Methanobactin reverses acute liver failure in a rat model of Wilson disease

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In Wilson disease (WD), functional loss of ATPase copper-transporting β (ATP7B) impairs biliary copper excretion, leading to excessive copper accumulation in the liver and fulminant hepatitis. Current US Food and Drug Administration– and European Medicines Agency–approved pharmacological treatments usually fail to restore copper homeostasis in patients with WD who have progressed to acute liver failure, leaving transplantation as the only viable treatment option. Here, we investigated the therapeutic utility of methanobactin (MB), a peptide produced by *Methylosinus trichosporum* OB3b, which has an exceptionally high affinity for copper. We demonstrated that ATP7B-deficient rats recapitulate WD-associated phenotypes, including hepatic copper accumulation, liver damage, and mitochondrial impairment. Short-term treatment of these rats with MB efficiently reversed mitochondrial impairment and liver damage in the acute stages of liver copper accumulation compared with that seen in untreated ATP7B-deficient rats. This beneficial effect was associated with depletion of copper from hepatocyte mitochondria. Moreover, MB treatment prevented hepatocyte death, subsequent liver failure, and death in the rodent model. These results suggest that MB has potential as a therapeutic agent for the treatment of acute WD.

Introduction

Wilson disease (WD) is an autosomal recessively inherited disorder of copper metabolism caused by ATPase copper-transporting β (ATP7B) gene mutations (1–3). ATP7B is a copper-transporting ATPase that mediates copper excretion into bile (1, 4), and its functional loss causes a major impairment of the ability of hepatocytes to maintain copper homeostasis at the cellular and systemic levels, resulting in persistent copper accumulation in the liver. This can lead, most likely as a result of spillover of liver copper, to deleterious effects on the brain and, in many cases, to chronic liver disease and also to fulminant hepatic failure (1, 5–8).

To restore copper homeostasis, the clinically used copper chelators D-penicillamine (D-PA) and trientine (TETA) or the candidate drug tetrathiomolybdate (TTM) are administered daily (1, 5). This lifelong therapy is effective if commenced before the onset of advanced hepatic or neurologic disease (6–8). The same holds true for zinc salts, which are primarily used in mild cases of WD to decrease copper absorption via the gastrointestinal tract or as copper maintenance therapy in chelator-treated patients (1, 5, 9). However, in circumstances of acute liver failure — caused by delayed diagnosis, treatment failure, or rapidly developing fulminant hepatitis — death is almost certain unless liver transplantation is performed (1, 5, 7, 10, 11). Poor treatment compliance may be another reason for the death of these patients.

In Wilson disease (WD), functional loss of ATPase copper-transporting β (ATP7B) impairs biliary copper excretion, leading to excessive copper accumulation in the liver and fulminant hepatitis. Current US Food and Drug Administration– and European Medicines Agency–approved pharmacological treatments usually fail to restore copper homeostasis in patients with WD who have progressed to acute liver failure, leaving transplantation as the only viable treatment option. Here, we investigated the therapeutic utility of methanobactin (MB), a peptide produced by *Methylosinus trichosporum* OB3b, which has an exceptionally high affinity for copper. We demonstrated that ATP7B-deficient rats recapitulate WD-associated phenotypes, including hepatic copper accumulation, liver damage, and mitochondrial impairment. Short-term treatment of these rats with MB efficiently reversed mitochondrial impairment and liver damage in the acute stages of liver copper accumulation compared with that seen in untreated ATP7B-deficient rats. This beneficial effect was associated with depletion of copper from hepatocyte mitochondria. Moreover, MB treatment prevented hepatocyte death, subsequent liver failure, and death in the rodent model. These results suggest that MB has potential as a therapeutic agent for the treatment of acute WD.

### Related Commentary: p. 2412

**Authorship note:** J. Lichtmannegger and C. Leitzinger are co-first authors. F. Eckardt-Schupp is deceased.

**Conflict of Interest:** During the revision phase of this manuscript, H. Zischka, J. Lichtmannegger, and Alan A. DiSpirito filed a patent application on the clinical utility of copper chelation using methanobactin (“Means and methods for treating copper-related diseases.” Patent application no. EP15201070.8).

**Submitted:** October 22, 2015; **Accepted:** April 12, 2016.

**Reference information:** *J Clin Invest.* 2016;126(7):2721–2735. doi:10.1172/JCI85226.
Given these issues, there is a clear, unmet medical need for alternative and innovative WD treatments. In this regard, 3 recent reports merit special attention. First, gene therapy via adeno-associated viral vectors designed for liver-specific ATP7B expression provided long-term correction of copper metabolism in Atp7b–/– mice (12). This is an exciting new route for treating patients with WD; however, as transgene expression upon vector injection is of patients with WD (11). All of the currently Food and Drug Administration–approved and European Medicines Agency–approved copper chelators have adverse effects, including bone marrow toxicity, nephrotoxicity, hepatotoxicity, anemia, and triggering of autoimmune disease (1, 5, 7). Because of the toxicity of D-PA, discontinuation of treatment is required in almost one-third of patients with WD (5).
rather slow, this drawback may hamper its use in acute WD (13).
A second study reported that the plant-derived decapeptide OSIP108 protected against copper-induced cell death in yeast and human HepG2 cells. Intriguingly, this positive effect was not due to a reduction of cellular copper but rather to effects on sphingomyelin homeostasis (14). A third study reported hepatocyte targeting of 1,154 Da (metal free) and an exceptionally high copper affinity showing that treatment was initiated before the onset of clinically manifest liver damage (18). We recently described copper-driven fulminant hepatitis in LPP Atp7b–/– rats that carry a homozygous 13-kb deletion in the Atp7b gene (hereafter referred to as Atp7b–/– rats) (16–18). Because of this recessive genetic defect, heterozygous Atp7b+/- rats have a normal phenotype. In Atp7b–/– rats, however, accumulating liver copper results in WD-like pathology. Clinical manifestations include elevated levels of serum aspartate aminotransferase (AST) and bilirubin, rapidly followed by liver failure and death (18). The early hepatocellular phenotype in these animals is characterized by increased mitochondrial copper accumulation, provoking progressive alterations of their structure (18). A similar mitochondrial phenotype has been reported in WD patients and in other WD animal models (19–22). In Atp7b–/– rats, mitochondrial changes could be reversed by several weeks of treatment with copper chelators. However, this therapy was only successful when treatment was initiated before the onset of clinically manifest liver damage (18).

