The Menkes and Wilson disease genes counteract in copper toxicosis in Labrador retrievers: a new canine model for copper-metabolism disorders

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ABSTRACT

The deleterious effects of a disrupted copper metabolism are illustrated by hereditary diseases caused by mutations in the genes coding for the copper transporters ATP7A and ATP7B. Menkes disease, involving ATP7A, is a fatal neurodegenerative disorder of copper deficiency. Mutations in ATP7B lead to Wilson disease, which is characterized by a predominantly hepatic copper accumulation. The low incidence and the phenotypic variability of human copper toxicosis hamper identification of causal genes or modifier genes involved in the disease pathogenesis. The Labrador retriever was recently characterized as a new canine model for copper toxicosis. Purebred dogs have reduced genetic variability, which facilitates identification of genes involved in complex heritable traits that might influence phenotype in both humans and dogs. We performed a genome-wide association study in 235 Labrador retrievers and identified two chromosome regions containing ATP7A and ATP7B that were associated with variation in hepatic copper levels. DNA sequence analysis identified missense mutations in each gene. The amino acid substitution ATP7B:p.Asp1453Gln was associated with copper accumulation, whereas the amino acid substitution ATP7A:p.Thr327Ile partly protected against copper accumulation. Confocal microscopy indicated that aberrant copper metabolism upon expression of the ATP7B variant occurred because of mis-localization of the protein in the endoplasmic reticulum. Dermal fibroblasts derived from ATP7A:p.Thr327Ile dogs showed copper accumulation and delayed excretion. We identified the Labrador retriever as the first natural, non-rodent model for ATP7B-associated copper toxicosis. Attenuation of copper accumulation by the ATP7A mutation sheds an interesting light on the interplay of copper transporters in body copper homeostasis and warrants a thorough investigation of ATP7A as a modifier gene in copper-metabolism disorders. The identification of two new functional variants in ATP7A and ATP7B contributes to the biological understanding of protein function, with relevance for future development of therapy.

KEY WORDS: ATP7A, ATP7B, Dog, Liver

INTRODUCTION

Copper is an essential trace element for a wide range of biochemical processes in the body. The P-type copper-transporting ATPases ATP7A (Vulpe et al., 1993) and ATP7B (Bull et al., 1993) are crucial for copper homeostasis. At the cellular level, these proteins have a biosynthetic role in the trans-Golgi network, where they facilitate incorporation of copper into proteins. Furthermore, they prevent toxic accumulation of cellular copper by redistribution to a vesicular compartment, resulting in excretion of copper through either the apical membrane (ATP7B) or the basolateral membrane (ATP7A). This redistribution involves metal-binding domains (MBDs) in cytoplasmic domains of the proteins and is regulated in part by their phosphorylation state (Voskoboinik et al., 2002).

Body copper homeostasis is maintained by balancing the rates of dietary copper absorption and biliary copper excretion. ATP7A is highly expressed in enterocytes, where it is involved in copper uptake across the basolateral membrane into the portal circulation for delivery to the liver, where copper is stored (Linder and Hazegh-Azam, 1996). Besides its function in intestinal epithelial cells, ATP7A was recently identified to play an important role in hepatic copper mobilization in response to copper demands of peripheral tissues (Kim et al., 2010). Excess hepatic copper is excreted into the bile and requires expression of functional ATP7B in hepatocytes (Gitlin, 2003).

The severe effects of copper-metabolism imbalance are illustrated by inherited disorders resulting from mutations in ATP7A and ATP7B.

Mutations in ATP7A result in a fatal, X-linked copper-deficiency disorder in infants known as Menkes disease. The disease is characterized by cerebral and cerebellar degeneration, failure to thrive, coarse hair and connective tissue abnormalities (Kaler, 2011). Wilson disease results from mutations in ATP7B (Gitlin, 2003) and is associated with copper accumulation in the liver and secondarily in the brain, resulting in hepatic cirrhosis and neuronal degeneration. The age of onset and the clinical manifestations vary greatly between individuals affected by Wilson disease. This lack of
genotype-phenotype correlation might be influenced by currently unidentified genetic modifiers. Other hereditary diseases leading to hepatic copper accumulation in infants include Indian childhood cirrhosis (Tanner, 1998) and endemic Tyrolean infantile cirrhosis (Müller et al., 1996). In these diseases, the causal genes are currently unknown, and dietary copper intake is believed to contribute significantly to disease progression.

In order to develop new treatment strategies for copper-metabolism disorders, several rodent models were investigated, including natural models, such as the mottled mouse (Grimes et al., 1997), the toxic milk mouse (Theophilos et al., 1996) and the Long–Evans cinnamon rat (Li et al., 1991), in addition to ATP7A (Wang et al., 2012) and ATP7B knockouts (Buiakova et al., 1999). Although rodent models are invaluable for studying diseases, the dog as a large-animal model represents a unique translational bridge between rodents and humans. The best-characterized canine model for copper toxicosis is the Bedlington terrier, in which severe hepatic copper accumulation is caused by a deletion in the COMMD1 gene (Van de Sluis et al., 2002). However, convincing evidence of involvement of the COMMD1 gene in human copper-metabolism disorders is lacking (Coronado et al., 2005; Lovicu et al., 2006; Müller et al., 2003).

The Labrador retriever dog breed was recently characterized as a new mammalian model for copper toxicosis, distinct from COMMD1-associated autosomal recessive copper toxicosis. In predisposed Labrador retrievers, hepatic copper accumulation induces hepatic cirrhosis, usually in middle-aged dogs. Female Labrador retrievers are at increased risk for the disease (Fieten et al., 2012b).

The genes underlying the disease in Labrador retrievers were not known. We hypothesized that identification of these genes would contribute to the understanding of copper-metabolism biology in dogs and other mammals. Given that the copper genome is highly conserved across species, this study might reveal genes that are candidate modifiers in human copper-metabolism disorders. Furthermore, elucidating the genetic background of copper toxicosis in Labrador retrievers contributes to the characterization of this new model for copper-metabolism disorders, necessary to exploit its possibilities fully.

