Homocystinuria, which typically results from cystathionine β-synthase (CBS) deficiency, is the most common defect of sulfur amino acid metabolism. CBS condenses homocysteine and serine to cystathionine that is then converted to cysteine. Individuals with homocystinuria have markedly elevated plasma levels of homocysteine and methionine and reduced concentrations of cystathionine and cysteine. Clinical disease manifestations include thromboembolism and neuropsychiatric, ocular, and skeletal complications. Here, we have shown that administration of PEGylated CBS into the circulation of homocystinuria model mice alters the extra- and intracellular equilibrium of sulfur amino acids, resulting in a decrease of approximately 75% in plasma total homocysteine (tHcy) and normalization of cysteine concentrations. Moreover, the decrease in homocysteine and the normalization of cysteine in PEGylated CBS–treated model mice were accompanied by improvement of histopathological liver symptoms and increased survival. Together, these data suggest that CBS enzyme replacement therapy (ERT) is a promising approach for the treatment of homocystinuria and that ERT for metabolic diseases may not necessitate introduction of the deficient enzyme into its natural intracellular compartment.

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individuals to enjoy an unrestricted or only mildly restricted diet and thus significantly improve their quality of life. To provide these benefits, a therapy must deal with the core deficiency underlying this condition, namely the aberrant CBS function. Accordingly, we envisioned that systemic introduction of CBS, in the form of enzyme replacement therapy (ERT), may be beneficial to homocystinuria patients.

ERT is well known for its successful application in lysosomal storage diseases (LSDs) and other genetic disorders. However, the use of ERT to treat CBSDH presents a number of specific obstacles that need to be addressed. First, in LSDs, for example, the exogenous enzyme is glycosylated and thus taken up by receptor binding to the sugar moiety, followed by uptake and trafficking into the lysosome. This natural uptake to the site of need is not a trait of CBS that can be exploited for homocystinuria. We asked whether introduction of a recombinant enzyme into the circulation, with no trafficking to its native intracellular environment, would favorably affect intracellular metabolism. We assumed that a significant decrease in extracellular homocysteine due to CBS administration can generate a concentration gradient, triggering a flux of homocysteine from intra- to extracellular spaces, where the administered enzyme can further process it, with the extracellular CBS thus serving as a homocysteine “sink.”

A second obstacle is rooted within the structure and mode of activation of native human CBS (hCBS), which exists as a tetramer composed of 4 identical monomers. This form of the enzyme has a high tendency toward aggregation, which poses a major constraint on purification efforts (13). Additionally, CBS is allosterically regulated by S-adenosyl-methionine (SAM), which binds to the autoinhibitory C-terminal regulatory domain and activates the enzyme by approximately 5-fold in vitro (14). SAM levels in CBS-deficient patients and controls are 1.38 μM and 0.093 μM, respectively (15), in both cases far below the $K_m$ value for CBS (17.7 μM) (16), which would prevent full activation of hCBS if administered to patients. However, it was previously determined that a human truncated CBS (htCBS), lacking 138 amino acids from the C-terminus (the regulatory domain), forms dimers and is constitutively activated, regardless of SAM. Moreover, this form of the enzyme tends to form fewer aggregates (17, 18). Together, these data imply that htCBS is better suited than the WT enzyme for use in the development of ERT to treat CBSDH.

In addition, key pharmacokinetic and pharmacodynamic considerations must be addressed. Therapeutic proteins, unlike small molecules, are exclusively delivered via parenteral routes, and treatment efficacy may be greatly impacted by their absorption, distribution, metabolism, and excretion (ADME) (19). High-molecular-weight compounds, such as enzymes, have limited tissue penetration capability and are thus mainly present in the plasma. These proteins may then be maintained in the circulation for a short period of time, as they are removed from the bloodstream by several mechanisms (19). Ideally, administered CBS is expected to maintain high activity in plasma for a period of time that is sufficient to have a steady and significant effect on sulfur amino acid metabolism. Accordingly, enhancing ERT efficacy may require additional modifications to the protein in order to increase the retention time in vivo. This may be achieved by PEGylation, the addition of polyethylene glycol (PEG) moieties onto the surface of the protein, a strategy that has become widely accepted. PEGylation was found to minimize proteolysis, immune response, and antigenicity, while increasing protein stability and size and reducing renal excretion (20).

Here, we present a body of work that characterizes several modified forms of htCBS and show for the first time to our knowledge that exogenously administered CBS, in which exposed cysteine residues are PEGylated, positively affects the aberrant concentration of sulfur metabolites in homocystinuric mice. Furthermore, our ERT restores metabolic control in the face of a standard diet, ameliorates histopathological changes in the liver, and improves survival of a KO mouse model. The delivery of PEGylated htCBS thus represents a potential new therapeutic approach for CBSDH.

