Hepatic pyruvate and alanine metabolism are critical and complementary for maintenance of antioxidant capacity and resistance to oxidative insult

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ABSTRACT

Objective: Mitochondrial pyruvate is a critical intermediary metabolite in gluconeogenesis, lipogenesis, and NADH production. As a result, the mitochondrial pyruvate carrier (MPC) complex has emerged as a promising therapeutic target in metabolic diseases. Clinical trials are currently underway. However, recent in vitro data indicate that MPC inhibition diverts glutamine/glutamate away from glutathione synthesis and toward glutaminolysis to compensate for loss of pyruvate oxidation, possibly sensitizing cells to oxidative insult. Here, we explored this in vivo using the clinically relevant acetaminophen (APAP) overdose model of acute liver injury, which is driven by oxidative stress.

Methods: We used pharmacological and genetic approaches to inhibit MPC2 and alanine aminotransferase 2 (ALT2), individually and concomitantly, in mice and cell culture models and determined the effects on APAP hepatotoxicity.

Results: We found that MPC inhibition sensitizes the liver to APAP-induced injury in vivo only with concomitant loss of alanine aminotransferase 2 (ALT2). Pharmacological and genetic manipulation of neither MPC2 nor ALT2 alone affected APAP toxicity, but liver-specific double knockout (DKO) significantly worsened APAP-induced liver damage. Further investigation indicated that DKO impaired glutathione synthesis and increased urea cycle flux, consistent with increased glutaminolysis, and these results were reproducible in vitro. Finally, induction of ALT2 and post-treatment with dichloroacetate both reduced APAP-induced liver injury, suggesting new therapeutic avenues.

Conclusions: Increased susceptibility to APAP toxicity requires loss of both the MPC and ALT2 in vivo, indicating that MPC inhibition alone is insufficient to disrupt redox balance. Furthermore, the results from ALT2 induction and dichloroacetate in the APAP model suggest new metabolic approaches to the treatment of liver damage.

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Keywords Acetaminophen; Diabetes; Glutathione; Liver; Metabolism; Oxidative stress

1. INTRODUCTION

Acetaminophen (APAP; a.k.a. paracetamol) is one of the most widely used antipyretic and analgesic drugs in the world. In a cross-sectional survey of the US population, for example, >20% of both adults and children reported use of APAP at least once in the preceding seven days [1,2] and similar data have been published from other countries [3–5]. However, APAP overdose can cause severe liver injury leading to acute liver failure (ALF) and death [6]. Oxidative stress is believed to be the central driver of APAP hepatotoxicity [7–9]. Cytochrome P450 enzymes (P450s), particularly CYP2E1, convert APAP to the reactive intermediate N-acetyl-p-benzoquinone imine (NAPQI) [10,11] which depletes the antioxidant glutathione and then binds to mitochondrial proteins [7,12–14]. The mitochondrial protein alkylation coincides with and likely causes reduced respiration [15] and an increase in mitochondrial reactive oxygen species (ROS) production [7,16]. The redox imbalance that results from combined glutathione depletion and increased ROS rapidly promotes phosphorylation and activation of the c-Jun N-terminal kinases (JNK) 1/2 [17,18], which translocate to mitochondria and further reduce mitochondrial respiration via SHP1-mediated inhibition of Src [19,20]. Other kinases are also activated by the oxidative stress and

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may exacerbate the injury [21–23]. Eventually, the mitochondrial membrane permeability transition occurs [24], mitochondria swell, and their membranes rupture [25]. The latter releases intermembrane and matrix proteins into the cytosol, including endonucleases which then cleave nuclear DNA [26]. Ultimately, the damaged hepatocytes die by necrosis [27,28]. Thus, anything that upsets mitochondrial metabolism or cellular redox regulation would be expected to alter APAP toxicity. The carbon backbone of pyruvate can enter the mitochondrial TCA cycle via either pyruvate carboxylation or oxidation and is an important substrate for gluconeogenesis, lipid synthesis, and NADH generation for the electron transport chain (ETC) [29]. Most pyruvate is produced in the cytosol as a product of glycolysis or conversion from lactate and is taken up by mitochondria via the mitochondrial pyruvate carrier (MPC) complex. Interestingly, in cultured cells and in hepatic tumors, genetic deletion of the MPC or pharmacological MPC inhibition not only blocks pyruvate oxidation [30,31], but also decreases glutathione synthesis and disrupts redox balance [32,33]. Thus, due to the role of mitochondrial dysfunction in APAP-induced liver injury and the critical resistance to APAP, we hypothesized that preventing mitochondrial pyruvate uptake by disrupting the MPC would worsen mitochondrial damage and oxidative stress after APAP overdose and therefore exacerbate APAP toxicity.