Whereas gene-mapping studies in human complex diseases require thousands of affected individuals and millions of single nucleotide polymorphism (SNP) markers, smaller numbers suffice for a genome-wide association study (GWAS) of inbred pedigree dogs owing to the reduced genetic and phenotypic heterogeneity. Therefore, the dog is a useful model for identifying genes in naturally occurring complex hereditary diseases that are phenotypically similar in dogs and humans (Boyko, 2011).

The aims of the present study were to identify gene variants involved in copper toxicosis in the Labrador retriever and to provide biological evidence for their involvement in the disease.

RESULTS

Dogs

The initial GWAS cohort consisted of 235 Labrador retrievers. The independent replication cohort consisted of 59 Labrador retrievers. The median copper score in the 294 Labrador retrievers included in the study was 1.5 (range 0-5). A summary of characteristics of the dogs, including age, sex and hepatic copper scores, is provided in Table S1.

Electron microscopic evaluation of copper toxicosis in Labrador retrievers

To complement the phenotype description of copper toxicosis in Labrador retrievers, liver biopsies of two Labrador retrievers affected with copper toxicosis and a control were assessed by electron microscopy. Ultrastructural changes in the biopsies of the affected dogs corresponded to copper-laden lysosomes, comparable to what is recognized by electron microscopy of the liver of human individuals with Wilson’s disease (Fig. 1).

Genome-wide association study of copper toxicosis in Labrador retrievers

After quality control of genotype data used for the GWAS, 35.7% of SNPs were removed because of a low minor allele frequency and 0.7% of SNPs had a call rate of less than 98%, leaving 109,496 SNPs and 235 dogs for population stratification was determined by a multidimensional scaling plot (Fig. S1), and inflation of the test statistic was inspected by a QQ-plot (Fig. S2). The heritability of hepatic copper score was estimated to be 0.48. The highest association signal using hepatic copper level as a quantitative trait in a linear mixed model was identified on chromosome 22, with a $P$-value of $2.4 \times 10^{-8}$ and a $\beta$ of 0.65 (s.e. 0.14; Fig. 2A). Inspection of the region showed a linkage disequilibrium (LD) block at the first 10 Mb of the chromosome. The crucial interval was determined to be at 22:152034-10753911, which contained 83 genes, including the candidate gene ATP7B, located at position 22:162769-225266 (Canfam 3.1; Fig. 2B).

A stratified analysis for males and females was performed (Fig. 2C,D). In the male subset, the crucial region was located at X:54322875-65713151 and contained 77 genes. The highest association signal [$P=1.1 \times 10^{-4}$ and a $\beta$ of $-0.57$ (s.e. 0.15)] was caused by six SNPs that formed an LD block of 8.2 Mb, including the candidate gene ATP7A at position X:60203319-60356690 (Fig. 2C).

Given the fact that ATP7A and ATP7B code for copper transporters, they were strong candidate genes for involvement in copper toxicosis; therefore, we focused on these genes for in-depth DNA sequence analysis. A list of observed DNA variants and their effect on the hepatic copper score is presented in Table 1.

The five nominally significant variations ($P_{\leq 0.05}$) were genotyped in an independent cohort of 59 Labrador retrievers (Table 1; Table S1). The association with hepatic copper score was replicated for the non-synonymous nucleotide substitution of ATP7A at X-chromosomal position 60279238 (ENSCAFT00000049745 ATP7A: c.980C>T), the intronic variation in ATP7A (position X:60338569) and the non-synonymous nucleotide substitution of ATP7B at chromosome 22, position 225112 (ENSCAF0000006859 ATP7B:c.4358G>A). When analysed in the complete data set ($n=294$), the $P$-values were similar to or lower than initially derived in the GWAS, $1.6 \times 10^{-7}$ and $4.9 \times 10^{-7}$ for ATP7A:c.980C>T and ATP7B:c.4358G>A, respectively (Table 1).

When inspecting the effect of the combined genotypes for ATP7A and ATP7B on hepatic histological copper levels (Table 2), we appreciated that for both sexes ATP7B:c.4358A was associated with increased hepatic copper levels in an additive way. This effect is most clear in female dogs, leading to a significant difference between copper levels of dogs homozygous for ATP7B:c.4358A and any of the other dogs. Concurrent presence of ATP7A:c.980T attenuated the copper accumulation effect in both sexes. This effect was most notable in males, leading to a significant difference between dogs with and without the mutant allele of ATP7A (Table 2).

The heritability for hepatic copper score in the model including the co-variates sex, ATP7A:c.980C>T and ATP7B:c.4358G>A was calculated to be 0.42. Thus, the explained genetic variability by the two non-synonymous mutations is $(0.48-0.42)/0.48$ 12.5%, of...
which 4.3% could be attributed to \( \text{ATP7A}:\text{c.980C>T} \) and 8.2% could be attributed to \( \text{ATP7B}:\text{c.4358G>A} \).

\( \text{ATP7A}:\text{c.980C>T} \) resulted in an amino acid substitution from a threonine to an isoleucine in the third MBD of \( \text{ATP7A} \) at position 327 (Ensembl: ENSCAFT00000049745, \( \text{ATP7A:p.Thr327Ile} \)). \( \text{ATP7B}:\text{c.4358G>A} \) caused an amino acid substitution from an arginine to a glutamine in the C-terminus of the \( \text{ATP7B} \) protein at position 1453 (Ensembl: ENSCAFT00000006859, \( \text{ATP7B:p.Arg1453Gln} \)).

Alignment of the amino acid sequence adjacent to canine \( \text{ATP7A:p.Thr327Ile} \) (\( \text{ATP7AT327I} \)) and \( \text{ATP7B:p.Arg1453Gln} \) (\( \text{ATP7BR1453Q} \)) indicated strong conservation of the domains in mammals (Fig. 3A,B).