**Results**

*Unmodified htCBS exhibits a short retention time in the circulation.* The pharmacokinetic properties of a pharmacologically active substance that is administered into the circulation are greatly affected by natural mechanisms of ADME. Rapid clearance of the injected molecule may greatly impact treatment efficacy, and thus longer circulating half-lives are desired, which may translate into less frequent administrations or smaller doses that in turn minimize the side effects. To determine the pharmacokinetics of htCBS, a single dose of 5 mg/kg body weight (BW) was injected via the s.c., i.v., or i.p. route into C57BL/6j mice. As shown in Figure 1A, administration of htCBS via the i.v. or i.p. route reached the highest CBS-specific activity values during the first 4 hours after injection (observed peak plasma levels of up to 123 and 76 mU/μl, respectively. See also Supplemental Table 1 for AUC values; supplemental material available online with this article; doi:10.1172/JCI85396DS1). For the s.c. mode of delivery, lower specific activity values were recorded (highest value of 34 mU/μl plasma). All routes of administration demonstrated similar values at 8 hours after injection. Most important, however, is that 24 hours after injection, only residual htCBS activity was detected, regardless of the route used. The pharmacokinetic parameters (Supplemental Table 1) showed that the half-life for htCBS in circulation was 2.7 hours, suggesting that after roughly 6 half-lives, less than 20 hours, htCBS levels should be undetectable. The bioavailability of htCBS was 50% for the s.c. route, suggesting that s.c. delivery may be a clinical option for this therapeutic approach. The longer apparent htCBS half-lives for s.c. and i.p. delivery (Supplemental Table 1) can be explained by a slow absorption phase that occurs during the elimination phase, thereby extending the mean residency time and apparent half-life. However, for our purpose, a much longer half-life was desirable, and thus we moved on to test PEGylated species of htCBS.

The rapid decline in htCBS activity in vivo may be attributed to a facilitated clearance from the circulation and may also be attributed to the loss of enzymatic activity of htCBS once introduced into the circulation. To test the latter, we incubated the htCBS enzyme in vitro in mouse plasma at 37°C for up to 96 hours and measured activity every 24 hours. As shown in Figure 1B, no significant loss of activity was recorded for up to 72 hours of incubation. A modest 25% loss of activity was recorded after 96 hours of incubation. Thus, we conclude that clearance from the circulation, rather than loss of activity in plasma, accounts for the rapid loss of htCBS activity in the circulation, as also shown below.
generate the ME020MA PEGhtCBS) or a high 40-kDa molecular-weight, 4-arm branched PEG (designated GL4-400MA, to generate the GL4-400MA PEGhtCBS). PEGylation did not affect the specific activity of the enzyme (Figure 1C). A dose of 5 mg/kg BW PEGylated htCBS (PEGhtCBS) was administered to C57BL/6J mice via the s.c. route, and CBS activity was monitored at different time points. The PEGylated enzymes showed significant activity in the circulation 24 and 48 hours after injection. As shown in Figure 1D, ME020MA PEGhtCBS showed specific activity of 41 and 14 mU/μl plasma, and GL4-400MA PEGhtCBS showed specific activity of 65 and 28 mU/μl plasma 24 and 48 hours after injection, respectively. As shown in Figure 1E for GL4-400MA PEGylation enhances the plasma half-life of htCBS in vivo. Conjugation of PEG to biopharmaceuticals has emerged as the most promising technique to positively impact the pharmacokinetics of a variety of parenterally administered substances. Accordingly, and since the modification of cysteine residues with maleimide-activated PEG is specific, simple, and forms a biologically inert thioether bond, we monitored the effect of cysteine-directed PEGylation of the enzyme on activity in vitro and on retention time in vivo (see also Supplemental Figure 2A, which shows most of the cysteine-directed PEG types used, presented with good retention times in vivo). For that purpose, htCBS was PEGylated with a low 2-kDa molecular-weight linear PEG (designated ME020MA, to

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generate the ME020MA PEGhtCBS) or a high 40-kDa molecular-weight, 4-arm branched PEG (designated GL4-400MA, to generate the GL4-400MA PEGhtCBS). PEGylation did not affect the specific activity of the enzyme (Figure 1C). A dose of 5 mg/kg BW PEGylated htCBS (PEGhtCBS) was administered to C57BL/6J mice via the s.c. route, and CBS activity was monitored at different time points. The PEGylated enzymes showed significant activity in the circulation 24 and 48 hours after injection. As shown in Figure 1D, ME020MA PEGhtCBS showed specific activity of 41 and 14 mU/μl plasma, and GL4-400MA PEGhtCBS showed specific activity of 65 and 28 mU/μl plasma 24 and 48 hours after injection, respectively. As shown in Figure 1E for GL4-400MA
PEGhtCBS, the activity in plasma was significantly extended for the PEGylated protein as compared with that of the non-PEGylated counterpart. Figure 1E, together with Supplemental Table 1, thus further attests to the rapid clearance of the non-PEGylated enzyme, supporting rapid clearance as the main mechanism for loss of activity of this enzyme.