2. MATERIALS AND METHODS

2.1. Animals

WT male C57BL/6J mice between the ages of 7–8 weeks were purchased from the Jackson Laboratory (Bar Harbor, ME, USA) and used for experiments at 8–10 weeks of age. The generation of mice harboring a floxed allele of the gene encoding ALT2 (Gpt2) derived from Gpt2 germline KO mice generated by the Knockout Mouse Project Repository (KOMP) (project ID CSD24977) has recently been described [34]. Similarly, the generation of Mpc2 floxed mice using ES cells obtained from the European Conditional Mouse Mutagenesis Program (EUCOMM) (International Knockout Mouse Consortium [IKMC] project 89918, catalog no. MAE1900) has been described [35]. Transgenic mice expressing Cre under control of the albumin promoter (B6.Cg-Speer6-ps1Tg[Nb-cre][21Mip]/J; Jackson Laboratory stock number: 003574) were bred with these animals to create liver-specific Cre mice. Combined liver-specific Gpt2/Mpc2 double KO (DKO) mice were then generated by intercrossing the two liver-specific gene knockouts. Littermate mice not expressing Cre (fl/fl) mice were used as controls. Male mice were used in all experiments because female mice are resistant to APAP hepatotoxicity [36]. All Mpc2 KO and DKO mice and their WT littermate controls were used at 10–16 weeks of age. All Gpt2 (ALT2) KO mice were used between 14 and 28 weeks of age. Animals were kept in a temperature-controlled 12 h light/dark-cycle room with ad libitum access to food and water. For BCLA time course and dose–response experiments, fed WT mice were treated with BCLA dissolved in 1× phosphate-buffered saline (PBS) at the indicated doses or PBS alone and liver tissue was collected at the indicated time points. For APAP experiments, mice were fasted overnight, then treated i.p. with 300 mg APAP per kg body mass (mg/kg) dissolved in PBS. Some WT mice were pre-treated i.p. with the specified doses of BCLA or PBS or with 100 mg/kg dexamethasone (Dex) for the indicated times before APAP. Others received 30 mg/kg MSDC, 318 mg/kg NAC, or appropriate vehicle 2 h after APAP. NAC and NAC vehicle (0.9% saline) were administered i.v. All other treatments were given i.p. Blood and liver tissue were collected at 6 h post-APAP.

Some liver pieces were fixed in 10% phosphate-buffered formalin for histology, while others were flash-frozen in liquid nitrogen and stored at –80 °C until biochemical analyses. All study protocols were approved by the Institutional Animal Care and Use Committees of the University of Arkansas for Medical Sciences and Washington University in St. Louis. The Animal Research: Reporting of In Vivo Experiments (ARRIVE) guidelines were consulted during manuscript preparation and we have attempted to conform to those recommendations.

2.2. Biochemistry

Alanine aminotransferase (ALT) was measured in serum using a kit from MedTest Dx, Inc. (Canton, MI, USA), according to the manufacturer’s instructions. Lactate dehydrogenase (LDH) was measured in serum as previously described [37]. For hepatic ALT activity, frozen liver tissue was homogenized in 25 mM HEPS buffer with 5 mM EDTA (pH 7.4), 0.1% CHAPS, and protease inhibitors using a bead homogenizer (Thermo Fisher Scientific, Waltham, MA). After centrifugation at 10,000 × g and 4 °C for 5 min, ALT was measured in the supernatant using the MedTest kit mentioned above. Protein concentration in the lysates was measured using the bicinchoninic acid (BCA) assay and was used to normalize hepatic ALT activity. Ketones were measured using the EnzyChrom Ketone Body Assay (BioAssay Systems, Hayward, CA USA) according to the manufacturer’s instructions.

2.3. Cell culture

Primary mouse hepatocytes were isolated from WT mice using a collagenase perfusion technique, as previously described [38], and used in experiments no more than 24 h after isolation. HepG2 cells overexpressing CYP2E1 were purchased from Hera Bioblas (Lexington, KY USA) and grown in Williams’ Medium E with 5% fetal bovine serum, glutamate, penicillin/streptomycin, and insulin. Regular HepG2 cells were purchased from the American Type Culture Collection (Manassas, VA USA) and grown in the same medium. Cells were treated as described in the results section.

2.4. Histology

Formalin-fixed tissue was embedded in parrafin wax and 5 μm sections were mounted on glass slides for hematoxylin and eosin (H&E) staining. H&E staining was performed using a standard protocol. Necrosis was quantified by two independent, blinded, fellowship-trained hepatobiliary pathologists. Slides were imaged using a Labomed Lx400 microscope with a 16 MP Atlas digital camera and Pixel Pro software (Laba America, Fremont, CA USA).

2.5. Western blotting

Western blotting was performed as we previously described [39]. The primary antibodies against phospho-JNK (Cat. No. 4668), JNK (Cat. No. 9252), and Mpc2 (Cat. No. 46141) were purchased from Cell Signaling Technology (Danvers, MA, USA) and used at 1:1,000 dilution in TBS-T with 5% milk. The Cyp2e1 antibody was purchased from Proteintech (Rosemont, IL, USA) (Cat. No. 19937-1-AP). The ALT2 antibody was from Sigma (St. Louis, MO, USA) (Cat. No. HPA051514). Secondary antibodies were purchased from Li-Cor Biotechnology (Lincoln, NE, USA) and Cell Signaling Technology. Total protein was stained with Coomassie blue. All blots were imaged using a Li-Cor Odyssey instrument (phospho-JNK and JNK) or an Invitrogen iBright FL1500 (all other blots and protein images).