**Functional evaluation of \( \text{ATP7B}^{R1415Q} \)**

\( \text{ATP7B}^{R1415Q} \) is partly mis-localized to the endoplasmic reticulum

To analyse how the canine \( \text{ATP7B:p.Arg1453Gln} \) might affect copper homeostasis at the cellular level, we expressed the mutant protein exogenously in a cell line. To investigate the consequences of canine \( \text{ATP7B:p.Arg1453Gln} \) substitution on the subcellular localization of \( \text{ATP7B} \), the conserved arginine (\( \text{Arg} \)) at position 1415 (Ensembl: ENST00000242839) of human GFP-tagged \( \text{ATP7B} \) was replaced by a glutamine (\( \text{Gln} \)). Both human \( \text{ATP7B:p.Arg1415-GFP} \) (\( \text{ATP7B}^{WT} \)) and \( \text{ATP7B:Gln1415-GFP} \) (\( \text{ATP7B}^{R1415Q} \)) were expressed in HeLa cells and and treated with either bathocuproine disulphonate (BCS) or 200 \( \mu \text{M CuSO}_4 \). Confocal microscopy revealed that in low-Cu\(^{2+}\) conditions, \( \text{ATP7B}^{WT} \)-GFP exhibited significant overlap with the \text{trans}-Golgi marker TGN46 as reported before (Barnes et al., 2009), whereas stimulation with \text{CuSO}_4 induced \( \text{ATP7B}^{WT} \)-GFP trafficking to the cell surface and numerous cytosolic vesicular structures (Fig. 4A; Fig. S3). In contrast, a significant amount of \( \text{ATP7B}^{R1415Q} \)-GFP was detected in the endoplasmic reticulum (ER) network-like membranes in both low- and high-Cu\(^{2+}\) conditions (Fig. 4A; Fig. S3), although substantial amounts of the protein were delivered to the Golgi and post-Golgi compartments. Morphometric analysis confirmed an increase in the percentage of cells that exhibited GFP signal in the ER upon expression of \( \text{ATP7B}^{R1415Q} \), whereas the number of cells with \( \text{ATP7B}^{WT} \)-GFP in the ER was very low (Fig. 4C). Partial mis-localization of \( \text{ATP7B}^{R1415Q} \) to the ER compartment was also observed in an HepG2 cell line (Fig. 4B,C). This indicated that the substitution of arginine to glutamine impairs (at least partly) appropriate compartmentalization of \( \text{ATP7B} \).

\( \text{ATP7B}^{R1415Q} \) fails to reach the canalicular domain of polarized HepG2 cells

To verify whether \( \text{ATP7B}^{R1415Q} \) can be targeted to the apical (canalicular) domain of hepatic cells, which is necessary to remove...
excess Cu$^{2+}$ from hepatocytes into bile, we grew HepG2 cells in the conditions that allow their maximal polarization (Slimane et al., 2003). Then, the cells were transfected with either ATP7B<sup>WT</sup>-GFP or ATP7B<sup>R1415Q</sup>-GFP and subsequently incubated with 200 $\mu$M CuSO$_4$. Stimulation with copper resulted in delivery of ATP7B<sup>WT</sup>-GFP to the apical surface domains, which was labelled with the canalicular marker multidrug resistance protein 2 (MRP2) (Fig. 4B; Fig. S4). On the contrary, ATP7B<sup>R1415Q</sup>-GFP failed to arrive at the canalicular surface of the cells (Fig. 4B; Fig. S4) and remained mainly in the intracellular cytoplasmic structures, as it has been already demonstrated for another ER-retained mutant, ATP7B<sup>H1069Q</sup> (Polishchuk et al., 2014). These results indicated that the mutant cannot be delivered efficiently to regular copper-excretion sites in polarized hepatocytes.

**ATP7B<sup>R1415Q</sup> exhibits copper-transporting activity**

To assess the ability of ATP7B<sup>R1415Q</sup> to transport copper, we transfected HeLa cells with the metal-responsive element (MRE)-luciferase reporter, a copper sensor based on the metallothionein-1 promoter that responds to bioavailable cytosolic copper and that was previously used to characterize cellular copper import (van den Berghe et al., 2007). Co-expression of MRE-luciferase reporter with empty GFP vector resulted in strong induction of luciferase activity (Fig. 4D) upon exposure of the cells to CuSO$_4$. The induction of this copper-dependent reporter was significantly reduced when ATP7B<sup>WT</sup> was co-expressed in the cells, whereas transfection of the partly active ATP7B<sup>H1069Q</sup> mutant resulted in only modest inhibition of MRE-luciferase activation (Fig. 4D). Cells expressing ATP7B<sup>R1415Q</sup> exhibited substantial reduction of
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</tr>
<tr>
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<td>C</td>
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<td>0.41</td>
<td>0.2</td>
<td>0.039</td>
<td>0.27</td>
<td>0.48</td>
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Summary of identified mutations in ATP7A and ATP7B in 96 Labrador retrievers that were randomly selected from the cohort of 235 Labrador retrievers used for the GWAS. Effect estimates for the non-reference allele on copper score, corrected for sex effect, standard errors and P-values are reported. Significant mutations (P<0.05) were replicated in an independent replication cohort consisting of 59 Labrador retrievers. Significantly replicated mutations were genotyped in the entire cohort of 294 Labrador retrievers, and allele frequencies, effect estimates and P-values are reported in the last column. AA, amino acid; All freq., allele frequency; Chr., chromosome; CF 3.1, CanFam 3.1 genome build; non-ref., non-reference; ref., reference; s.e., standard error; SNP, single nucleotide polymorphism; indel, small insertion or deletion.
reporter activity (Fig. 4D), while expressing a similar amount of protein to ATP7BWT-transfected cells (Fig. S5). This observation indicates that ATP7BR1415Q is still capable of transporting copper out of the cytosol and that observed differences are not the result of differences in protein expression.

Taken together, our findings indicate that accumulation of copper upon ATP7BR1415Q expression occurs mainly as a result of failure of the mutant to reach the appropriate copper-excretion compartments rather than as a result of a reduction in its copper-translocating activity.

Table 2. Hepatic histological copper score in relation to ATP7A and ATP7B genotype in male and female Labrador retrievers

<table>
<thead>
<tr>
<th>ATP7B:c.4358G&gt;A genotypes</th>
<th>Males</th>
<th>Females</th>
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</thead>
<tbody>
<tr>
<td>AA</td>
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<tr>
<td>2.25 (0-4) a</td>
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<td>AG</td>
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<tr>
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<td>2 (0-3.5) z</td>
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<tr>
<td>GG</td>
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<td>n=11</td>
</tr>
<tr>
<td>1.25 (0-3.5) a,b</td>
<td>1.5 (0-3.5) z</td>
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</tbody>
</table>

<table>
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<th>Females</th>
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<td></td>
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<tr>
<td>2 (0-3.5) z</td>
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<td>CT</td>
<td>n=0</td>
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<tr>
<td>0.5 (0-2.5) z</td>
<td></td>
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</table>

Combinations of genotypes of individual dogs for ATP7A:c.980C>T and ATP7B:c.4358G>A are indicated. Data are presented for male dogs and female dogs separately. The number of dogs (n) and median and range (in parentheses) for hepatic copper score per genotype category are summarized. Significant differences between genotype categories are indicated by letters in males (a,b,c) and females (y,z). Genotypes for which fewer than five animals were present were omitted from the statistical analysis.