Here, we present the potential clinical utility of copper chelation using methanobactin (MB) to treat acute WD. MB is a posttranslationally modified peptide from the methanotrophic proteobacterium Methylosinus trichosporium OB3b, with a MW of 1,154 Da (metal free) and an exceptionally high copper affinity (23–25). MB to sequester extracellular copper (25, 27, 28). MB is permeable for their copper-dependent methane oxidase (27, 28) and excrete MB through the bacterial outer membrane, is reinternalized across the cytoplasmic membrane via a TonB transporter (29), and delivers copper either to methane oxidases or to storage proteins like copper storage protein 1 (CSP1) (30).

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Table 1. Mitochondrial copper content progressively increases with disease state in livers from Atp7b–/– rats

<table>
<thead>
<tr>
<th>Group (genotype)</th>
<th>Age (d)</th>
<th>Weight (g)</th>
<th>AST (U/l)</th>
<th>Bilirubin (mg/dl)</th>
<th>Cu (ng/mg protein)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control (+/−)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F, N = 15</td>
<td>83–146</td>
<td>170 ± 13</td>
<td>115 ± 28</td>
<td>&lt;0.5</td>
<td>14 ± 2</td>
</tr>
<tr>
<td>M, N = 9</td>
<td>81–95</td>
<td>281 ± 36</td>
<td>128 ± 37</td>
<td>&lt;0.5</td>
<td>15 ± 1</td>
</tr>
<tr>
<td>Affected (−/−)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F, N = 4</td>
<td>85–92</td>
<td>165 ± 16</td>
<td>138 ± 37</td>
<td>&lt;0.5</td>
<td>1,353 ± 137</td>
</tr>
<tr>
<td>M, N = 11</td>
<td>66–93</td>
<td>253 ± 25</td>
<td>151 ± 33</td>
<td>&lt;0.5</td>
<td>1,188 ± 196</td>
</tr>
<tr>
<td>Disease onset (−/−)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F, N = 4</td>
<td>85–93</td>
<td>150 ± 3</td>
<td>331 ± 45</td>
<td>&lt;0.5</td>
<td>1,385 ± 58</td>
</tr>
<tr>
<td>M, N = 6</td>
<td>80–107</td>
<td>275 ± 45</td>
<td>244 ± 55</td>
<td>&lt;0.5</td>
<td>1,337 ± 155</td>
</tr>
<tr>
<td>Diseased (−/−)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F, N = 1</td>
<td>84</td>
<td>162</td>
<td>590</td>
<td>0.5</td>
<td>1,389</td>
</tr>
<tr>
<td>M, N = 4</td>
<td>91–107</td>
<td>274 ± 20</td>
<td>412 ± 124</td>
<td>1.3 ± 1.1</td>
<td>1,389 ± 79</td>
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<tr>
<td>Moribund (−/−)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F, N = 3</td>
<td>97–106</td>
<td>140 ± 5</td>
<td>454 ± 96</td>
<td>14.2 ± 5.5</td>
<td>1,667 ± 478</td>
</tr>
<tr>
<td>M, N = 1</td>
<td>95</td>
<td>255</td>
<td>570</td>
<td>8.0</td>
<td>1,261</td>
</tr>
</tbody>
</table>
F, female; M, male.
Identical tissue damage features were observed in the livers of untreated WD patients and diseased Atp7b–/– livers (Figure 1A). Fibrosis was observed in all WD patients’ livers, and beginning fibrosis was observed in livers from diseased Atp7b–/– rats (Supplemental Figure 1A). These characteristics were absent from heterozygous Atp7b+/– control livers but steadily progressed in Atp7b–/– rats (Supplemental Figure 1B).

Another striking analogy between livers from patients with WD and Atp7b–/– rats was the structural damage of mitochondria. Transparent vacuoles of varying sizes containing amorphous but Identical tissue damage features were observed in the livers of untreated WD patients and diseased Atp7b–/– livers (Figure 1A). Fibrosis was observed in all WD patients’ livers, and beginning fibrosis was observed in livers from diseased Atp7b–/– rats (Supplemental Figure 1A). These characteristics were absent from heterozygous Atp7b+/– control livers but steadily progressed in Atp7b–/– rats (Supplemental Figure 1B).

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also electron-dense material, separated inner and outer membranes, marked differences in electron densities, and cristae dilations (Figure 1B and Supplemental Figure 1C) were observed as the typical WD mitochondrial phenotype (19). Importantly, we found highly comparable copper levels in liver homogenate and mitochondria obtained from diseased Atp7b−/− rats and untreated WD patients’ livers (Figure 1C). However, lower copper content was present in the homogenate from the explanted livers and iso-