2.6. Plasma metabolomics for amino acids and pyruvate

Amino acids were extracted from 20 μL of plasma with 200 μL of methanol containing Try-d8 (1600 ng), Phe-d8 (160 ng), Tyr-d4 (160 ng), Leu-d3 (320 ng), Ile-13C6,15N (160 ng), Met-d3 (160 ng),
Val-d8 (400 ng), Glu-d3 (2400 ng), Thr-13C4 (320 ng), Ser-d3 (400 ng), Asp-d3 (3200 ng), Ala-d4 (1600 ng), Asn d3,15N2 (3200 ng), Gly-d2 (16000 ng), Gln-13C5 (3200 ng), Pro-d7 (800 ng), Cit-d4 (80) ng, His 13C6 (400 ng), Arg-13C6 (1600 ng), Orn-13C5 (400 ng), Lys-d8 (400 ng), and pyruvic acid-13C5 (80 ng) as the internal standards for Try, Phe, Tyr, Ile, Met, Val, Glu, Thr, Ser, Asp, Ala, Asn, Gly, Gln, Pro, Cit, His, Arg, Orn, Lys, and pyruvic acid (Pyr) respectively. The sample aliquots for pyruvic acid were derivatized with o-phenylenediamine to improve mass spectrometric sensitivity. Quality control (QC) samples were prepared by pooling aliquots of study samples and injected after every four study samples to monitor instrument performance throughout these analyses.

The analysis was performed on a Shimadzu 20AD HPLC system and a SIL-20AC autosampler coupled to 4000qtrap mass spectrometer (AB Sciex) operated in positive multiple reaction monitoring (MRM) mode. Data processing was conducted with Analyst 1.6.3 (Applied Biosystems). The relative quantification data for all analyses were presented as the peak ratio of each analyte to the internal standard. The Metabolomics Facility of Washington University School of Medicine performed the analysis.

2.7. qPCR
RNA was extracted from liver tissue using the RNeasy Mini Kit (Qiagen, Venlo, Netherlands), mRNA levels for Glicc (Fwd, 5' to 3'; ATC TGC AAA GGC GGC AAC; Rev, 5' to 3'; ACT CCT CTG CAG CTG CTC), Gsr (Fwd, 5' to 3'; ATG CCT ATG TGA GCC GCC TG; Rev, 5' to 3'; TGC GAA TGT TGC ATA GCC GT), Gpd2 (Fwd, 5' to 3'; AAT CTA CTG AGG TAA TCC GA; Rev, 5' to 3'; GGG CTG TTT AGT AGG TTT GGG TA) and Mpc2 (Fwd, 5' to 3'; ACC TAC CCA CGA CTC ATG GAT; Rev, 5' to 3'; TGT AAA GCG GCC TCA ATT TCT) were measured by real-time quantitative PCR using the Applied Biosystems PowerSYBR Green PCR Master Mix (Thermo Fisher, Waltham, MA USA) according to the manufacturer’s instructions.

2.8. Statistical analyses
Sample sizes for the pharmacologic treatment experiments were determined a priori and are indicated in the caption for each figure. In each case, the difference in serum ALT between the vehicle control and drug-treated groups was the primary comparison of interest and the experiments were designed to achieve a minimum of 80% power at α = 0.05 using our historical ALT data to predict variation. Sample sizes for the KO and DKO animals were determined by litter size, frequency, and availability for this project. Secondary analyses were carried out on the same samples. Statistical significance was assessed using either Student’s t-test or one-way analysis of variance (ANOVA) with post-hoc Dunn’s test or Tukey’s test. For non-normally distributed data, the Mann–Whitney U-test or the Kruskal–Wallis test were used. All statistical analyses were performed using SigmaPlot 12.5 (Systat, San Jose, CA, USA) or Prism 10.0 (GraphPad, Boston, MA, USA). P-values are indicated in the figures or figure captions.

3. RESULTS
3.1. MPC inhibition and deletion have no effect on APAP hepatotoxicity
Prior work has suggested that enhanced hepatic pyruvate metabolism after APAP overdose may have hepatoprotective effects. For example, genetic deletion of pyruvate dehydrogenase kinase 4 (PDK4), which inhibits mitochondrial pyruvate oxidation, protected mice from APAP-induced injury [40]. To further test the effects of enhancing pyruvate oxidation, we post-treated mice with either vehicle or the PDK inhibitor dichloroacetate (DCA) 2 h after APAP in order to increase pyruvate flux through the pyruvate dehydrogenase complex (PDHC). Consistent with the prior work using PDK4 KO mice, DCA reduced APAP hepatotoxicity as indicated by serum ALT (Figure 1A). Given this finding, one might predict that loss of mitochondrial pyruvate metabolism would exacerbate APAP-induced liver injury. The mitochondrial pyruvate carrier (MPC) catalyzes the transport of pyruvate into the mitochondrial matrix and is composed of two proteins (MPC1 and MPC2) that form a heterodimeric complex in the inner mitochondrial membrane [29]. Thus, we treated liver-specific MPC2 KO mice with APAP to determine whether loss of mitochondrial pyruvate metabolism might affect the response to APAP mediated hepatotoxicity. Consistent with prior reports [35,41], MPC2 deletion resulted in instability and degradation of MPC1 (Figure 1B), effectively leading to complete loss of the MPC complex. However, MPC deficiency had no effect on APAP hepatotoxicity at 6 h after treatment, within the timeframe when mitochondrial metabolism is most critical [7,8]. There were no significant differences in serum ALT or hepatic necrosis, total glutathione, oxidized glutathione, APAP-protein binding, or JNK activation between MPC2 KO mice and WT littermate controls (Figure 1C–K).