Taken together, our findings indicate that accumulation of copper upon ATP7B[R1415Q] expression occurs mainly as a result of failure of the mutant to reach the appropriate copper-excretion compartments rather than as a result of a reduction in its copper-translocating activity.

Fig. 3. Protein domains of ATP7A and ATP7B involved in copper toxicosis in Labrador retrievers. Overview of the ATP7A (A) and ATP7B (B) proteins with the N-terminus, metal-binding domains (MBDs), actuator domain (A), nucleotide-binding domain (N), phosphorylation domain (P) and the C-terminus. The green asterisk indicates the position of the mutations. (A) Alignment of the region containing ATP7A[T327h] (in green) shows a strong conservation of this amino acid position in the human, rat, mouse, cow, cat and horse. The copper-binding site XMXCXXC (boxed), predicted α-helix (red) and β-sheets (blue) are indicated. (B) Alignment of the region containing ATP7B[R1455Q] (in green) shows a strong conservation of this amino acid position in the human, rat, mouse, cow, cat and horse. The predicted α-helix (red) and β-sheets (blue) are indicated.
Canine dermal fibroblasts expressing ATP7A<sup>T327I</sup> accumulate more copper than wild-type fibroblasts

To ascertain whether ATP7A:9.Thr327Ile (ATP7AT327I) influenced cellular copper accumulation, a copper uptake and retention assay (Moller et al., 2011) was performed using copper 64 isotope (<sup>64</sup>Cu) in dermal fibroblasts.

Fibroblasts with ATP7AWT were derived from five female and three male dogs, and fibroblasts with the variant ATP7AT327I were...
derived from four female and three male dogs, and used for the $^{64}$Cu experiments. For the $^{65}$Zinc experiments, fibroblasts from three ATP7A$^{WT}$ dogs (two females, one male) and three ATP7A$^{T327I}$ dogs (one female, two males) were used. All dogs used for the fibroblast studies that were homo- or hemizygous for ATP7A: c.980C were homo- or hemizygous for the reference allele for the SNP in intron 18 (position 60338569), and all dogs that were homo- or hemizygous for ATP7A:c.980T were homo- or hemizygous for the non-reference allele.

The model for $^{64}$Cu accumulation in fibroblasts that included the fixed factors time and mutation had the lowest Akaike’s information criterion (AIC). Overall, a significant increase of copper at time points 22 and 30 h was present. The fibroblasts derived from the ATP7A$^{T327I}$ dogs accumulated significantly more copper than fibroblasts derived from ATP7A$^{WT}$ dogs (Fig. 5A).

To establish whether the differences in $^{64}$Cu levels between fibroblasts were the result of abrogated efflux, a chase method was used, after preloading of cells with copper for 22 h. The average export rate of fibroblasts with the ATP7A$^{T327I}$ variant was significantly lower compared with the average export rate in fibroblasts derived from ATP7A$^{WT}$ (Fig. 5B). Western blots of dermal fibroblasts derived from ATP7A$^{WT}$ and ATP7A$^{T327I}$ dogs did not show differences in ATP7A protein expression (Fig. S6).

As a control, fibroblasts from a human fibroblast cell line (ATP7A$^{WT}$) and a human Menkes disease-affected individual (Menkes; both n=1) were included in the experiments. Fibroblasts derived from the human with Menkes disease accumulated considerably more $^{64}$Cu than the ATP7A$^{WT}$ fibroblasts (Fig. 5C) and showed abrogated efflux (Fig. 5D).

Handling of an alternative trace metal in canine fibroblasts was not influenced by the ATP7A genotype. Incubation of fibroblasts with $^{65}$Zn resulted in similar patterns of accumulation of isotope over time (Fig. S7A), and the release kinetics of newly accumulated $^{65}$Zn were comparable between the fibroblasts derived from ATP7A$^{WT}$ dogs and dogs with ATP7A$^{T327I}$ (Fig. S7B). In summary, this experiment showed that fibroblasts derived from dogs with the ATP7A$^{T327I}$ accumulate more copper than fibroblasts derived from dogs with ATP7A$^{WT}$ and that copper accumulation might be the result of decreased copper excretion by ATP7A$^{T327I}$ fibroblasts.

**Copper-induced trafficking is not changed in cells expressing ATP7A$^{T327I}$**

To investigate whether the observed decrease in copper excretion was a result of aberrant trafficking, immunofluorescence studies were performed in canine dermal fibroblasts derived from ATP7A$^{WT}$ and ATP7A$^{T327I}$ dogs. Both variants reside in the...
Golgi in low-copper circumstances. After 2 h incubation with copper, both ATP7AWT and ATP7AT327I exhibited substantial redistribution from the Golgi to the small post-Golgi vesicles and plasma membrane (Fig. S8).

To understand whether basolateral targeting of ATP7AT327I was compromised, apicobasal distribution of both ATP7AWT and ATP7AT327I was investigated in filter-grown polarized Madin–Darby canine kidney (MDCK) cells. The MDCK cells were transfected with GFP-tagged versions of either ATP7AWT or ATP7AT327I and exposed to CuSO4 for 4 h. Confocal microscopy revealed efficient delivery of ATP7AWT to the lateral cell surface domain of MDCK located below the occludin-labelled belt of tight junctions. As did the wild-type protein, ATP7AT327I exhibited efficient targeting to the lateral walls of MDCK cells (Fig. S9), indicating that its polarized delivery in epithelial cells remained unaffected.