Figure 3. MB depletes copper from liver mitochondria, hepatocytes, and whole liver. (A) MB, but not D-PA or TETA, extracted copper from Atp7b−/− mitochondria (2 mM each, 30-minute incubation, N = 3). (B) MB was less toxic to the mitochondrial respiratory complex IV than TTM (MB: N = 3, n = 9; TTM: N = 1, n = 3). (C) Copper-preloaded HepG2 (N = 3) and WD patient-derived HLCs (1 of 2 preparations) were de-coppered by MB. (-), untreated control; (+), 24-hour treatment with 300–500 μM MB. (D) Dose-dependent MB uptake into HepG2 cells at 2 and 24 hours (N = 3). (E) Toxicity of MB versus TTM in HLCs (neutral red assay; N = 3, n = 9). CCCP, positive control. (F) MB-treated (500 μM) HepG2 cells showed only intermediate phases of ΔΨ loss (6 hours, 250 μM CCCP). Nuclei, blue; mitochondria with ΔΨ, orange-red; mitochondria without ΔΨ, green. Arrows indicate cells with low ΔΨ (N = 2). Scale bars: 50 μm. (G) Cumulative copper excretion into bile following a 2-hour Atp7b−/− liver perfusion. Only MB (0.7 μM) forced high copper amounts into bile compared with TTM (0.8 μM), D-PA (2.2 μM), and TETA (1.8 μM). N = 3. Note the different scales for MB (right, blue axis) and buffer control, D-PA, TETA, and TTM (left, black axis). (H) All chelators except TTM transported copper to the perfusate (conditions as in G). N = 2. (I) Only MB markedly reduced liver copper levels during Atp7b−/− liver perfusion, but not D-PA, TETA, or TTM (conditions as in G). N = 3. One-way ANOVA with Tukey’s multiple comparisons test (A, B, D, E, and I); unpaired t test with Welch’s correction (C). *P < 0.05 and ****P < 0.0001 versus control (A–C and E); ****P < 0.0001 versus respective MB concentration (B and E). **P < 0.01 versus 0.05 mM MB; ***P < 0.05 versus 0.1 mM MB; and ****P < 0.05 versus 0.3 mM MB (D). Co, buffer-treated control.
Figure 4. Acute liver failure is efficiently avoided by a short-term in vivo treatment with MB. (A) Overt liver damage was reduced in MB-treated but not D-PA- or TETA-treated Atp7b−/− livers. Scale bars: 100 μm; H&E staining (symbols as in Figure 1 and Supplemental Figure 1). Daily doses were: 150 mg (130 μmol) MB/kg BW; 100 mg (540 μmol) D-PA/kg BW; and 480 mg (2,190 μmol) TETA/kg BW. Three-day MB treatment: 2 experiments; 5-day MB treatment: 5 experiments; D-PA and TETA treatments: 4 experiments. (B) AST values decreased in 2 of 3 and 4 of 6 Atp7b−/− rats treated for 3 or 5 days with MB, respectively, but not in untreated Atp7b−/− (N = 6; 3 affected and 3 diseased) or short-term D-PA− or TETA-treated rats (N = 4; 3 affected and 3 diseased). Treatment started in rats at 82–90 days of age. (C) Mild reduction of whole-liver and significant reduction of mitochondrial copper level in short-term MB-treated rats (3-day MB treatment, N = 3; 2 affected and 1 disease onset, aged 88–89 days; 5-day MB treatment, N = 5; 5 affected, aged 89–95 days) but not in untreated rats (N = 4; 2 affected, 2 diseased, aged 90–91 days) or D-PA− or TETA-treated rats (N = 4; 3 affected, 1 disease onset, aged 86–89 days). One-way ANOVA with Tukey’s multiple comparisons test. *P < 0.05 versus untreated controls. (D) Massively reduced numbers in mitochondria with severely impaired structure (type 4, arrows) were isolated from MB-treated rats but not from untreated (Figure 2A) or D-PA− or TETA-treated Atp7b−/− rats (quantification in Supplemental Figure 4A). Scale bars: 1 μm. (E) Treatment of Atp7b−/− control rats with MB (i.p. once daily on 2 consecutive days; N = 4) did not change whole-liver or mitochondrial copper levels. Untreated control, N = 3. (F) Upon i.p. injection, MB was only detectable in the serum for half an hour, indicating a very short systemic residence time (n = 2). U, untreated.
lated mitochondria of the D-PA-pretreated WD patients (Figure 1C). This coincided with more heterogeneous impairment of the mitochondrial structure (Figure 1B), which probably results from zonal heterogeneities originating from massive fibrosis within these livers (Figure 1A).

**Increasing copper load impairs mitochondrial membrane integrity and function.** Mitochondrial copper content progressively increased with disease state in livers from Atp7b–/– rats (Figure 1C and Table 1). This was paralleled by increasingly severe membrane deficits, as demonstrated directly at the level of freshly isolated mitochondria (Figure 2) and described as follows: (a) a drastic decrease in structurally normal rat liver mitochondria (types 1 and 2) from Atp7b–/– livers compared with control livers and a corresponding increase in the number of structurally altered organelles (types 3 and 4, Figure 2A); (b) a significant alteration of the mitochondrial membrane “fluidity” at the polar head groups of the membrane lipid-water interface (TMA-DPH), but not at the membrane inner lipid phase (DPH) (Supplemental Table 1), as revealed by membrane polarization measurements with the fluorophores 1,6-diphenyl-1,3,5-hexatriene (DPH) and trimethylammonium-DPH (TMA-DPH) (33, 34); (c) a large-amplitude swelling of control mitochondria upon induction of the mitochondrial permeability transition (MPT) (18, 35) by either calcium or copper that was significantly reduced in mitochondria from diseased and disease-onset Atp7b–/– rats (Figure 2B); (d) a significant impairment of the capacity of Cys-A to block calcium-induced MPT in Atp7b–/– versus control mitochondria (Figure 2C and Supplemental Table 2); (e) a jeopardization of the time stability of the inner mitochondrial transmembrane potential (ΔΨ) and loss of ΔΨ in Atp7b–/– mitochondria at earlier time points compared with control mitochondria (Figure 2D and Supplemental Table 3); and (f) a progressive and significant impairment of the ATP production capacity of Atp7b–/– mitochondria (Supplemental Figure 4F).

**The bacterial peptide MB rapidly depletes accumulated mitochondrial copper.** We compared the capability of MB with that of the copper chelators D-PA, TETA, and TTM to remove copper from freshly isolated Atp7b–/– mitochondria. While D-PA and TETA were without effect, TTM showed a tendency to remove copper, and MB significantly decreased the copper levels associated with Atp7b–/– mitochondria (Figure 3A). We obtained similar results with mitochondria from WT rats artificially preloaded with copper (Supplemental Figure 2, B and C). Furthermore, MB was significantly less toxic than TTM, as shown by the impaired vital copper-dependent mitochondrial respiratory complex IV (Figure 3B and Supplemental Figure 2D).

**MB efficiently de-coppers hepatocytes with low cell toxicity.** At the cellular level, overnight MB treatments significantly reduced cellular copper levels in HepG2 cells that were either copper preloaded at concentrations that showed no toxicity (Figure 3C and Supplemental Figure 2F) or in HepG2 cells that had a normal copper load (Supplemental Figure 2E). Moreover, in an attempt to test the efficacy of MB on samples from patients with WD, we reprogrammed urinary epithelial cells from these patients into induced pluripotent stem cells (iPSCs) that differentiated into hepatocyte-like cells (HLCs) (Supplemental Figure 2, G–J) and found comparable copper depletion levels in these HLCs following treatment with MB (Figure 3C).