As shown in Supplemental Table 2, the half-life increased from 2.7 hours for the non-PEGylated htCBS to 16.7 and 30.4 hours for htCBS PEGylated with ME020MA and GL4-400MA, respectively. This means that the time of exposure after a single dose would be 5- to 10-fold longer for the PEGylated forms. Additionally, the bioavailability after s.c. administration ranged from 50% to 80%, suggesting that the s.c. route is a reasonable one for clinical studies.

A repeated injection regimen was also tested. C57BL/6J mice were injected 3 times with 5 mg/kg BW of htCBS on 3 consecutive days, and blood samples were collected every 24 hours, as well as after 4 hours after the first and third injections. Figure 2 shows an activity peak of 20 mU/μl plasma 4 hours after injection. However, 24 hours after each injection, only residual activity could be demonstrated for the non-PEGylated enzyme, whereas for the GL4-400MA PEGhtCBS, significantly higher activity levels (up to 84 mU/μl plasma) for a substantial period of time were observed, demonstrating the superiority of the PEGylated enzyme compared with its non-PEGylated counterpart.

PEGhtCBS administration positively impacts plasma levels of total homocysteine and cystathionine concentrations and restores normal cysteine levels. The ultimate goal for the use of htCBS to treat CBSDH is to lower the toxic total homocysteine (tHcy) load and elevate cystathionine and cysteine levels. These changes are expected to prevent, reverse, or delay the onset of CBSDH symptoms. Homocystinuria research, however, has long been hindered by the lack of a suitable animal model. Mice with a completely knocked-out CBS gene die within 2 to 3 weeks after birth. The human-only (HO) mice, which were previously generated in our laboratory, are different in that, in addition to being KOs for the mouse gene, they are transgenic for the human gene and express low levels of hCBS, which allows their survival to adulthood (21). HO mice exhibit severe elevations of tHcy, methionine, S-adenosylmethionine, and S-adenosylhomocysteine and present with a concomitant decrease in plasma and hepatic levels of cysteine. Accordingly, these mice exhibit characteristics that, in several aspects, recapitulate the disease as it occurs in humans.

Since diurnal variation may obscure the htCBS-dependent metabolic changes, we followed the transsulfuration metabolites throughout a 24-hour cycle. Figure 3A shows that tHcy levels changed considerably, and thus it was decided to administer the enzyme and perform blood collections at the time of the lowest tHcy levels (15:00), when the effect of the enzyme could be isolated from the natural tHcy clearance mechanisms (data for each mouse are presented in Supplemental Figure 1 for all metabolites).

In order to evaluate the impact of the administered PEGhtCBS on sulfur amino acid metabolites in vivo, we treated HO mice with either 5 mg/kg BW of GL4-400MA PEGhtCBS or the non-PEGylated enzyme for 5 consecutive days. Treatment was then stopped for 10 days and restarted on day 14 for an additional 5 consecutive days of once-daily injections. Figure 3B describes the changes in tHcy levels for individual HO mice, and Figure 3C depicts the average tHcy and cystathionine levels. Initial tHcy levels varied greatly among individual HO mice (100–285 μM). However, once treated with the PEGylated enzyme, a significant drop in tHcy plasma concentrations (up to a 250 μM reduction, representing an 87% drop for mouse 435) was apparent and was remarkably similar for all mice treated. As shown in Figure 3C, the mean concentration of tHcy 24 hours after the second injection was 55 μM, a 70% decrease from the initial 181 μM concentration levels. Similarly, tHcy levels were in the range of 38 μM to 45 μM 24 hours after the fourth and fifth injections, respectively. A blood sample collected on day 11 demonstrated that the effect was not completely diminished 7 days after the last injection, with tHcy values at 112 μM (62% of the initial value). On day 14, tHcy levels were approaching pretreatment levels, and renewing the injection regimen for an additional 5 consecutive days resulted in a drop of tHcy levels to 60 μM. Cystathionine levels were also greatly increased following the injections, inversely reflecting the changes in tHcy levels. As shown in Figure 3C, whenever tHcy levels were reduced, cystathionine levels were increased and vice versa. Cystathionine levels rose to as high as 43 μM (24 hours after the fifth injection; day 5), representing an approximately 9-fold increase from the

Figure 2. Repeated injection of the PEGylated, but not the non-PEGylated, htCBS shows buildup of CBS activity in vivo. C57BL/6J mice were injected at 0, 24, 48 hours (indicated by arrows) with 5 mg/kg BW of the non-PEGylated htCBS (n = 5) or with GL4-400MA PEGylated htCBS (n = 5) and were bled at the indicated time points. Data are presented as the mean ± SEM and were compared using an unpaired Student’s t test. **P ≤ 0.01 and ***P ≤ 0.001.
Although using the GL4-400MA PEGylated enzyme resulted in a significant improvement in biochemical changes, this specific PEG is more complex in nature (4 arms), expensive, and, to our knowledge, has not been used in any approved drug. Accordingly, we tested several different PEG chemistries (Supplemental Figure 2), and our data indicate that CBS PEGylated with PEGs of lower than 20-kDa molecular weight demonstrated a more rapid loss of CBS activity in plasma compared with CBS enzymes modified initially 4.8 μM at time 0 (the reference ranges for cystathionine in WT and Cbs<sup>−/−</sup> heterozygous mice are 2.5 ± 0.3 μM and 5.0 ± 0.8 μM, respectively) (22). Importantly, as shown in Figure 3E, treatment with PEGhtCBS normalized the cysteine levels, starting at 145 μM and rising to 225 μM during the treatment periods. In contrast to the effect of the PEGylated enzyme, no changes in tHcy or cystathionine levels were observed upon administration of non-PEGylated htCBS (Figure 3, D and E).