**DISCUSSION**

We performed a GWAS in 235 Labrador retrievers to identify causal genes for copper toxicosis. We identified two chromosomal loci that were positively and negatively associated with hepatic copper levels. The two loci respectively harboured the genes coding for the copper transporters ATP7B and ATP7A. Subsequent Sanger sequencing in 96 Labrador retrievers and validation of mutations in an independent cohort of 59 Labrador retrievers identified a non-synonymous SNP in ATP7B associated with an increase in hepatic copper levels, and a non-synonymous and intronic SNP in ATP7A associated with a decrease in hepatic copper levels. Functional follow-up studies for the non-synonymous mutations provided biological evidence for their involvement in a disturbed copper metabolism.

Copper toxicosis in Labrador retrievers and Wilson disease are similar with regard to the amount of hepatic copper accumulation, presence of iron accumulation and progression to liver cirrhosis. Electron microscopy of liver sections of affected Labrador retrievers revealed ultrastructural changes that are similar to those observed in humans with Wilson disease. We show, for the first time, that Labrador retrievers and humans with Wilson disease share ATP7B as the causal gene. However, histopathological differences between copper toxicosis in Labrador retrievers and Wilson disease are present and include predominant centrolobular copper accumulation in Labrador retrievers versus a perportal distribution in humans with Wilson disease. Furthermore, fatty degeneration and Mallory bodies are not recognized in liver histopathology from Labrador retrievers. In both diseases, the age of onset is usually in middle or older age, but neurological symptoms and Kaiser–Fleisher rings have not been recognized in Labrador retrievers.

Dietary intake of copper seems important in disease progression in Labrador retrievers (Fieten et al., 2014, 2012a) and, in this regard, Labrador retriever copper toxicosis is comparable to endemic Tyrolean infantile cirrhosis and Indian childhood cirrhosis. The observed differences among copper toxicosis diseases in humans and Labrador retrievers are intriguing, and further studies are needed to identify whether they can be attributed to general species differences in copper metabolism or to other, yet to be identified gene mutations in either species.

ATP7B is highly expressed in hepatocytes, where it is located in the trans-Golgi network in low-copper conditions. With increasing intracellular copper levels, ATP7B traffics to endo-lysosomal vesicular structures and to the canalicular surface of hepatocytes (Guo et al., 2005; Polishchuk et al., 2014; Schaefer et al., 1999) to facilitate copper excretion into the bile. The C-terminus has an important role in regulating copper-responsive trafficking. Mutations of residues in this region are suggested to contribute to aberrant Golgi retention of ATP7B in high-copper circumstances (Braiterman et al., 2011). Our fluorescence studies did not show a significant difference in GFP signals in both HeLa and polarized HepG2 cells between the wild type and ATP7BR1415Q after copper treatment. However, ATP7BR1415Q showed increased retention in the ER and failure to reach the canalicular surface in polarized hepatic cells in high-copper conditions, which might occur because of mis-folding of the protein. Mis-folding and aberrant retention in the ER have been previously reported in Wilson disease-associated ATP7B mutations (Gupta et al., 2011; van den Bergh et al., 2009). The luciferase assay demonstrated that the copper-transport capacity of ATP7BR1415Q showed only a modest impairment, indicating that the functional effect of ATP7BR1415Q in hepatic copper accumulation in Labrador retrievers results from mis-localization of the protein, rather than from severely decreased copper-transport capacity.

In Labrador retrievers, there is a strong female predisposition for disease (Fieten et al., 2012b). X-chromosomal mapping in a combined data set can be hampered by hemizygosity in males, and thereby, possibly cause disturbance of association signals. Therefore, we analysed males and females separately. In the region harbouring ATP7A, we identified a suggestive signal of association with lower hepatic copper levels in male dogs. We hypothesized that mutations in ATP7A might modify copper accumulation, and this gene was therefore analysed by DNA sequence analysis. The identified missense mutation ATP7A:c.980C>T results in the amino acid substitution ATP7A:p.Thr327Ile in MB3 of ATP7A and was negatively associated with hepatic copper levels. MB3 is located in the cytoplasm and has the ability to receive copper from the Atox1 chaperone (Strausak et al., 2003). MB3 of ATP7A is only metallated in elevated copper conditions (Banci et al., 2010) and is thought to be an important domain for regulating ATP7A activity via cell-signalling pathways (Veldhuis et al., 2011). It has been shown that ATP7A:p.T327 is an important phosphorylation site of ATP7A (Veldhuis et al., 2009).

To investigate whether ATP7AT327I influenced cellular copper accumulation, we studied copper uptake and retention in dermal fibroblasts derived from Labrador retrievers hemi- or homozygous for either ATP7AWT or ATP7AT327I. Mutant fibroblasts accumulated significantly more copper and showed a decreased efflux rate without a change in protein expression levels, indicating a functional impairment of ATP7A. We also investigated whether this could be a result of aberrant trafficking or sorting of the mutant upon copper stimulation. Here, we used wild-type and ATP7AT327I dermal fibroblasts and polarized MDCKs transfected with either ATP7AWT or ATP7AT327I. In both experiments, correct trafficking to the plasma membrane and delivery to the basolateral membrane (in polarized MDCKs) upon copper stimulation was observed. These results are in line with previous observations that the phosphorylation site ATP7A:pT327 is not involved in aberrant trafficking of ATP7A (Veldhuis et al., 2009), but is rather predicted to stabilize a conformational change in ATP7A, exposing the CxxC copper-binding region of MB3, which might affect catalytic activity of ATP7A (Veldhuis et al., 2011). This might indicate that catalytic activity, rather than aberrant trafficking of ATP7A, is responsible for the observed decrease in hepatic copper levels in Labrador retrievers.

In humans, disease-causing mutations in the third metal-binding domain of ATP7A were not reported (Kaler, 2011). For ATP7B, 15 (three silent, 12 Wilson disease-causing) variants in MB3 are
described. Two mutations, c.915T>A (Loudianos et al., 1998) and c.918-931del (Davies et al., 2008), lead to a premature stop and a concomitant absence of ATP7B:p.Ser306 (corresponding to ATP7A:p.Thr327). Both of these mutations cause Wilson disease.

Another single base-pair substitution, C>T, located in intron 18 (position 60338569, CF 3.1), was also inversely related to hepatic copper score. In the dogs from which the fibroblasts were derived, the SNP in intron 18 and ATP7A:c.980C>T were in complete LD, and therefore we cannot rule out the possibility that the intrinsic variation exerts an effect on ATP7A gene function; however, we consider it less likely. The alleles are both pyrimidines within a stretch of pyrimidines, are not situated close to an intron-exon junction, and the mutation is not expected to activate a cryptic splice site. In the total data set, the intronic SNP was in high LD ($r^2 = 0.92$, $D’ = 0.97$) with ATP7A:c.980C>T, making it likely that the observed association is a result of the fact that this SNP is in LD with ATP7A:c.980C>T.