Using a specific MB mAb, we found that MB was taken up dose dependently into HepG2 cells (Figure 3D) and that cytotoxicity occurred only at millimolar MB concentrations (Figure 3E). At the mitochondrial level, MB concentrations of 500 μM reduced the mitochondrial membrane potential only partially (Figure 3F). Thus, MB efficiently de-coppers hepatocytes, without major toxic side effects.

**MB directs liver copper into bile.** We further investigated the copper-removing efficiency of MB at the whole-organ level (Figure 3, G–I, Supplemental Figure 3). During a 2-hour perfusion of Atp7b–/– livers, MB released 10-fold greater amounts of copper into bile than did TTM (Figure 3G and Supplemental Figure 3B), the major physiological copper excretion route (36). D-PA and TETA did not provoke any detectable release of copper into bile (Figure 3G). All chelators, except TTM, caused an increased presence of copper in the perfusate (Figure 3H), which may be linked to the ability of TTM to precipitate copper intracellularly (37). The release of copper into the perfusate was partly due to hepatocellular death, as the cellular damage marker lactate dehydrogenase (LDH) paralleled the copper release curves (Supplemental Figure 3C). Of note, only MB treatment had already markedly reduced the Atp7b–/– liver copper levels after 2 hours of perfusion (Figure 3I).

**Short-term MB application reverses liver damage in vivo.** We assessed the efficiency of short-term MB treatment in Atp7b–/– rats at the age of liver disease onset (85–90 days). Animals received either MB (i.p.) for 3 or 5 days or the clinically used copper chelators D-PA or TETA (administered orally for 4 days).

MB administration strongly reduced the histopathological damage markers in Atp7b–/– livers (Figure 4A), in contrast to that observed with D-PA or TETA treatments. The latter 2 chelators were unable to circumvent the increase in serum AST levels (indicative of progressive liver damage, Figure 4B). In contrast, in 6 of 9 MB-treated Atp7b–/– rats, AST levels decreased (Figure 4B), and the animals regained BW (Table 2). Importantly, after 5 days of MB treatment, 2 Atp7b–/– animals with onset disease and 1 diseased Atp7b–/– rat were rescued from liver dysfunction (AST <200 U/l, Figure 4B and Table 2).

In comparison with untreated, age-matched Atp7b–/– rats and in contrast to D-PA or TETA treatments, MB induced a marked reduction in total liver copper levels, which was especially significant in the mitochondrial compartment (Figure 4C). This mitochondrial de-coppering was paralleled by ultrastructural and biochemical restoration of Atp7b–/– mitochondria (Figure 4D and Supplemental Figure 4F). Severely impaired mitochondria (type 4, Figure 2A) were almost absent in isolates from MB-treated Atp7b–/– rats, but not in isolates from D-PA– or TETA-treated animals (Figure 4D and Supplemental Figure 4A). Importantly, the impaired ATP production capacity of Atp7b–/– mitochondria was fully restored by MB, a result that was significantly different from that observed with D-PA or TETA treatments, which were without therapeutic effect (Supplemental Figure 4F).

**Toward a new WD treatment regimen using MB.** From a clinical perspective, we initiated experiments to scrutinize the potential therapeutic use of MB.

First, concerning drug safety, 4 heterozygous Atp7b–/– rats were treated i.p. with MB on 2 consecutive days. We observed no signs of toxicity, as BW, serum AST levels, and bilirubin values...
a clinically relevant copper maintenance therapy in WD (9). All MB-treated animals showed restoration of normal serum AST levels that lasted for at least 2 weeks, after which AST levels rose again (Figure 5A). At the time of analysis, 1 animal was still healthy, and 2 animals manifested different stages of liver disease (Figure 5, A and C). The degree of liver damage correlated with mitochondrial, but not whole-liver, copper levels (Figure 5B), as well as with structural (Figure 5D) and functional defects in mitochondria (Supplemental Figure 4B).

Fourth, we conducted a first test involving replacement of daily chelation therapy by a regimen consisting of intense treatment cycles interrupted by observation periods (Table 3). Five Atp7b–/– rats as well as 5 age- and sex-matched Atp7b +/– controls (untreated) were included, and pairs of rats were sacrificed on experimental days 1, 8, 29, 36, and 85, respectively. On experimental day 1, all animals were healthy, with the sacrificed Atp7b–/– rat showing a pronounced liver and mitochondrial copper load and slightly impaired mitochondrial function (87% ATP production capacity) compared with its Atp7b+/– control (pair 1). The 4 remaining Atp7b–/– rats were subjected to the first treatment cycle consisting of 3 daily i.p. MB injections for 5 days. All animals remained healthy, and this resulted in a 40% reduction in the copper load on experimental day 8 (pair 2), which increased to starting levels after an additional 3 weeks of observation (pair 3, day 29). Fol-

### Table 2. Short-term treatments with 5× MB (once daily, i.p. or i.v.) or with 16× MB (twice daily, i.p.) reduce the mitochondrial copper load