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with PEGs of higher molecular weight. For that reason, we opted to continue with the linear 20-kDa PEG, designated ME200MA0B (Supplemental Figure 2B). In addition, from a drug manufacturing perspective, this PEG yields a preparation of lower viscosity as compared with that of the 40-kDa PEGs and is cheaper.

Next, we monitored the long-term effect of PEGhtCBS administration. The enzyme was administered to 8 HO mice twice per week for the first 2 weeks and then 3 times per week for the remaining 6 weeks. The dose was increased to 7.5 mg/kg BW in order to maximize the effect. Blood samples were collected 24
hours after the first weekly injection and 72 hours after the last weekly injection. As shown in Figure 4A, plasma tHcy levels were reduced from 212 μM at time 0 to an average in the range of 62 to 103 μM 24 hours after the injections and were within a range of 141 to 189 μM 72 hours after the last weekly injection. From day 28 on, the 72-hour post-injection values were significantly lower than the time 0 value (P ≤ 0.02). Cystathionine and cysteine levels were positively influenced as well. Cystathionine levels increased from 6 μM to as high as 30 μM (Figure 4B), and cysteine levels were constantly above 200 μM 24 hours after injection (Figure 4C). From day 24 on, fluctuations in cystathionine concentrations were much less pronounced compared with those observed in the first 3 weeks of treatment. We also analyzed liver, kidney, and brain tissues to determine the levels of nonprotein-bound homocysteine (free and disulfide-linked forms), known to be significantly elevated in HO mice compared with the 1–5 nmol/g levels detected in WT mice (22). As shown in Figure 4D, a significant reduction of 44%, 63%, and 47% in nonprotein-bound homocysteine was detected in liver, kidney, and brain tissues, respectively.

A C15S-mutated form of htCBS exhibits no aggregation and a uniform PEGylation pattern. Although it was previously reported that htCBS is less prone to aggregation as compared with the full-
length protein (17), it still forms tetramers and higher aggregates. Such soluble aggregates may stimulate immune defense mechanisms and lead to, e.g., amyloid deposition (23), but also represent a technological challenge for process development. Batch-to-batch variability, multiple forms as a result of oligomerization or aggregation, and inconsistent PEGylation represent major difficulties for drug development efforts. As shown in Figure 5A, different batches of htCBS showed different dimer/tetramer ratios, with varying degrees of higher aggregates. An in-gel activity assay demonstrated that these aggregates had CBS activity (Figure 5B). Aggregation causes variable PEGylation patterns, as shown in Figure 5C. PEGylation resulted in 2 distinct bands, presumably reflecting a varying degree of PEGylation, as aggregation may obscure the available PEGylation sites. Accordingly, because of a more pronounced aggregation of batch 28 (Figure 5A), htCBS PEGylation resulted in a more pronounced lower band, i.e., fewer PEGylated species, on SDS-PAGE (Figure 5C), whereas PEGylation of batch 112, composed mostly of tetramers and dimers (Figure 5A), resulted in a more pronounced upper band as compared with that observed in batch 28 (Figure 5C). Incubation of both enzyme batches with the reducing agent tris(2-carboxyethyl)phosphine (TCEP) converted many of the tetramers and higher aggregates to a dimeric form (Figure 5D). These data imply that aggregation is driven by the formation of disulfides between exposed cysteines on the surface of htCBS dimers. Thus, we hypothesized that TCEP treatment may also result in a more extensive PEGylation, as more dimers are generated from higher-molecular-weight forms, exposing additional potential PEGylation sites. Indeed, as shown in Figure 5E, the equilibrium between the upper and lower PEGylation bands on an SDS-PAGE was significantly shifted toward the upper band following TCEP treatment as compared with PEGylation in the absence of TCEP. In order to gain more control and reproducibility in the PEGylation process, we mutated the C15 residue, which was previously shown to be the most reactive cysteine on the surface of CBS (17), to serine, thereby generating the C15S htCBS mutant. Figure 5F depicts a Coomassie-stained native gel, in which 2 main bands (dimers and tetramers) were observed for htCBS, whereas the C15S htCBS mutant had a single band, which, as expected, was not affected by treatment with TCEP. This observation was further substantiated by size-exclusion HPLC (SEC-HPLC). The single peak for the dimeric C15S htCBS mutant eluted at 8.62 minutes, while the main peak for the htCBS eluted at 7.9 minutes, corresponding to a tetramer (Figure 5H). As shown in Figure 5G, the absence of aggregates in the C15S htCBS resulted in a more consistent PEGylation pattern between different batches with, almost exclusively, a single PEGylated species appearing on SDS-PAGE.