$ATP7B$:c.4358G>A showed an additive effect on hepatic copper levels, which was most significant in female dogs (Fig. 2). Copper-accumulating effects of $ATP7B$:c.4358G>A were modified by the presence of ATP7A:c.980C>T, and this effect was most clear in male dogs (Table 2). In accordance with observations in humans with Menkes disease, where aberrant ATP7A function results in an impaired intestinal copper uptake, we suggest that in Labrador retrievers ATP7A327I might result in decreased intestinal copper uptake, thereby attenuating hepatic copper accumulation because of ATP7BR1453Q. Other possible mechanisms influencing copper homeostasis in Labrador retrievers and, more specifically, the interplay between ATP7A and ATP7B might include regulation by hormones, inflammatory cytokines, growth factors or a yet unidentified small molecule acting as a regulator of systemic copper metabolism (Kim et al., 2010).

In humans with Wilson disease, a wide variation in clinical symptoms and hepatic copper accumulation is noticed, which is currently unexplained (de Bie et al., 2007; Lee et al., 2011; Riordan and Williams, 2001; Shah et al., 1997). When we translate our observations in the Labrador retrievers to the human disease, we might think of ATP7A as a possible modifier gene in Wilson disease. Variations in ATP7A with a small effect on protein function might reduce the rate of copper accumulation in humans with Wilson disease, thereby modifying the disease phenotype, for example by delaying the age at onset of clinical symptoms. In fact, part of the apparent discrepancy between the genetic incidence of Wilson disease and the number of affected individuals diagnosed (Coffey et al., 2013) might be a result of a slower rate of copper accumulation that does not reach toxic levels in the life span of some individuals. In a recent pilot study, two SNPs in ATP7A were genotyped in humans with Wilson disease. No association was identified between the ATP7A polymorphisms and the presence of Wilson disease or clinical phenotypic parameters (Przybylkowski et al., 2014). However, these results clearly do not exclude ATP7A as a candidate modifier, because it was a very limited investigation of ATP7A and a small data set.

In conclusion, the present study illustrates the potential of canine inbred populations for the unravelling of complex hereditary diseases. We identified the involvement of the copper transporter ATP7B in copper toxicity in the Labrador retriever, identifying this breed as the first naturally occurring non-rodent mammalian model for Wilson disease. The concurrent identification of a functional mutation in ATP7A suggests a role for the Menkes gene as a disease modifier in copper-metabolism disorders. Owing to its long life span and large body size, the Labrador retriever will be an invaluable large-animal model for the development of new therapies, including gene therapy, beneficial to both human and canine affected individuals.

**MATERIALS AND METHODS**

**Dogs**

Samples for the initial GWAS and DNA sequencing study were collected between 2003 and 2009, and samples for the replication cohort were collected between 2009 and 2013 at the Faculty of Veterinary Medicine, Utrecht University, with informed consent of the owners. The procedures were approved by the Utrecht University Ethical Committee as required under Dutch legislation. In addition, procedures were reviewed and approved by the WALTHAM Centre for Pet Nutrition. Each dog underwent a physical examination. An EDTA blood sample of 4 ml was used for DNA extraction from peripheral white blood cells by a salt extraction procedure (Miller et al., 1988). For diagnostic purposes, liver biopsies were obtained by the Menghini technique, ultrasound guided with a Tru-cut device and 14 gauge needle, during laparoscopy or laparotomy (Rothuizen et al., 2006). Local anaesthetics (infiltration of the skin and subcutaneous tissues with lidocaine) were applied for Tru-cut liver biopsies. When necessary, dogs were sedated with propofol (dose to effect) during the liver biopsy procedure. For liver biopsies obtained by laparoscopy or laparotomy, dogs underwent general anaesthesia. The anaesthesia protocols used were individually designed based on the anaesthetic risk classification of the American Association of Anesthesiologists. Tissue specimens were embedded in paraffin, and 4 μm thick sections were mounted on glass slides. Sections were stained with haematoxylin and eosin, rubineac acid (Uzman, 1956) for copper and reticulin (Gordon and Sweerts, 1936). All samples were evaluated and graded by one board-certified pathologist (T.S.G.A.M.v.d.I.) according to World Small Animal Veterinary Association standards (Van Den Ingh et al., 2006). Hepatic copper accumulation was histologically graded on a scale from zero to five as described previously (Fieten and Rothuizen, 2014).

**Electron microscopy**

For routine electron microscopic analysis, liver specimens were obtained from two Labrador retrievers affected with copper toxicosis and one healthy dog of mixed breed. Biopsies were fixed in 1% glutaraldehyde in 0.2 M HEPES buffer. Small blocks of liver tissue were post-fixed in uranyl acetate and in OsO4. After dehydration through a graded series of ethanol, the tissue samples were cleared in propylene oxide, embedded in the Epoxy resin (Epon 812) and polymerized at 60°C for 72 h. From each sample, thin sections were cut with a Leica EM UC7 ultra-microtome. Thin sections were investigated further using a FEI Tecnai-12 (FEI; Eindhoven, The Netherlands) electron microscope equipped with a Veleta CCD camera for digital image acquisition.

**Genome-wide association study**

DNA samples from 235 Labrador retrievers were genotyped using Illumina’s Canine HD BeadChip, containing ~170,000 SNPs. Data were analysed in the R-package (R Development Core Team, 2011) ‘GenABEL’ version 1.6-6 (Aulchenko et al., 2007). Quality-control thresholds were set using the function ‘check.marker’. A threshold of 98% successful genotypes per individual and 98% successful genotypes per SNP was used. SNPs for which at least 20 heterozygotes were typed were included in the analysis. Furthermore, a check for sex mismatch (based on genotypes at the X chromosome), duplicate sample checks and checks for heterozygosity outiers were performed. The phenotype copper score was analysed as a quantitative trait, and sex was modelled as a fixed factor. Stratification and cryptic relatedness were adjusted for by mixed models implemented in the function ‘mmscore’ (Chen and Abecasis, 2007). A correction for residual genomic control was applied. The heritability of hepatic copper score with correction for sex was calculated based on genotype kinship matrix with the function ‘polygenic_hglm’ based on the packages ‘hglm’ (Ronnegard et al., 2010) and ‘GenABEL’. The autosomes and pseudo-autosomal region of the X chromosome were analysed in male and female dogs together. Another additional stratified analysis for males and females was performed to facilitate inspection of the X chromosome.
Genome-wide suggestive association criteria were set to an uncorrected P-value of $P \leq 5 \times 10^{-5}$. Boundaries for the crucial intervals were determined for a significance level of $P \leq 5 \times 10^{-4}$. Crucial intervals were screened for the presence of possible candidate genes.