Although uptake via the MPC is the primary source of mitochondrial pyruvate, we have previously shown that cytosolic and mitochondrial alanine-pyruvate cycling catalyzed by alanine transaminase enzymes can partially circumvent the loss of MPC activity [35]. Indeed, shRNA-mediated knockdown of alanine transaminase 2 (ALT2), which catalyzes the transamination of alanine and α-ketoglutarate to form pyruvate and glutamate in the mitochondrial matrix, further attenuates mitochondrial pyruvate flux in MPC2-deficient hepatocytes [35]. In the present study, we found that ALT2 protein abundance was 1.9 ± 0.2-fold greater in livers from MPC2 KO mice compared to WT mice after APAP treatment (Figure 1B and I), suggesting that mitochondrial alanine-pyruvate cycling might be compensating for loss of the MPC during APAP hepatotoxicity as well.

Though we cannot rule out that injury may have also contributed to ALT2 induction at this point, the data induction of ALT2 in APAP toxicity may have been protective either way.

3.2. ALT2 inhibition and deletion have no effect on APAP hepatotoxicity
The idea that increased ALT2 might have prevented increased injury in the MPC2 KO mice led us to determine if increasing mitochondrial alanine flux is protective against APAP-induced injury more generally. We pre-treated WT mice for 24 h with a large dose of dexamethasone (Dex) followed by APAP. Large doses of Dex are known to strongly induce ALT2 expression in the liver [42,43]. Importantly, western blotting confirmed an increase in hepatic ALT2 with Dex treatment and the gold-standard for injury, histology, revealed that Dex administration was indeed protective (Figure 2A–O).

We next used a pharmacological approach to suppress mitochondrial alanine metabolism. β-chloro-α-alanine (BCLA) is an alanine analog that inhibits the enzyme’s activity [44]. Although it has been extensively used in studies in vitro, we are not aware of any data in the literature characterizing the dose—response and kinetics of BCLA inhibition of ALT activity in the liver in vivo. Thus, we first performed timing and dose—response studies to optimize dosing. The results revealed that BCLA can inhibit ALT activity in the liver in mice, that the effect is greater at 24 h post-BCLA treatment than an earlier time point, and that >50% inhibition can be achieved with a dose of 100 mg/kg (Suppl. Fig. 1). Based on these data, we adopted a 24 h single-dose (Figure 2D–H) and 48 h two-dose pre-treatment regimen with 100 mg/kg BCLA to study the role of ALT in APAP toxicity. Consistent
Neither MPC inhibition nor liver-specific knockout worsen liver injury or glutathione depletion after APAP overdose. (A) WT male mice 8–10 weeks old were treated with 300 mg/kg APAP followed by 250 mg/kg dichloroacetate (DCA) or PBS vehicle (Veh) 2 h later. Serum ALT was measured at 6 h post-APAP. (B–K) Male livers-specific MPC KO mice and WT littermates 10–16 weeks old were treated with 300 mg/kg APAP or PBS vehicle. Serum and liver tissue were collected 6 h later. (B) MPC1, MPC2, β-actin, and ALT2 were measured in liver homogenates by immunoblot. (C) Serum ALT was measured. (D) 5 μm-thick liver sections were glass-mounted and stained with hematoxylin and eosin (H&E) to measure necrosis. Representative images are shown at 100× magnification. (E) Necrosis was quantified in the liver sections. (F) Total glutathione (GSH + GSSG) was measured in liver tissue. (G) Oxidized glutathione was measured in liver tissue and expressed as a proportion of the total glutathione. (H) APAP-protein adducts were measured in liver tissue. (I) Densitometric analysis was performed on the ALT2 immunoblot in (B). (J) Phosphorylated and total JNK (p-JNK and JNK, respectively) were measured in liver homogenates by immunoblot. (K) Densitometry of K. Comparison of two means was performed by two-tailed t-test. Comparison of three or more groups was performed by ANOVA with post-hoc Tukey’s test for multiple comparisons. *p < 0.05, ****p < 0.0001. In all panels, lines and error bars indicate mean ± SE. Circles show individual data points and sample sizes. Box plots show medians and IQR. Open circles: Veh. Filled circles: APAP.
Figure 2: Neither ALT2 inhibition nor ALT2 knockout worsen liver injury after APAP overdose. (A–C) WT male mice 8–10 weeks old were treated with 100 mg/kg dexamethasone (Dex) followed by 300 mg/kg APAP 24 h later. Liver tissue was collected 6 h post-APAP. (A) ALT2 induction in liver homogenates was measured by immunoblot. (B) 5 μm-thick liver sections were glass-mounted and stained with hematoxylin and eosin (H&E) to measure necrosis. Representative images are shown at 100× magnification. (C) Necrosis was quantified in the liver sections. (D–H) WT male mice 8–10 weeks old were treated with 100 mg/kg BCLA followed by 300 mg/kg APAP 24 h later. Serum and liver tissue were collected 6 h post-APAP. ALT activity was measured in (D) serum and (E) liver homogenates. (F) 5 μm-thick liver sections were glass-mounted and stained with H&E to measure necrosis. Representative images are shown at 100× magnification. (G) Necrosis was quantified in the liver sections. (I–M) Male liver-specific ALT2 KO mice and WT littermate controls 14–28 weeks old were treated with 300 mg/kg APAP or PBS vehicle. Serum and liver tissue were collected at 6 h post-APAP. (I) ALT2 was measured in liver homogenates by immunoblot. (J) Serum ALT was measured in APAP and PBS treated WT and KO mice. (K) Serum LDH was measured in APAP and PBS treated WT and KO mice. (L) 5 μm-thick liver sections were glass-mounted and stained with H&E to measure necrosis. Representative images are shown at 100× magnification. (M) Necrosis was quantified in the liver sections. Comparison of two means was performed by two-tailed t-test. Comparison of three or more groups was performed by ANOVA with post-hoc Tukey’s test for multiple comparisons. *p < 0.05, ***p < 0.001. In all panels, lines and error bars indicate mean ± SE. Circles show individual data points and sample sizes. Box plots show medians and IQR.
with its action as an ALT inhibitor. BCLA reduced both serum and hepatic ALT activity after APAP overdose (Figure 2D and E). However, like MPC2 deletion, BCLA had no effect on injury as indicated by the alternative serum liver injury biomarker LDH and by hepatic necrosis (Figure 2F–H). Results from the two-dose 48 h BCLA pre-treatment regimen were nearly identical (data not shown).