We also tested the effect of PEGC15S at 1, 4, and 24 hours after injection (Supplemental Figure 4). Interestingly, whereas the effect on tHcy and cysteine was observed only 24 hours after injection (Supplemental Figure 4, A and C), a significant increase in cystathionine was already observed at 1 and 4 hours after injection (Supplemental Figure 4B).

**PEGC15S and betaine operate synergistically in vivo.** It is reasonable to assume that ERT for homocystinuria, once approved, will be administered on a background of protein restriction and betaine supplementation. Thus, it is important to determine the effect of ERT therapy combined with betaine. Accordingly, we compared the effect of betaine, PEGC15S, and a combination of betaine plus PEGC15S treatments on metabolites of the transsulfuration pathway.

The betaine-only and betaine-plus-PEGC15S groups were treated with betaine throughout the 18 days of the experiment, and the latter group was injected with PEGC15S on days 14 and 15. Mice in the PEGC15S-only treatment group were maintained on water with no betaine and injected on days 14 and 15 as well. As shown in Figure 6A, 14 days of betaine treatment resulted in a 29% and 32% decrease in tHcy for the betaine-treated groups. The betaine-only treatment group maintained these levels of tHcy out to day 18. Injection of PEGC15S on the background of
ment with PEGC15S alone. As shown in Figure 6A, the difference between these 2 groups became statistically significant 48 hours after the second injection (on day 17; \( P \leq 0.05 \)). Thus, we conclude that the combination of PEGC15S and betaine allows the maintenance of lower tHcy levels for an extended period of time.

Betaine treatment did not result in an increase in cystathionine levels over the 18-day experimental period (Figure 6B). In contrast, betaine resulted in a more pronounced reduction of 74%, 77%, 76%, and 60% on days 15, 16, 17, and 18, respectively. Treatment with PEGC15S alone resulted in a reduction of 69%, 67%, 52%, and 33% on days 15, 16, 17, and 18, respectively. According to these data, injection of PEGC15S with or without betaine supplementation was far superior to betaine treatment alone. In addition, the combination of betaine plus PEGC15S was superior to treat-
an increase in these levels was observed for the 2 groups that were treated with PEGC15S (without or with betaine). Interestingly, the combination of PEGC15S plus betaine resulted in an increase in cystathionine levels, which, starting on day 16 and beyond, were significantly lower than those in the group that received only the PEGC15S enzyme. This is likely due to the fact that treatment with PEGC15S alone channels homocysteine to condense with serine to form cystathionine. Combination with betaine diverts some of the available homocysteine through the remethylation pathway to produce methionine, and thus less homocysteine is available for PEGC15S to be converted to cystathionine.

PEGC15S rescues KO mice from early death and improves liver disease. The majority of CBS-KO mice die within 2 to 3 weeks after birth. Thus, we hypothesized that administration of PEGC15S to KO pups early after birth would rescue them. The mice were injected twice per week with PEGC15S or PBS for 5 weeks. During the first 21 days, the lactating mothers had drinking water containing 2% betaine (an estimated daily intake of 4 g/kg). As shown in Figure 7A, on day 21, only 18% of the mice survived in the PBS-injected group as opposed to 93% of those that were injected with the enzyme. No PBS-injected animal survived to day 24, while an 86% survival rate was recorded for the PEGC15S-injected group. From day 24 and beyond, the number of enzyme-injected animals began to decline as well, with 45% survival on day 29 and 33% on day 35 (Figure 7A).

It was previously demonstrated that KO mice suffer from severe liver damage (22). Prolonged survival of KO mice may thus be correlated with improvement in liver function after CBS treatment. Accordingly, we performed histological analysis of liver sections from animals that were injected twice weekly with PEGC15S for a period of 35 days. Animals were sacrificed on day 36 and liver samples processed for histological analysis by optical microscopy. PBS-injected animals did not survive to day 35, but for 2 animals, we were able to take liver samples for processing immediately after death on days 17 and 24, respectively (Figure 7B). The analyses were carried out in a blinded fashion. PBS-injected animals developed severe hepatopathies characterized by pronounced microvesicular to macrovesicular steatosis and substantial death of hepatocytes, with minimal or no fibrosis. Multiple mono or oligo cellular necroses of hepatocytes dispersed throughout the liver lobule, accompanied by a prominent inflammatory resorptive reaction, were found in 1 animal, whereas massive confluent necroses of hepatocytes mainly in perportal zones with a sparse inflammatory infiltrate dominated a feature in the other. Signs of liver parenchyma regeneration were minimal in both animals. Two PEGC15S-injected animals showed moderate changes in the liver, with minor signs of parenchyma damage, but prominent regeneration, and were correctly recognized by the examiner as the ones receiving therapy. Changes indicative of liver parenchyma regeneration (slightly irregular architecture of the liver lobule, basophilic cytoplasm of hepatocytes, frequent mitoses and binucleated hepatocytes, and nuclear pleiomorphy and enlargement with prominent nuclei) were the most conspicuous feature. The overall morphology was consistent with that of increased proliferation and transcription in hepatocytes. Furthermore, rare focal necroses of hepatocytes and mild steatosis of a microvesicular type were detected, and fibrosis was minimal.