<table>
<thead>
<tr>
<th>Treatment (genotype)</th>
<th>Age (d)</th>
<th>Weight (g)</th>
<th>AST (U/l)</th>
<th>Bilirubin (mg/dl)</th>
<th>Cu (ng/mg protein)</th>
<th>Mitochondrial Cu depletion (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Start</td>
<td>End</td>
<td>Start</td>
<td>End</td>
<td>Homogenate</td>
<td>Mitochondria</td>
</tr>
<tr>
<td><strong>MB 2× i.p. (+/-)</strong></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>F, rat 1</td>
<td>89</td>
<td>188</td>
<td>181</td>
<td>108</td>
<td>&lt;0.5</td>
<td>13</td>
</tr>
<tr>
<td>F, rat 2</td>
<td>89</td>
<td>170</td>
<td>171</td>
<td>123</td>
<td>&lt;0.5</td>
<td>16</td>
</tr>
<tr>
<td>F, rat 3</td>
<td>161</td>
<td>209</td>
<td>212</td>
<td>139</td>
<td>&lt;0.5</td>
<td>11</td>
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<tr>
<td>F, rat 4</td>
<td>161</td>
<td>213</td>
<td>216</td>
<td>127</td>
<td>&lt;0.5</td>
<td>15</td>
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<tr>
<td><strong>MB 3× i.p. (-/-)</strong></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>F, rat 1 (A)</td>
<td>89</td>
<td>144</td>
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<td>131</td>
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<td>1,307</td>
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<tr>
<td>M, rat 2 (A)</td>
<td>88</td>
<td>261</td>
<td>261</td>
<td>192</td>
<td>&lt;0.5</td>
<td>881</td>
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<tr>
<td>F, rat 3 (A)</td>
<td>88</td>
<td>259</td>
<td>260</td>
<td>106</td>
<td>&lt;0.5</td>
<td>840</td>
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<tr>
<td><strong>MB 5× i.p. (-/-)</strong></td>
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<td></td>
<td></td>
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<tr>
<td>F, rat 1 (A)</td>
<td>91</td>
<td>157</td>
<td>154</td>
<td>116</td>
<td>&lt;0.5</td>
<td>–</td>
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<tr>
<td>M, rat 2 (Do)</td>
<td>90</td>
<td>197</td>
<td>205</td>
<td>480</td>
<td>&lt;0.5</td>
<td>806</td>
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<tr>
<td>M, rat 3 (Do)</td>
<td>90</td>
<td>231</td>
<td>242</td>
<td>270</td>
<td>&lt;0.5</td>
<td>851</td>
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<tr>
<td>F, rat 4 (Do)</td>
<td>95</td>
<td>160</td>
<td>164</td>
<td>369</td>
<td>&lt;0.5</td>
<td>915</td>
</tr>
<tr>
<td>F, rat 5 (A)</td>
<td>89</td>
<td>171</td>
<td>178</td>
<td>83</td>
<td>&lt;0.5</td>
<td>728</td>
</tr>
<tr>
<td>M, rat 6 (A)</td>
<td>89</td>
<td>258</td>
<td>269</td>
<td>97</td>
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<tr>
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<td>135</td>
<td>143</td>
<td>295</td>
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<td>163</td>
<td>273</td>
<td>&lt;0.5</td>
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</table>

*Mitochondrial copper depletion relating to the respective untreated group (Table 1). A, affected; Do, disease onset; D, diseased; Mo, moribund.
die within a few days. In contrast, following the “acute rescue regimen,” animal 3 regained 29% of its BW, had a drastic reduction of AST and bilirubin to normal levels, and hepatic copper depletion associated with massive structural and functional mitochondrial recovery (Table 2 and Supplemental Figure 4, D–F).

Discussion
Mitochondrial copper overload occurs to a similar extent in WD patients and Atp7b−/− rats (Figure 1). This overload causes mitochondrial impairments (Figure 2) that lead to liver damage (Figures 1 and 5 and Supplemental Figures 1 and 4). Failure to respond to established treatments in human WD has been associated with a failure to deplete mitochondrial copper stores (38). Thus, reversal of mitochondrial copper overload seems to be a prerequisite for successful treatment. Clinically used copper chelators slowly deplete copper stores, are ineffective in some patients with chronic WD, and fail to rescue many patients with acute, fulminant WD–induced hepatitis and liver failure (1, 5, 10). Here, we show that MB rapidly and effectively depleted excess hepatic copper stores at mitochondrial, cellular, and organ levels (Figure 3 and Supplemental Figures 2 and 3). In vivo, MB reversed WD–associated hepatitis and was effective even when applied in acute, fulminant liver disease (Figures 4 and 5; Supplemental Figure 4; and Table 2).
The increasing copper burden directly affected the structural, physical, and biochemical integrity of mitochondrial membranes (Figure 2). Atp7b+/− liver mitochondria showed considerable matrix condensations, intermembrane space deposits, and cristae dilatations (Figure 2A). Using fluorescence polarization, we observed a steady increase in TMA-DPH polarization values in Atp7b+/− mitochondrial membranes and an increasing copper load (Supplemental Table 1). Thus, excessive mitochondrial copper deposition physically alters the mitochondrial membrane lipid-water interface (33, 34, 39), plausibly by direct interaction with mitochondrial membrane proteins (18). This results in a significantly lower extent of expansion upon induction of the MPT (ref. 40 and Figure 2B). Furthermore, the capacity of Cys-A to block calcium-induced MPT was significantly impaired in Atp7b+/− liver mitochondria (Figure 2C and Supplemental Table 2). Cys-A normally inhibits MPT by blocking inner mitochondrial membrane binding of its target protein Cyp-D (40, 41). Biochemically, Atp7b+/− mitochondria lost their ability to sustain their inner transmembrane potential (ΔΨ) with a progressively greater copper load (Figure 2D and Supplemental Table 3), as well as their capacity to produce ATP (Supplemental Figure 4F). Since impaired mitochondrial membranes can seal the point of no return toward death (42), we conclude that efficient therapeutic measures against WD must rely on the reduction of mitochondrial copper concentrations.

Hepatic copper transport and homeostasis require various membrane transporters, a subset of intracellular copper chaperones (43), and low-molecular-weight binding partners like glutathione (44), but also as-yet uncharacterized intracellular ligands (4, 45). Copper enters the cell via high-affinity transporters like copper transporter 1, and it is believed that differential copper affinities of the binding partners direct the metal to its final cellular destination (44). Copper can enter the Cu-metallothionein (MT) storage pool, be routed to copper-dependent enzymes (e.g., Cu/Zn SOD or the mitochondrial cytochrome c oxidase [Cco]), or be transported out of the cell via ATP7B (4, 43). As the latter transporter is dysfunctional in WD, a large part of the intracellular copper accumulates in the cytosolic MT storage pool (46, 47). However, in the Long-Evans Cinnamon (LEC) rat model of WD, it was shown that the cytosol from livers just before hepatitis still contained Cu-unsaturated MTs and thus may still act as antioxidants (47). On the contrary, excess mitochondrial copper loads and severe mitochondrial destruction were observed at the onset of hepatitis in the LEC and Atp7b+/− rats (18), arguing for the pivotal need to reverse this mitochondrial copper overload. Given that mitochondrial Cco, the intermembrane Cu/Zn SOD, and the mitochondrial copper chaperones present with exceptionally low Cu+ dissociation constants, there is a considerable driving force of copper into mitochondria (20, 44). Consequently, efficient mitochondrial copper removal may be achieved by compounds with high copper affinities. MB has an exceptionally high copper affinity (28) of more than 10−25 M, which is orders of magnitude higher than the affinities of known copper-binding proteins and ligands in eukaryotic cells (44). Thus, MB could possibly reverse mitochondrial copper overload.