BCLA did not completely block hepatic ALT activity (Figure 2E). In addition, it is possible that the drug cannot reach ALT2 within the mitochondrial matrix. Thus, to ensure complete inhibition of ALT2, we next used liver-specific ALT2 KO mice (Figure 2I). However, like MPC2 KO, there was no significant effect of ALT2 deletion on APAP-induced liver injury, as indicated by serum ALT (Figure 2J) and LDH (Figure 2K) and by hepatic necrosis (Figure 2L,M). Although it may seem surprising that ALT2 KO itself did not reduce serum ALT activity, this observation is consistent with reports that ALT1 is the dominant isoform released into serum [45–47], accounting for ≥90% of ALT activity in circulation after injury [47]. Thus, we continued to use serum ALT as our primary measure of liver injury. We also confirmed all ALT results with histological analysis. Altogether, these data indicate that neither MPC2 deletion nor ALT2 deficiency alters APAP hepatotoxicity but do not rule out the possibility of complementary effects.

3.3. Combined MPC2/ALT2 deficiency worsens APAP-induced liver injury

To determine if deletion of both MPC2 and ALT2 worsened APAP hepatotoxicity, we generated liver-specific ALT2/MPC2 double KO (DKO) mice (Suppl. Fig. 2). In contrast to mice with only single deletion of MPC2 or ALT2, DKO mice had greater serum ALT activity and hepatic necrosis after APAP overdose (Figure 3A–C). Importantly, they also had lower total hepatic glutathione levels (Figure 3D). Glutathione reaches its nadir with nearly complete loss by approximately 0.5 h post-APAP and then recovers over the following 6 h [12]. Thus, these 6 h data suggest impaired glutathione re-synthesis, possibly due to the diversion of glutamate to α-ketoglutarate (αKG) and away from glutathione synthesis as we had initially hypothesized would happen with MIP inhibition. Furthermore, these effects occurred despite less APAP-protein binding in the DKO mice (Figure 3E). Because protein alkylation is the initiating event in APAP hepatotoxicity, even greater injury may have been observed had the level of protein binding been the same as in WT. Indeed, when we normalized serum ALT to hepatic APAP-protein adducts, the difference in means was even greater with a 4.9 ± 0.2-fold increase in ALT in the DKO animals (mean ± SE) (Figure 3F).

The reason for the reduced adducts in DKO animals may have been depletion of NADPH, which is necessary for the P450 catalysis that produces NAPQI because the P450 catalytic cycle uses electrons from NAPDH transferred via the enzyme P450 reductase. The protein levels of CYP2E1 — the primary enzyme that converts APAP to NAPQI [48] — were the same between genotypes (Fig. G,H), demonstrating that the lower NAPQI production and protein binding was not due to altered CYP2E1 expression. Furthermore, lower NADPH levels have been reported in MPC1-deficient cells in vitro [53] and we observed NADPH depletion in our DKO animals after APAP overdose as well (Figure 3I,J).