Discussion

In systems biology, the robustness of a biological system is defined by its ability to function properly in the face of perturbations. Redundancy of elements in the system is one of the mechanisms by which such robustness is achieved (24). The biological transsulfuration system seems to lack proper redundancy with its components and is therefore prone to mutational perturbations, as one of its elements, the CBS enzyme, is the only component in this system that can process homocysteine to cystathionine. A limited system redundancy partly exists, as homocysteine, the first metabolite that funnels into the pathway, can alternatively be converted to methionine through the remethylation pathway, thus relieving the homocysteine load. In addition, cysteine, a downstream product, can be obtained directly from the diet. Nevertheless, these pathways are limited in their capacity to maintain normal levels of metabolites, and lack of CBS activity manifests with detrimental consequences for patients if left untreated. Thus, a therapeutic approach for CBSDH must correct the perturbation rather than reinforce the alternative pathways. For most of the B6-nonresponsive CBSDH patients, the only current option is to reduce the intake of methionine by a strict low-methionine diet supplemented with cysteine in the amino acid mixture and by decreasing the homocysteine concentration with betaine supplementation. Currently, since gene therapy, the ultimate potential cure for metabolic diseases such as CBSDH, is still plagued with safety issues in the form of vector-induced genomic perturbations (insertional mutagenesis, genotoxicity) (25), ERT using htCBS seems to be the best hope for these patients to ease the symptoms of the disease while ameliorating the demanding dietary measures.

ERT requires chronic administration of the pharmacologically active substance, and thus lowered costs and suitability for self-administration are of importance. Therefore, s.c. administration is favored. However, as we show here (Figures 1 and 2), delivering htCBS via the s.c. route not only resulted in lower activity in the circulation compared with that seen with other routes, but, like what was observed with the other routes, showed almost no activity 24 or 48 hours after injection. Therefore, protein modification using PEG, a strategy that has been in the forefront of protein modification in the pharmaceutical industry for the past 25 years (20), was vital to achieve an acceptable retention time. Indeed, PEGylation of the htCBS enzyme improved the pharmacokinetics and, subsequently, the pharmacodynamic profile of the enzyme in vivo (Figures 1-3).

In healthy individuals, tHcy levels are in the range of approximately 5 to 15 μM (15), 98% of which is in the form of disulfides or is protein bound. Only 2% of the tHcy exists as a nonbound, free, reduced aminothiol that can serve as a substrate for CBS (26, 27). This balance is dramatically altered in CBSDH patients, with free, reduced homocysteine reaching 10%–25% of the tHcy values that are observed in these patients (up to ~400 μM) (28). In both cases, free, reduced homocysteine comprises only a fraction of tHcy. Nevertheless, the data presented in this study show that the presence of PEGylated htCBS in the circulation resulted in a 70–250 μM (70%–90%) decrease in tHcy levels (Figure 3). Thus, the initial levels of free homocysteine available for the enzyme (10%–25% of the total) cannot solely account for the decrease in tHcy levels. Therefore, additional pools of free homocysteine likely
become available to the enzyme, as the balance between bound and unbound homocysteine is affected upon PEGC15S administration. Alternatively, as free homocysteine becomes scarce as a result of the enzyme activity, the balance between free homocysteine and homocysteine adducts (in the form of protein-bound homocysteine or disulfides) changes to favor the generation of free homocysteine. Together, this implies that, upon administration of the PEGylated enzyme, the balance between bound and unbound homocysteine is shifted toward the free homocysteine that is processed by the enzyme in circulation. Thus, the PEGylated enzyme serves as a “sink” that constantly and systematically relieves the flux of homocysteine from extra- and intracellular pools.

It is noteworthy that even a reduction of 70% to 90% from the initial tHcy values was not sufficient to restore normal tHcy levels. This raises the question whether the significant drop in tHcy levels is enough to be of therapeutic utility in CBS DH patients. It was previously shown (29) that in mice with at least 54 times higher tHcy levels than those in WT mice, several phenotypic features were observed including facial alopecia, osteoporosis, reduction in mean survival, and other signs. However, no such signs were observed in mice that had tHcy levels of approximately 30 times the normal values. The authors concluded that homocysteine elevation is pathogenic only above a certain threshold level. Of particular interest are the results obtained in a multicenter observational study in humans (30), in which it was found that various treatment combinations failed to restore normal homocysteine values. The authors concluded that homocysteine levels in CBSDH patients and that B6 nonresponders had tHcy levels that were 6–8 times higher than the upper limit in the normal population. Nevertheless, these treatment strategies, achieving an imperfect biochemical control, significantly (relative risk, 0.09; P < 0.0001) reduced the risk of thromboembolism in these patients. Collectively, these data indicate that in order to ameliorate the clinical manifestation of CBSDH, a reduction of tHcy to below a threshold level, estimated at 80 to 100 μM, is sufficient.