In agreement with this, MB was able to efficiently deplete accumulated copper at the mitochondrial, cellular, and whole-organ levels to a considerable extent via excretion into bile, and no significant cell toxicity or mitochondrial dysfunction was induced (Figure 3 and Table 2). Of note, we found that a higher degree of liver damage (LDH release) coincided with elevated copper mobilization upon liver perfusion with chelating agents (Supplemental Figure 3C). This finding may be of clinical relevance, as worsening of the clinical presentation is reported in up to 20% of WD patients with neurological symptoms, despite sufficiently dosed chelation treatment with D-PA or TETA (5, 48, 49). Such deterioration may be explained by chelator-induced liver copper mobilization possibly leading to secondary copper toxicity to the brain (48, 50, 51). Thus, in severe WD states, liver copper mobilization
into the bloodstream may be adverse — if the low-affinity chelator used succeeds in extracting copper from hepatic stores — yet releases the bound metal before its excretion. Therefore, treating WD with high-affinity chelators such as MB, which has a copper affinity several orders of magnitude greater than that of D-PA (52), may constitute a pivotal advantage. In agreement with this notion, in preliminary studies, we observed that administering MB for 2 to 8 days (i.p.) did not elevate total brain copper levels. In addition, several studies (e.g., refs. 53, 54) reported improvement of neurological symptoms upon liver transplantation, strengthening the view that some of the excess brain copper may be excreted upon restoration of a physiological copper excretion into bile. MB causes a highly efficient excretion of copper into bile (Figure 3G). Taken together, these results support the prediction of an improvement of neurological symptoms upon MB therapy; however, this needs to be substantiated in future studies.

The superior in vitro and ex vivo de-coppering efficiency of MB led us to investigate a new treatment strategy in Atp7b–/– rats. Instead of the typical WD treatment regimen involving long-term oral administration of copper chelators, we used a short but intense de-coppering treatment with repetitive injections of MB (Figures 4 and 5, and Table 2). In contrast to D-PA and TETA, short-term i.p. MB administration significantly de-coppered Atp7b–/– liver mitochondria, which was paralleled by a regression of mitochondrial structural deficits and liver damage (Figure 4). In addition, following a 5-day MB regimen and subsequent treatment discontinuation, all animals remained healthy for at least 2 weeks (Figure 5). This delayed disease reappearance cannot be attributed to the zinc-enriched diet given from the start of MB treatment, since hepatic copper reaccumulation occurred despite zinc supplementation. In fact, in a further experiment with repeated MB treatment cycles and drug-free intervals without zinc supplementation (Table 3), the disease onset was delayed to beyond 166 days of age, an approximate doubling of the disease-free period in the Atp7b–/– rat. Importantly, the disease status clearly correlated with mitochondrial, but not whole-liver, copper content (Figure 5B). Therefore, the parameters for mitochondrial copper content, structure, and biochemical functionality not only serve as early response markers for disease progression in WD patients and Atp7b–/– rats (Figures 1 and 2 and Supplemental Figure 4F), but also as diagnostic biomarkers of treatment efficiency (Figure 4, C and D) and predictive markers of recurrence of liver damage (Figure 5, B and D, and Table 3). We observed that both i.p. and i.v. MB administration efficiently depleted mitochondrial copper levels (Table 2). At present, the acid sensitivity of MB (55) precludes its oral administration (in contrast to current clinical pharmacotherapy), a limitation that may, however, be overcome by using biocompatible polymers or other galenic formulations (56). Nevertheless, parenteral MB administration has fast and reliable effects, and injecting MB twice daily proved to be particularly efficient, reversing the disease phenotype even in animals that had signs of advanced liver dysfunction (Table 2 and Supplemental Figure 4D–F). Further, concerning the temperature stability of MB, we observed a time-dependent destabilization of MB at 37°C (Supplemental Figure 4C), which was effectively avoided, however, upon prestabilization by zinc (Supplemental Figure 4C) and did not interfere with the de-coppering properties of MB (Figure 3C and Supplemental Figure 2E). Hence, zinc-stabilized MB may be advantageous for applications requiring prolonged persistence at elevated temperatures.

With regard to drug safety, we observed no signs of MB toxicity in Atp7b–/– rats treated up to 3 times per day (Table 3) or in heterozygous Atp7b+/– control rats treated for 2 consecutive days (Figure 4E). However, one potential issue could be an immune response against the MB peptide. While it cannot be completely excluded, this appears unlikely, because small compounds like MB, with a MW of approximately 1,000, are considered nonimmunogenic (57). Moreover, no toxicity was observed in animals that underwent repetitive cycles of intense MB treatments (Table 3), and the first ELISA analyses testing sera of these treated animals for “anti-MB” Abs were negative. Nevertheless, this issue will need further attention.

In conclusion, we report what we believe to be an innovative therapeutic strategy for treating WD that consists of a short but intense de-coppering treatment with MB that proved effective, even at the acute liver failure stage in a WD rat model. In total, 24 Atp7b–/– rats at various stages of disease were highly effectively treated once, twice, or thrice daily, either by i.p. or i.v. injection of MB (Tables 2 and 3, and Figure 5). All treated animals were rescued, and liver damage reappeared only after several weeks of drug-free phases. The treatment of patients with bacterial siderophores is not without precedent, as exemplified by deferoxamine from Streptomyces pilosus, which is used for treating iron poisoning and hemochromatosis (58). We therefore propose MB as an additional candidate in this pharmaceutical class and as the lead drug of choice to develop a more efficient and safe treatment of WD.

Methods

Patient-derived samples
Livers from 4 WD patients with liver failure who had undergone liver transplantation at the University Hospital Heidelberg were included in this study. Two patients (patients 1 and 2) had received no prior copper chelation therapy, while 2 patients (patients 3 and 4) presented with liver failure after D-PA treatment. Upon explantation, the WD patients’ livers were shock-frozen in LN2 and stored at –80°C. Thawed samples were immediately fixed using formalin (Bio Optica Milano) for histological analysis or glutaraldehyde (Electron Microscopy Sciences) for electron microscopic analysis. WD patients’ urinary epithelial cells were collected from freshly donated mid-stream urine.