3.4. The mechanism of exacerbated injury in DKO mice may be related to diversion of glutamate away from glutathione

Compensatory glutaminolysis at the expense of glutathione synthesis has been previously reported in MPC1-deficient hepatocytes in vitro [32] and in other conditions in which mitochondrial pyruvate metabolism was impaired [49]. Thus, we compared basal total hepatic glutathione between DKO and WT mice and between HepG2 cells treated with the combination of another MIP inhibitor, UK-5099, and BCLA or vehicle to mimic combined MPC and ALT2 KO. Interestingly, glutathione was 22 ± 5% (mean ± SE) lower in liver tissue from DKO animals (Figure 4A) and almost undetectable in the UK/BCLA co-treated cells (Figure 4B). Targeted plasma metabolomics also revealed elevated urea cycle intermediates (citrulline, arginine, and ornithine) in the untreated DKO mice (Table 1). While steady state metabolite concentration is not equivalent to metabolic flux, the results are consistent with increased disposal of ammonia from glutamate and other amino acids that are being utilized to compensate for loss of pyruvate and alanine entry into the TCA cycle. Unfortunately, urea, potentially a more direct endpoint for urea cycle activity, could not be measured in these samples because APAP overdose also causes kidney damage, resulting in blood urea concentrations 2–3-fold greater than control values [50]. It would be difficult to disentangle the contributions of the urea cycle and kidney damage to elevated urea levels. Alternatively or in addition to increased ammonia disposal, altered expression of glutathione synthesizing or metabolizing enzymes could explain glutathione depletion in the DKO animals. To address these possibilities, we measured mRNA for glutamate cysteine ligase (Gcl), the rate-limiting enzyme in glutathione synthesis, and for glutathione reductase (Gsr) in liver tissue from WT, MPC2 KO, ALT2 KO, and DKO mice. Importantly, there were no differences in the expression of either between any of the genotypes (Suppl. Fig. 3).

Importantly, post-treatment of cells with UK/BCLA after glutathione depletion with buthionine sulfoximine (BSO) completely prevented glutathione re-synthesis over the next 24 h (Figure 4C). Treatment with 500 μM BSO for 24 h almost fully depleted glutathione. Cells post-treated with Veh control for an additional 24 h after removal of BSO recovered approximately 30% of control glutathione content, but cells post-treated with the combination of 25 μM UK-5099 and 250 μM BCLA instead of vehicle failed to re-synthesize any (Figure 4C). Furthermore, APAP toxicity was dramatically increased in HepG2 cells overexpressing CYP2E1 when APAP was co-administered with UK and BCLA (Figure 4D,E), similar to the DKO mice. Importantly, CYP2E1-overexpressing HepG2 cells were previously validated for the study of APAP hepatotoxicity, with formation of APAP-protein adducts, mitochondrial damage, and cell death occurring similar to primary mouse and human hepatocytes [51], and our preliminary dose—response and time course data were consistent with those earlier results (Suppl. Fig. 5).

Collectively, the data presented are consistent with increased conversion of glutamate and other amino acids to α-keto acids to feed the TCA cycle, leading to reduced glutamate availability for glutathione synthesis.

3.5. NAC still protects in MPC2 KO and DKO mice

NAC is currently the standard-of-care treatment for APAP overdose in humans. It is generally believed to protect against APAP hepatotoxicity by providing cysteine for re-synthesis of glutathione which can then scavenge NAPQI and ROS. However, the doses of NAC given to APAP overdose patients (140 mg/kg loading, followed by additional lower doses) greatly exceed those required to replenish glutathione. In addition, it has been reported that similar doses can increase flux through the PDHC in mice [52]. It has therefore been proposed that large doses of NAC like those administered clinically protect against APAP in part by increasing pyruvate oxidation [52,53] — an effect that would require the MPC. To test this hypothesis more directly than in prior work, we examined NAC protection in DKO mice and WT littermate controls. Interestingly, NAC protected equally well in two genotypes (Figure 5A), despite using the same large dose (318 mg/kg) employed in the prior work [53]. To our surprise, NAC also supported glutathione re-synthesis in the DKO mice similar to the WT animals (Figure 5B). These data suggest that while excess NAC may or may not
increase PDHC flux, this is not required for NAC-mediated protection after APAP hepatotoxicity.