A low-methionine diet can be effective in establishing and maintaining metabolic control in patients. This effect, however, is hindered by the lack of dietary compliance, particularly among late-diagnosed patients, resulting in a concomitant increase in homocysteine, accompanied by development of symptoms, especially in children (10). Betaine may help to achieve metabolic control in less compliant patients, and its combination with the diet may represent the best available treatment at present. However, current therapies for vitamin B₆ nonresponders do not increase cystathionine levels, and cysteine supplementation is warranted. In contrast, the murine data presented in this study show that administration of PEGhtCBS resulted in a significant reduction of tHcy levels, while the mice were maintained on a standard diet of 19% protein content (as opposed to the low-methionine diet used to treat human patients). This effect was far superior to the effect of betaine alone under these conditions, and combination of the enzyme with betaine showed that both can work synergistically to maintain low tHcy levels for an extended period of time (Figure 6). In addition to achieving metabolic control of homocysteine levels, administration of PEGhtCBS resulted in a significant increase in cystathionine levels and normalization of cysteine levels.

The role of cystathionine, other than serving as a substrate for CGL to produce cysteine, is unknown. Interestingly, CBS-KO mice suffer severe growth retardation and hepatopathy, and most die within 3 weeks after birth, even while the mothers are on betaine (21). The HO mice, which carry 2 copies of the human CBS gene, have high levels of tHcy but nevertheless suffer only minor consequences, if any. The only difference in metabolites between these 2 murine models is the level of cystathionine, which is significantly elevated in HO mice (~10 μM) as compared with that in KO mice (~1 μM), which may imply a positive role for cystathionine. Indeed, it has been suggested (31) that cystathionine may protect against ER stress-induced lipid accumulation, tissue injury, and apoptotic cell death. Moreover, high levels of cystathionine in brain tissues, along with rigorous control over CGL and CBS expression during mammalian development (32), are in line with an important role for cystathionine other than that of a precursor to cysteine. One may thus argue that the clinical manifestations of CBSDH may not only be attributed to high levels of the toxic homocysteine but, additionally, to the lack of cystathionine and cysteine and to elevated methionine levels (33). Low intracellular cystathionine levels may be corrected by uptake of this amino acid via a transporter mechanism such as the cysteine-glutamate transporter (34). Accordingly, the marked increase in cystathionine documented here following administration of PEGhtCBS may also be beneficial in ameliorating CBSDH symptoms.

Importantly, we observed normalization of cysteine levels following administration of the enzyme, which may present another advantage for CBS ERT. Cysteine biosynthesis from cystathionine is catalyzed solely by CGL. The latter is only present inside the cells and does not exist as a plasma protein (data not shown). Additionally, the extracellular cystathionine produced by the injected enzyme may also enter the cells (34) and serve as an intracellular substrate for CGL. In addition, it was recently suggested that a reduction of tHcy levels may impede the formation of cysteine-homocysteine adducts, which are rapidly cleared via urine excretion (35), or the formation of albumin-bound cysteines, thus resulting in higher levels of cysteine in plasma (35). Normalizing cysteine levels without cysteine supplementation may present another advantage of CBS ERT.

The positive effect of CBS administration was not limited to a change in metabolic levels in plasma and tissues, but showed a significant impact on liver disease and survival of the affected mice. It was previously shown that the majority of KO pups die within 2 weeks after birth and that growth retardation and severe liver disease are the most prominent outcomes of the null CBS condition (22, 36). This is in concert with the liver damage documented in this study for the PBS-injected KO animals and in sharp contrast to the CBS-treated mice, in which signs of liver regeneration substantially dominated over signs of liver parenchyma damage, indicating a capacity of the liver to cope with the metabolic defect.

The data presented in this study provide a proof of concept for PEGhtCBS as a promising ERT for homocystinuria, one that ameliorates the manifestations and metabolic abnormalities that characterize the disease. Clearly, translating a successful academic project into a therapy for human patients requires, among other elements, process upscaling, high-yield production, consistent batches, and pharmacokinetic and toxicological analyses in higher species. These studies were performed successfully and will be published elsewhere. In addition, the successful produc-
tion, approval, and use in humans of several recombinant enzyme products such as ADAGEN (ERT to treat adenosine deaminase deficiency and SCID), Pulmozyme (to treat cystic fibrosis), pegloticase (to treat gout), and more (37, 38), attests to the applicability of enzyme therapies in humans. Our proposed ERT thus provides a therapeutic strategy for CBSDH patients that allows for the prevention or treatment of symptoms and relaxes the need to maintain a strict diet.