**DNA sequence analysis and validation**

Exons and intron-exon boundaries from positional candidate genes were selected for Sanger sequencing in 96 individual dogs randomly selected from the 235 Labrador retrievers used in the initial GWAS. Variants that were significantly associated with hepatic copper level were genotyped in an independent replication cohort of 59 Labrador retrievers and in all 294 dogs. Statistical analysis was performed with R version 2.11 (R Development Core Team, 2011). A linear regression model with sex as a covariate was used to test the association of the mutations identified with Sanger sequencing in the candidate genes with hepatic copper score. The mutations were modelled in an additive way.

Genotypes for the mutations that were significant after statistical testing in the replication cohort were genotyped by Sanger sequencing in the total cohort of 294 dogs. Differences in hepatic copper scores with respect to combined $ATP7A$ and $ATP7B$ genotype were determined using a Wilcoxon rank-sum test within males and females separately.

Genotypes for $ATP7B$ c.4358G>A and $ATP7A$c.980C>T were added as independent covariates, and heritability was calculated by using the function ‘polycigen_hglm’ (Ronnegard et al., 2010) in GenABEL. The explained genetic variability was calculated by the following formula: (heritability of model without mutations as a covariate minus heritability including mutations as covariates/heritability without mutations as a covariate). In all heritability calculations, sex was added as a covariate.

**Functional evaluation of $ATP7B$R1453Q**

Alignment of human and canine $ATP7B$ protein sequence confirmed that the Arg at position 1453 in the canine protein corresponds to the Arg at position 1415 in the human $ATP7B$. The human $p$Adlox-$ATP7B$-GFP construct was used, and site-directed mutagenesis was performed to make $p$AdloxATP7B-p:Arg1415Gln-GFP. Both constructs were used for cell transfection and subsequent immunofluorescence and luciferase assays.

**DNAs and site-directed mutagenesis**

To prepare $ATP7B$-p:Arg1415Gln, the $p$Adlox-$ATP7B$-GFP construct (provided by S. Lutsenko, Baltimore, MD, USA) was used as the template, and site-directed mutagenesis was performed according to the manufacturer’s instructions for point mutagenesis using the QuikChange II XL site-directed mutagenesis kit (Stratagene, La Jolla, CA, USA), using the following oligonucleotides: 5′-GGGACTCCCCCCAGGCCACACCATG-GGACC-3′ and 5′-GGGACTCCCCCCAGGCCACACCATG-GGACC-3′. Given that $ATP7A$ sequences are unstable in high-copy plasmids (reference PMID: 9571140), we used commercial DNA synthesis to generate silent mutations in the mouse $ATP7A$ because it is one of the most common Wilson disease mutations in the Caucasian population and proved to be functionally active. HeLa cells were plated in a 12-well plate and transfected with pGL3-E1b-TATA-4MRE reporter (van den Berge et al., 2007; provided by Bart van de Sluis, Groningen, The Netherlands) and then co-transfected with either empty vector (pEGFP-c1) or $p$AdloxATP7B-GFP or $p$AdloxATP7B-H1069Q-GFP or $p$AdloxATP7B-p:Arg1415Gln-GFP expression plasmids using the TransIT™-LT1 transfection protocol. After 24 h, cells were treated with 200 $\mu$M CuSO$_4$ for 24 h. Cells were subsequently harvested in a Passive Lysis Buffer (Promega, Madison, WI, USA) according to the manufacturer’s instructions. Firefly luciferase and Renilla luciferase activities were measured with a Dual-Luciferase® reporter assay system (Promega) on a GloMax® 96 Microplate Luminometer (Promega) according to the manufacturer’s instructions. Relative light units were calculated by dividing firefly measurements by Renilla measurements.

All values were normalized to the $ATP7B$ protein expression levels determined by ImageJ (Collins, 2007) analysis of western blot in the respective specimens. Western blot analysis was performed on lysates obtained by transfection of the cells with pEGFP-c1, $p$Adlox-$ATP7B$-GFP, $p$Adlox-$ATP7B$-H1069Q-GFP and $p$Adlox-$ATP7B$-p:Arg1415Gln-GFP expression plasmids. Lysates were prepared via the addition of lysis buffer (0.5% Triton X-100, 20 mM Tris-HCl (pH 7.4), 150 mM NaCl, 1 mM EDTA (pH 8), 0.5% NP-40, 10% glycerol and 1× protease inhibitor cocktail (Sigma S882011) for 10 min at room temperature. The mixture was placed into a microfuge tube, kept on ice for 10 min, and then spun at 17,000 $g$ for 15 min at 4°C in a microcentrifuge. The quality of transfer was verified by Ponceau staining, and molecular weights were inferred by comparison to prestained markers (New England Biolabs, Ipswich, MA, USA).

**Functional evaluation of $ATP7A$T327I**

Dermal fibroblast cells were derived from Labrador retrievers selected for the presence or absence of an $ATP7A$c.980C>T substitution resulting in the $ATP7A$T327I amino acid substitution in homozygous or hemizygous form. 
The the human dermal fibroblast cell HB156 from an individual with clinical Menkes disease (cell line: DD0355, no. 91071704) were purchased from the European Collection of Cell Cultures (ECACC). ECACC is managed as part of the Health and Safety Executive (HSE) in the UK. The human HDFn dermal fibroblast line (Invitrogen, Bleiswijk, The Netherlands) was used as a control cell line.