Animals
The LPP rat strain was provided by Jimo Borjigin (University of Michigan, Ann Arbor, Michigan, USA) (16), and Lou/c rats were originally from Harlan Winkelmann. Rats were maintained on an ad libitum Altromin 1314 diet (Altromin Spezialfutter) and tap water. The basic copper content of this standard diet was 13 mg/kg. Heterozygous Atp7b+/– rats served as controls in this study.

Animal treatments

In vivo treatments. LPP rats were treated with MB by daily i.p. injections for 3 or 5 consecutive days or by twice-daily i.p. injections for 8 consecutive days at a dose of 150 mg/kg BW, or for 4 days via D-PA-contain-
ing drinking water at a dose of 100 mg/kg BW/day or TETA-containing drinking water at 480 mg/kg BW/day (assuming 40 ml water intake per day for a rat weighing 250 g), respectively (59, 60). Based on a mean copper content of 250 μg/g wet weight (ww) in Atp7b−/− rats at the start of treatment (18), Atp7b−/− rats of 8 g ww contain approximately 31.5 μmol copper. Single MB doses were chosen in amounts equimolar to this copper amount. As for D-PA, the dose and route of administration used in this study were previously reported to successfully prevent the onset of hepatitis in LEC rats in long-term regimens (59). Subchronic toxicity studies in rats have revealed no toxicity of TETA at a dose of 3,000 ppm via the drinking water (60). With respect to the mean liver copper content, studies in rats have revealed no toxicity of TETA at a dose of 3,000 ppm via the drinking water (60).

Liver perfusion

Atp7b−/− livers (from animals aged 79–83 days) were perfused via the portal vein in a single-pass manner with Krebs-Ringer bicarbonate solution containing 5 mM glucose, gassed with 95% O2 and 5% CO2, and kept at 37°C (61, 62). The right lateral liver lobe was ligated, and its copper content served as a preperfusion control. After bile duct cannulation, a 20-minute bile sample was collected before the copper chelators were continuously added to the perfusion medium. Bile and outflow perfusates (from which LDH was measured) were collected at 10-minute intervals (61, 62). D-PA/HCl (20 mg/108 μmol); TETA/2HCl (20 mg/91 μmol); TTM/2 NH4 (10 mg/38 μmol); and MB (40 mg/35 μmol) were each dissolved in 50 ml 0.9% NaCl and continuously added to the perfusion medium. Bile and outflow perfusates (from which LDH was measured) were collected at 10-minute intervals (61, 62).

Histological examination, plasma/serum AST, and bilirubin

Formalin-fixed, paraffin-embedded liver samples were cut into 4-μm-thick sections and stained with either H&E (Carl Roth GmbH + Co. KG) or Masson’s trichrome (Merck KG) for analysis of fibrotic tissue. AST activity and bilirubin concentration in animal plasma or serum were measured with a Reflotron system (Roche Diagnostics).

Mitochondrial analyses

Mitochondria were derived from either frozen, explanted WD patients’ livers or freshly prepared rat liver homogenates, as described previously (35, 63), and purified by differential and density gradient centrifugation using either Percoll (GE Healthcare Europe) or Nycodenz (Axis-Shield). Fresh rat liver mitochondria were used for respiratory measurements, chelator treatments, analyses of swelling (MPT), transmembrane potential (∆ψm), polarization experiments, and ATP synthesis and were fixed with glutaraldehyde for subsequent electron microscopic analyses. Stored frozen mitochondria were used for respiratory complex IV activity and metal analyses.

Mitochondrial functional integrity was assessed by standard respiratory measurements in a Clark-type oxygen electrode (Oxygraph; Hansatech Instruments) (18). ATP synthesis was analyzed by the ATP Bioluminescence Assay Kit (Roche) (18). Mitochondrial swelling was measured by light scattering at 540 nm, and ∆ψm was followed by Rh123 fluorescence quenching in a 96-well plate reader (BioTek, Synergy 2; Bad Friedrichshall) (64). For calculation, in cases with onset but not-yet finished MMP depletion, end values were set to 180 minutes. Polarization was measured in DPH and TMA-DPH–dyed mitochondria (33, 34). In brief, mitochondria (3 mg/ml) were incubated for 30 minutes at 37°C with either DPH or TMA-DPH (50 μM and 20 μM, respectively). Parallel and perpendicular fluorescence was assessed in duplicate at ex 366 nm and em 425 nm. Polarization was calculated (65) in mP using the following formula: P = (I1 – G × I2)/(I1 + G × I2); G = 0.89.

In vitro treatment of isolated mitochondria with chelators. Freshly isolated Atp7b−/− mitochondria with elevated copper levels were subjected to 30-minute chelator treatments with either 2 mM D-PA, TETA, TTM, or MB and subsequently repurified by a Nycodenz-gradient to separate liberated copper from mitochondria. For validation, mitochondria from Atp7b−/− control rats were incubated with 1 mM DTT (Roche) for 5 minutes at room temperature, and thereafter, Cu2+ (Copper II Chloride Dihydrate; Sigma-Aldrich) was added at final concentrations of 200–600 μM for 20 minutes. Copper-loaded mitochondria were then repurified by Nycodenz-gradient centrifugation and subsequently treated with chelators as described above.

Cell culture

All cell lines were from ATCC unless otherwise indicated.

HepG2 cells were kept in MEM (Life Technologies) with 2% FCS (Biochrom). We found that Zn-MB was time stable at 37°C, in contrast to metal-free MB (Supplemental Figure 4C). Therefore, Zn-MB, generated by preparing a 20-mM MB solution and adding an equimolar concentration of Zn2+ (Zinc acetate; Sigma-Aldrich) solution under pH control, was used in cell culture experiments.

A neutral red cell toxicity assay (Sigma-Aldrich) was performed as described elsewhere (66). HepG2 cells (2 × 104) were incubated with 1 mM MB for 24 hours with medium alone (negative control), 0.05 to 1 mM zinc-MB or TTM, 0.002–1 mM copper-histidine, or 0.25 mM carbonyl cyanide m-chlorophenylhydrazone (CCCP) as a mitochondrial toxic positive control and were subsequently analyzed.