Methods

Construction of an hCBS expression vector. A full-length (551 amino acids) hCBS coding sequence was optimized for bacterial expression and cloned into the pUC57 vector following digestion with the EcoRV restriction enzyme (GenScript USA). The hCBS sequence was then amplified by PCR using primers A1 and A2 (see Supplemental Experimental Procedures for the list of primers used) to generate a sequence coding for the truncated enzyme (amino acids 1–413). The PCR product was then digested with the restriction enzymes Ncol and Xhol and ligated into the pET-28a(+) vector that was digested with the same enzymes. Cloning into the Ncol site of pET-28a(+) results in a G→C mutation as compared with the native CBS sequence. A Site-Directed Mutagenesis Kit (Agilent Technologies) using primers B1 and B2 was used to restore the WT sequence. The same strategy was used to generate the C15S mutant by using primers C1 and C2. All sequences were verified by DNA sequencing.

Expression and purification. A detailed description can be found in the Supplemental Experimental Procedures. In short, E. coli BL-21 GOLD (DE3) (Agilent Technologies) or HMS174 (DE3) (EMD Millipore) was transformed with the CBS-encoding plasmid and grown in Terrific Broth. Protein expression was induced at OD600 = 0.6–0.7 with isopropyl β-D-thiogalactopyranoside (IPTG) and harvested and lysed 16 hours later. The lysate was then processed through a multistep chromatographic procedure.

PEGylation. Activated PEG molecules were purchased from NOF America, and PEGylation was carried out according to the manufacturer’s instructions. For example, coupling of maleimide PEGs to the SH groups of htCBS (5 mg/ml) was performed in a 100 mM phosphate buffer (pH 6.5) overnight at 4°C. The molar ratio between PEG and the CBS subunit was either 10:1 or 5:1.

Animals. C57BL/6j and CBS-KO mice were obtained from The Jackson Laboratory. HO mice were previously generated in our laboratory (21). Animals were maintained on the 2920X extruded standard diet (Envigo). Mice were routinely genotyped by using quantitative real-time PCR as outlined in the Supplemental Experimental Procedures. A single-use lancet for submandibular bleeding was used for blood collection into Capject T-MLHG lithium heparin (12.5 IU) tubes with gel (Terumo). Tubes were then centrifuged at 1,200 × g for 10 minutes, followed by collection of plasma into 1.5-ml tubes and storage at −80°C.

In-gel CBS activity staining and CBS activity assay. Protein samples were separated using Native Page (Bio-Rad). The gel was then incubated for 15 to 30 minutes in a staining solution (100 mM Tris-HCl, pH 8.0, 20 mM L-cysteine, 50 mM 2-ME, 0.1 mM PLP, and 0.2 mM lead nitrate). The reaction was stopped by submerging the gel in 7% acetic acid.

A detailed description of the CBS activity assay can be found in the Supplemental Experimental Procedures and in ref. 39. In short, CBS activity was determined by a radioisotope assay with 14C-labeled serine as a substrate. The 4C-cystathionine formed in the reaction was separated from the labeled substrate by overnight descending paper chromatography.

Determination of metabolite concentrations. Plasma metabolites were determined by stable-isotope dilution gas chromatography/mass spectrometry as previously described (40). Measurements of Hcy and nonprotein-bound homocysteine and other aminothiols as well as amino acids in tissue were performed as described previously (21, 41).

The analysis was performed in a blinded fashion, without knowledge of the animal treatment regimen.

SEC-HPLC. CBS preparations were separated on a Yarra SEC-3000, 300 × 7.8 mm SEC column (Phenomenex). The column was calibrated and operated in 100 mM sodium phosphate (pH 6.8), at a flow rate of 1 ml/min at room temperature.

Histology. Liver tissue sections were stained with H&E to evaluate the histopathological changes. Masson’s trichrome staining was performed for detection of fibrosis, and steatosis was verified using Oil Red O staining for detection of apolar lipids in fixed, frozen sections. A detailed description can be found in the Supplemental Experimental Procedures.

Statistics. All data are presented as the mean ± SEM and were compared using a paired or unpaired, 2-tailed Student’s t test. When comparing 3 different experimental arms, data were analyzed by ANOVA, followed by Tukey’s post-hoc analysis. A P value of less than 0.05 was considered significant. The log-rank test was used for comparison of survival curves, and significance was calculated using the χ² test.

Study approval. All animal procedures were approved by the IACUC of the University of Colorado Denver, which is an Association for Assessment and Accreditation of Laboratory Animal Care-accredited (AAALAC-accredited; no. 00235); Public Health Service-assured (no. A 3269-01); and US Department of Agriculture–licensed (USDA-licensed; no. 84-R-0059) institution.

Author contributions

EMB designed the studies, conducted the experiments, acquired and analyzed the data, and wrote the manuscript. TM generated preliminary data, provided reagents, participated in the design of the studies, acquired data, performed data analysis, and revised the manuscript. IP managed animal husbandry, assisted with the execution of animal studies, and prepared samples for analyses. RSC developed the purification procedure and provided reagents. HH performed the histological analysis of tissues and revised the manuscript. JK determined the levels of metabolites in tissues and revised the manuscript. VK coordinated the histological and metabolic analysis of tissues and revised the manuscript. JPK supervised the work, participated in the study design and analysis, and wrote the manuscript.

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