Fibroblast cultures were routinely maintained in DMEM (Invitrogen) supplemented with 1-glutamine (2 mM), 10% FBS, MEM non-essential amino acids (100 µM) and penicillin-streptomycin at 37°C in a humidified atmosphere of 5% CO2 in air. For experiments, canine cells were used between passage 4 and 9.

Radioisotope experiments

Methods for radioisotope experiments were based on those described previously (Speer et al., 2007). In short, 64Cu was prepared using metallic copper wire (1 mg) irrigated overnight to provide an induced activity of approximately 16.6 Ci/g (Reactor Institute, Delft, The Netherlands). Four hours after irrigation and 30 min before the start of the study, the copper wire was dissolved in 100 µl 10.3 M HNO3 and subsequently neutralized with 0.5 M NaOH. DMEM was added to give a final concentration of 10 mM copper. Fibroblasts were cultured from skin-punch biopsies, obtained with local anaesthesia (lidocaine infiltration).

Fibroblast cells were plated in triplicate in six-well plates in DMEM+10% FBS. Cells were incubated with 3 µM 64Cu for 6, 22 or 30 h. Culture medium was removed after treatment, and the cells were washed four times with Hanks solution containing 3 µM CuCl2. Following wash steps, 325 µl of 0.2% SDS was added to lyse the cells. Cellular copper loading was determined from counts of the cell lysate (450 and 800 keV for 1 min; Packard B5003 gamma counter; Packard BioScience Benelux, Groningen, The Netherlands). For efflux studies, after copper treatment for 22 h as described above, cells were washed four times with Hanks solution containing 3 µM CuCl2 before incubation in fresh medium containing no additional copper for 8 h. Cellular copper loading was assessed at 0, 0.5, 1.5, 3, 6 and 8 h as described in the previous paragraph. Copper loading was normalized to the protein concentration of the cell lysate (Bio-Rad Protein Assay Kit; Bio-Rad, The Netherlands). 65Zn (PerkinElmer; 1 mCi chloride) was added to the cells was incubated with blocking/permeabilizing solution (0.5% bovine serum albumin, 0.1% saponin and 50 mM NH4Cl in PBS) for 20–30 min. Primary and secondary antibodies were diluted in blocking/permeabilizing solution and added to the cells overnight and for 45 min, respectively.

Co-localization between ATP7B variants and organelle markers was visualized using anti-TGN46 antibody (Golgi marker), VAP-A antibody (ER marker), Na+K+-ATPase antibody (plasma membrane marker) and LAMPI antibody (late endosome/lysosome marker) followed by the Alexa Fluor 568 secondary antibody. Antibodies against the tight junction protein occludin were used to label the border between apical and basolateral domains in MDCK cells. To visualize canalicular domains, polarized HepG2 cells expressing ATP7BWT or ATP7B1415Q were labelled with anti-MRP2 antibody followed by the secondary anti-mouse Alexa Fluor 568. Canine dermal fibroblasts were treated with BCS or CuSO4 as described above, fixed with 2.5% glutaraldehyde, and then stained with a solution containing 2.5% osmium tetroxide and 0.1% ruthenium red for 1 h and 90 min, respectively. Canine fibroblasts were then fixed in 2.5% glutaraldehyde and 1% osmium tetroxide, washed and dehydrated in a graded series of ethanol followed by propylene oxide. Prepared cells were embedded in Durcupan (TAAB) and sectioned (80 nm) using an ultramicrotome (Ultratome IV, Leica). Sections were post-stained with lead citrate and uranyl acetate and viewed with a transmission electron microscope (Zeiss EM 912, Jena, Germany).

Immunofluorescence

The intracellular distribution of GFP-tagged ATP7B or ATP7A proteins was observed 24 or 48 h after transfection. Cells were treated with 500 µM BCS overnight and fixed directly or washed and incubated for 2 h with 200 µM CuSO4 before fixation in 4% paraformaldehyde. Then the cells were incubated with blocking/permeabilizing solution (0.5% bovine serum albumin, 0.1% saponin and 50 mM NH4Cl in PBS) for 20–30 min. Primary and secondary antibodies were diluted in blocking/permeabilizing solution and added to the cells overnight and for 45 min, respectively. Staining for organelle markers was visualized using anti-TGN46 antibody (Golgi marker), VAP-A antibody (ER marker), Na+K+-ATPase antibody (plasma membrane marker) and LAMPI antibody (late endosome/lysosome marker) followed by the Alexa Fluor 568 secondary antibody. Antibodies against the tight junction protein occludin were used to label the border between apical and basolateral domains in MDCK cells. To visualize canalicular domains, polarized HepG2 cells expressing ATP7BWT or ATP7B1415Q were labelled with anti-MRP2 antibody followed by the secondary anti-mouse Alexa Fluor 568. Canine dermal fibroblasts were treated with BCS or CuSO4 as described above, fixed and stained for endogenous ATP7A with anti-ATP7A antibodies and with anti-Golgin97 to reveal TGN46. The images were acquired using a 63×1.4 NA oil immersion objective at an LSM710 or LSM700 confocal microscope with appropriate filter sets (Carl Zeiss, Jena, Germany). The intracellular distribution of GFP-tagged ATP7B protein was analysed in 10 randomly selected fields, and the percentage of the cells exhibiting ATP7B in ER, Golgi and/or plasma membrane vesicles was quantified. A Wilcoxon rank-sum test was used to test differences in percentages of GFP positivity within the ER, Golgi and/or plasma membrane. A P-value <0.05 was considered significant. For polarized MDCK cells, confocal z-stacks through the entire cell monolayer were taken, and xz sections were generated using Zeiss Zen software.

Statistical analysis

Counts per minute normalized to the protein concentration of the cell lysate (in counts per minute per milligram) were analysed using linear mixed-effect modelling with the R package ‘lme4’. The outcome variable was transformed by taking its natural logarithm in order to fulfill the criteria of normality and constant variance of residuals. Restricted maximum likelihood was used to estimate the fixed effects of duration and their interactions. A stepwise backward method was used to determine the model of best fit, based on Akaike’s information criterion. Estimates were reported with their 95% confidence intervals. For the efflux studies, the amount of 64Cu at the start of the experiment was set to 100%, and at the time points 0.5, 1.5, 3, 6 and 8 h the amount of 64Cu was calculated as a percentage of the initial value. P-value <0.05 was considered significant.

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**Supplementary information**
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