4. DISCUSSION

There is much interest in the development of MPC-targeted drugs to treat metabolic conditions in humans, including diabetes and liver disease [54,55], with some early-stage clinical trials completed [56] and a phase 3 trial currently underway (ClinicalTrials.gov ID: NCT03970031). Because pyruvate can be converted to glucose through a series of steps that begins with conversion to oxaloacetate within mitochondria, inhibiting mitochondrial pyruvate uptake could reduce hyperglycemia and its downstream adverse effects. Indeed, obesity- and streptozotocin-induced hyperglycemia is attenuated in mice with liver-specific MPC KO and after MPC inhibitor treatment [35,57]. MPC inhibitors also hold promise in treating nonalcoholic steatohepatitis (NASH), as MPC deletion or inhibition protects mice from liver injury and fibrosis in preclinical NASH models [57–59]. However, given the central
Figure 4: Combined inhibition of the MPC and ALT2 reduces glutathione synthesis in vivo and in vitro. (A) Basal total glutathione was measured in liver tissue homogenates from untreated male 10–16 weeks old liver-specific MPC2/ALT2 DKO mice. (B) HepG2 cells were treated with 25 μM UK-5099 and 250 μM BCLA or DMSO vehicle (Veh) control for 24 h and total glutathione was measured. (C) HepG2 cells were treated with either vehicle or 500 μM BSO for 24 h and total glutathione was measured. Other HepG2 cells were treated with 500 μM BSO for 24 h followed by vehicle or the combination of 25 μM UK-5099 and 250 μM BCLA for another 24 h and total glutathione was measured in the cells. (D–E) HepG2 cells overexpressing CYP2E1 were co-treated with vehicle, 25 μM UK-5099 and 250 μM BCLA, APAP and vehicle, or APAP and 25 μM UK-5099 and 250 μM BCLA for 48 h. (D) Serum LDH release was measured in the cells. (E) Dead cells were stained with propidium iodide and the cells were imaged by either phase contrast or fluorescence imaging (red fluorescence indicates propidium iodide positivity). Scale bars: 200 μm for 100× magnification, 50 μm for 400×. Comparison of two means was performed by two-tailed t-test. Comparison of three or more means was performed by ANOVA with post-hoc Tukey’s test for multiple comparisons. ****p < 0.0001. In all panels, lines and error bars indicate mean ± SE. Circles show individual data points and sample sizes. Box plots show medians and IQR. For the cell culture studies (B–E), each data point shows the mean of multiple technical replicates from one experiment. Experiments were performed across multiple days.
importance of pyruvate in mitochondrial intermediary metabolism, there are valid concerns regarding the safety of MPC inhibition. For example, alterations in glutathione synthesis and redox status that have been observed in cell culture studies [32,33] could predispose patients to oxidative insults [55]. The cause of these changes in glutathione during MPC inhibition appears to be an increase in glutaminolysis (conversion of glutamine to α-KG through glutamate). Entry of amino acids, particularly alanine and glutamine/glutamate, into the TCA cycle can compensate for MPC deficiency [35, 60–62]. However, increased TCA flux of glutamine and glutamate could also lead to decreased glutamate availability for glutathione synthesis, which could be detrimental in some contexts. Indeed, two recent studies have demonstrated that MPC inhibition leads to impaired glutathione synthesis and recycling of oxidized glutathione in cultured cells [32,33]. Interestingly, however, these effects were not observed in liver tissue from MPC1 KO mice in vivo [32]. Based on our data, the explanation for this discrepancy between cells and mice is likely compensation for loss of MPC activity by ALT2 in the animals. ALT2 was roughly 2-fold elevated in our MPC2 KO mice and there was no evidence of worsened glutathione depletion in those animals after APAP overdose. However, when both MPC2 and ALT2 were absent, hepatic glutathione decreased at baseline compared to WT mice and re-synthesis after APAP-induced depletion was severely impaired. Furthermore, the phenotype we observed in cells treated with both UK-5099 and the ALT inhibitor BCLA was much more severe than in prior studies using the MPC inhibitor alone. Altogether, our results suggest that loss of both the MPC and ALT2 is required to alter glutathione synthesis in hepatocytes in vivo. Thus, MPC inhibitors are likely insufficient to sensitize patients to oxidative insults, except possibly in rare cases of individuals with one of several recessive Gpi2 polymorphisms that drastically reduce enzyme activity [63,64]. Although one might think fatty acid oxidation (FAO) can serve as a source of acetyl-CoA for the TCA cycle and potentially also compensate for MPC loss, FAO is inhibited in APAP toxicity [65–68] resulting in lower ketones (Suppl. Fig. 6) among other effects and there was no difference in serum ketones between DKO and WT mice in the present study (Suppl. Fig. 6). Furthermore, FAO cannot serve as an anaplerotic substrate for the TCA cycle to support anabolic functions like lipogenesis or gluconeogenesis. It is therefore unlikely that FAO can fully compensate for impaired mitochondrial pyruvate and alanine metabolism during APAP-induced injury.