For immunofluorescence staining, 2 × 104 HepG2 cells were incubated with medium alone, 500 μM MB, or 250 μM CCCP in black 96-well plates with a clear glass bottom (PerkinElmer). Staining was done using 1.6 μM Hoechst 33342 (excitation wavelength [ex] 360–400 nm; emission wavelength [em] 410–480 nm); 300 nM MitoTracker Deep Red (ex 620–640 nm, em 650–760 nm); and 1 μM nonyl acridine orange (NAO) (ex 460–490 nm, em 500–550 nm) for 40 minutes at 37°C. After a washing step, fluorescence was analyzed using an Operetta System (PerkinElmer).

Cellular de-coppering by MB was assessed in HepG2 cells pretreated for 24 hours with either medium or 15 μM copper-histidine and after a subsequent 24-hour treatment with 500 μM MB. After washing steps, copper levels in 2.5 × 104 HepG2 cells were determined by inductively coupled plasma optical emission spectrometry (ICP-OES). Cellular MB uptake was determined from HepG2 cell lysates incubated for 2 or 24 hours with MB at different concentrations by a competitive ELISA using an anti-MB mAb.

MB Ab generation and competitive ELISA

Lou/c rats were immunized with a mixture of OVA-coupled MB (50 μg) (Squarix), 5 nmol CPG oligonucleotide (TIB MOLBIOL; Syntheselabor),
Generation of HLCs from WD patients

WD patients’ urinary epithelial cells were pelleted at 400 ×g for 10 minutes from freshly donated mid-stream urine (68). Cells were cultured in culture medium (DMEM/F12; Lonza Group) supplemented with 10% FBS (PAA Laboratories); 0.1 mM nonessential aa (NEAA) (Sigma-Aldrich); 0.1 mM β-mercaptoethanol (Gibco); and 1 mM Glutaredoxin (New England BioLabs) to ensure optimal growth. The cultures were then trypsinized and passaged 1:3 until they reached confluence. The medium was changed every 3 days, and the cultures were monitored for signs of contamination. The HLCs were then collected and used for further experiments.

Metal content determination

Copper levels in liver homogenates, cell lysates, and mitochondrial preparations were analyzed by ICP-OES (Thermo Fisher Scientific) after wet-ashing of samples with 65% nitric acid (Merck KGaA). Copper levels were measured in triplicate and expressed as micrograms per gram of tissue.

Electron microscopic analysis

Electron microscopic analysis of liver tissues and mitochondria was done as previously described (35).

Mitochondria structural analyses

For structural analyses, isolated mitochondria were categorized as follows: type 1 (normal structured mitochondria of the “condensed” type; ref. 71); type 2 (mitochondria with minor alterations such as slightly increased cristae); type 3 (mitochondria with massively increased cristae); and type 4 (mitochondria with massive matrix condensations, matrix vacuolization, detachments of the inner boundary membrane, and severe cristae deformations).

Miscellaneous

MB was isolated from the spent media of Methylosinus trichosporium OB3b as previously described (72). Endotoxin in MB was detected by a kinetic chromogenic method (Charles River Laboratories) and was on average 4.5 IU/mg. Copper-histidine solutions (1:3) were prepared as described previously (73). Protein quantification was done by the Bradford assay. Cytochrome C oxidase activity in isolated mitochondria was determined as described elsewhere (74).

Chemicals

D-P A was a gift of Heyl Pharma; TETA was from Sigma-Aldrich; and TTM (98% pure) was a gift of KT. Suzuki (Chiba University, Chiba Japan). CCCP was obtained from Sigma-Aldrich; DPH, TMA-DPH, MitoTracker Deep red, and NAO were obtained from Molecular Probes; and Hoechst 33342 was obtained from Life Technologies.

Statistics

Throughout this study “N” equals the number of analyzed animals and “n” the number of measurements. Data are presented as the mean ± SD. A Grubbs outlier test (ESD method, extreme studentized deviate) was performed if more than 4 technical replicates were available. A significance level of α = 0.05 was used. In detail, data from Figure 1C, Figure 2, A and B, Supplemental Figure 4F, and Supplemental Tables 1–3 were corrected for outliers. Statistical significance was analyzed using 1-way ANOVA with Tukey’s multiple comparisons test when comparing 3 or more sample sets (GraphPad Prism 7; GraphPad Software). For comparison between 2 groups the unpaired 2-tailed Student’s t test with Welch’s correction was used (GraphPad Prism 7). A P value of less than 0.05 was considered statistically significant.

Study approval

All patients provided informed consent, and the study was approved by the ethics committee of the Medical University of Heidelberg, Germany (for the study of liver samples from patients with WD) and by the
Ethis-Kommission der Ärztekammer Westfalen-Lippe und der Westfälischen Wilhelms-Universität Münster, Germany (for the study of WD patients’ urinary epithelial cells).

Animal experiments were approved by the government authorities of the Regierung von Oberbayern, Munich, Germany, and all animals were treated according to the guidelines for the care and use of laboratory animals of the Helmholtz Center Munich.

Author contributions

JL and CL designed and performed experiments, analyzed data, and wrote the manuscript. RW, S. Schmitt, S. Schulz, CE, TR, and YK performed experiments. DJ, FN, BKS, and PS performed histochemical and transmission electron microscopy analyses. AADS, NB, and BSB produced MB samples. AF and EK established primary anti-MB Abs and performed ELISA analyses. GD, FPR, SH, FES, NAD, and JA analyzed data. VS, CN, and HHJS constructed and performed HLC experiments. UM and DNG examined patients and collected clinical samples. GK designed experiments and wrote the manuscript. KHW examined patients, collected clinical samples, and designed experiments. HZ designed experiments, analyzed data, wrote the manuscript, and directed this study.

Acknowledgments

We are deeply indebted to the late Friedericke Eckardt-Schupp, who continuously supported this work. The authors would like to thank E. Samson for technical assistance and M. Atkinson and E.E. Rojo for critical reading of the manuscript. This study was supported in part by the Deutsche Forschungsgemeinschaft (DFG) grant RU742/6-1 (to H. Zischka) and STR 1160/1-2 (to B.K. Straub).

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