Another novel finding from our study is that inducing expression of ALT2 with a large dose of dexamethasone can reduce APAP-induced liver injury. The mechanism of protection with ALT2 induction could be enhanced glutathione re-synthesis due to the glutamate/glutamine-sparing effect of pyruvate, as similar large doses of other glucocorticoids have been shown to increase hepatic glutathione [69]. Although treatment of liver damage was not the focus of this study and 24 h pre-treatment with Dex is not clinically feasible in the context of acute APAP overdose, this result adds to a growing body of literature indicating that improving pyruvate entry and flux in the TCA cycle is protective against liver injury. For example, it was recently demonstrated that mice deficient in PDK4, which phosphorylates and inhibits the PDHC, had much lower liver injury and oxidative stress after APAP overdose than WT controls [40]. This finding is consistent with the present data showing that a chemical inhibitor of PDK activity, DCA, also reduced liver injury after APAP. Importantly, greater glutathione re-synthesis was observed in the PDK4-deficient mice [40], which is consistent with a glutathione-sparing effect of increased pyruvate oxidation. Another group reported that exogenous pyruvate and ethyl pyruvate are also protective against APAP and NAPQI toxicity in mice and HepG2 cells, respectively [70]. Thus,
targeting pyruvate metabolism may be a promising novel approach to treat APAP overdose in patients in combination with NAC. The primary mechanism by which NAC protects against APAP hepatotoxicity is providing cysteine for the rapid re-synthesis of glutathione [14,71,72]. However, it is often claimed that NAC is beneficial in APAP overdose patients even when administered late after overdose, when APAP bioactivation and ROS production have already occurred [73,74]. There is also modest evidence that NAC is beneficial for patients with non-APAP acute liver damage, presumably unaccompanied by glutathione depletion [75]. An alternative hypothesis to explain how NAC protects in these scenarios is via increased pyruvate oxidation. For example, Zwingmann and Bilodeau demonstrated that NAC increases glucose flux through the PDHC and TCA cycle and protects against oxidative stress even in glutathione-replete mice [52]. The mechanism by which NAC had this effect was unclear from their study but could have been conversion of the excess cysteine to pyruvate. It was later demonstrated that when equimolar doses of NAC and the tripeptide glutathione were administered to mice separately after APAP overdose, glutathione was more effective than NAC at reducing liver injury and it increased TCA cycle flux to a greater degree [53]. Furthermore, increasing the NAC dose 3-fold to provide the same number of amino acids as glutathione equalized their effects [53]. Based on this, it was concluded instead that amino acids derived from NAC are used to synthesize TCA cycle substrates which then enhance TCA cycle flux [53]. This latter hypothesis is consistent with our observation that the same large dose of NAC used in those studies protected equally well in WT and DKO mice that cannot use cytosolic cysteine-derived pyruvate. In addition, the fact that NAC also restored hepatic glutathione in DKO mice is consistent with the possibility that some excess cysteine was instead converted to glutamate, which could then be used in both glutathione re-synthesis and the TCA cycle. However, a definitive mechanism by which NAC protects late after APAP overdose or in glutathione-replete subjects cannot be determined from our data and additional experiments are needed.

Finally, it is notable that ALT activity has been the primary serum biomarker of liver injury since 1955 [76,77] but its metabolic and physiological roles in liver injury and disease are rarely considered, even though it is an important metabolic enzyme within the tissue [77]. Interestingly, it was recently reported that hepatic ALT1 and ALT2 contribute to hyperglycemia in diabetic mice by converting alanine to pyruvate which is then used in gluconeogenesis [34,78], providing possibly the first evidence that ALT2 has pathophysiological significance in liver-related metabolic diseases. The current study adds to our emerging awareness that ALT is more than a passive biomarker of injury by demonstrating its complementary role with the MPC in maintenance of mitochondrial intermediary metabolism and preservation of redox balance.

5. CONCLUSIONS

Overall, our data suggest that hepatocytes have metabolic redundancies to preserve TCA cycle flux and defend against oxidative stress even in the context of impaired mitochondrial pyruvate metabolism (see Graphical Abstract), which is an important metabolic substrate of the TCA cycle. However, if all sources of mitochondrial pyruvate are impeded, hepatocytes switch to metabolism of amino acids to maintain flux at the cost of reduced synthesis of the antioxidant glutathione. As the amino acid glutamate is shunted into the TCA cycle, less is available for glutathione synthesis, rendering hepatocytes more susceptible to oxidative insult. Conversely, stimulation of mitochondrial pyruvate flux was protective. These findings have implications for the safety and future development of MPC inhibitors to treat metabolic diseases and for future treatment of APAP hepatotoxicity.

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AUTHOR CONTRIBUTIONS


DECLARATION OF COMPETING INTEREST

LPJ is Chief Medical Officer of Acetaminophen Toxicity Diagnostics (ATD), LLC, which is developing a diagnostic test for APAP overdose. MR McGill is a consultant for ATD and has received research funding from GlaxoSmithKline which sells APAP. BNF is a shareholder and member of the scientific advisory board of Cirius Therapeutics, which is developing an MPC inhibitor to treat nonalcoholic steatohepatitis. The remaining authors declare that they have no conflicts related to the content of this article. These companies had no role whatsoever in the funding, conceptualization, design, execution, or reporting of the study.

DATA AVAILABILITY

Data will be made available on request.

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ABBREVIATIONS

ALF acute liver failure
ALT alanine aminotransferase
APAP acetaminophen
Dex dexamethasone
DKO double ALT2/MPC knockout
GSH reduced glutathione
GSSG oxidized glutathione
JNK c-Jun N-terminal kinase
KO knockout
LDH lactate dehydrogenase
LDH lactate dehydrogenase
MPC  mitochondrial pyruvate carrier
NAPQI N-acetyl-p-benzoquinone imine
P450  cytochrome P450
PDHC pyruvate dehydrogenase complex
PKD4 pyruvate dehydrogenase kinase 4
TCA tricarboxylic acid

APPENDIX A. SUPPLEMENTARY DATA

Supplementary data to this article can be found online at https://doi.org/10.1016/j.molmet.2023.101808.

REFERENCES


Original Article


