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A liver-targeting Cu(I) chelator relocates Cu in hepatocytes and promotes Cu excretion in *ATP7b*-/- mice

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Electronic Supplementary Information (ESI) available: detailed experimental section. Supplementary nano XRF images (Fig. S1 and S2). Supplementary data for the ⁶⁵Cu experiments on mice (Table S1 and Fig. S3)]. See DOI: 10.1039/x0xx00000x

Abstract.

Copper chelation is the most commonly used therapeutic strategy nowadays to treat Wilson's disease, a genetic disorder primarily inducing a pathological accumulation of Cu in the liver. The mechanism of action of Chel2, a liver-targeting Cu(I) chelator known to promote intracellular Cu chelation, was studied in hepatic cells that reconstitute polarized epithelia with functional bile canaliculi, reminiscent of the excretion pathway in the liver. The interplay between Chel2 and Cu localization in these cells was demonstrated through confocal microscopy using a fluorescent derivative and nano X-ray fluorescence. The Cu(I) bound chelator was found in vesicles potentially excreted in the canaliculi. Moreover, injection of Chel2 either intravenously or subcutaneously to a murine model of Wilson's disease increased excretion of Cu in the faeces, confirming in vivo biliary excretion. Therefore, Chel2 turns to a possible means to collect and excrete hepatic Cu in the faeces, hence restoring the physiological pathway.

Introduction.

A possible involvement of metal ions has been brought to the forefront in many neurodegenerative diseases without knowing whether the abnormal presence of metal ions is the cause or a consequence of the disease, for instance in Alzheimer's disease. In some cases however, when the homeostasis of an essential metal ion is disrupted by a genetic defect, the excess of the given metal is known to be responsible for the disease; this is the case for iron in hemochromatosis and aceruloplasminemia, manganese in hypermanganesemia or copper in Wilson's disease. Systemic metal chelation is the usual therapy for most of these diseases, in order to lower the dietary absorption of the metal ions and, hopefully, to deplete the excess stores by excreting the metal ions, often in urine. One striking observation is that these diseases promote a high increase in the liver content of the metal with disrupted homeostasis. Some metal ions also promote neurological symptoms with abnormal magnetic resonance images of the brain.

Wilson's disease is an autosomal recessive disease resulting from mutations of the ATP7B gene, whose product is a Cu(I)-transporting ATPase, a membrane protein that pumps copper out of the cytoplasm.

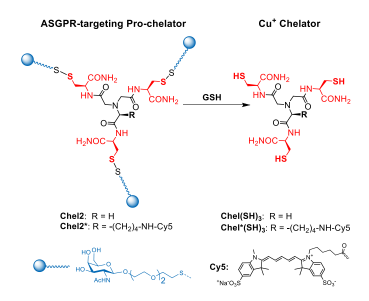
Under basal condition, the so-called Wilson protein is in the trans-Golgi network for dispensing copper to enzymes that use it as a co-factor. In case of excess copper in the cell, the Wilson protein changes its localization for pumping copper into intracellular vesicles that will eventually excrete it out of the cells. The *ATP7B* gene is expressed mainly in the liver and the brain and to a lesser extent in kidney, lung, placenta, mammary gland and intestine. Because vertebrates get rid of the excess copper by transferring it into the bile and then the faeces, the Wilson protein is responsible for the detoxification of the whole organism. Therefore, mutations in *ATP7B* genes primarily induce a pathological accumulation of copper in the liver. Over the years, the liver undergoes fibrosis, cirrhosis and in some cases, acute hepatitis that requires a liver transplantation for the patient to survive. In addition, about 40% of the Wilson's disease patients develop neurological symptoms and 20%, psychiatric symptoms with no obvious signs of chronic liver disease. Among the patients with a neurological presentation, those who underwent a liver transplantation generally improved their condition. Their liver was loaded with copper, even though these patients did not necessarily exhibit typical signs of a liver condition.

The heterogeneous clinical presentations and the late onset of the neurological signs make Wilson's disease difficult to diagnose from scratch, even though there is a clear MRI signature in the brain and many patients have a large copper overload at the time of diagnosis. The chelators prescribed all over the world are d-penicillamine and triethylenetetramine, two molecules that allow some copper removal of the body. Unfortunately, they also have adverse effects which preclude their use, in up to 30% of the cases for d-penicillamine. One major drawback of these molecules is that they are systemic and used at high dosage. They are only moderately specific for the excess copper stored in hepatocytes, especially when one has in mind the very high affinity and specificity found in proteins devoted to copper homeostasis. 12

Given these observations, we have developed in the past decade bio-inspired chelators mimicking the Cu(I) binding sites from two cysteine-rich proteins involved in copper homeostasis. ^{12,13} One chelator mimics a metallochaperone binding site with a CuS₂ coordination. ¹⁴ and the other one mimics a metallothionein binding site with a CuS₃ coordination. ^{15,16} To protect the Cu(I) binding sites until they reach their target, the chelator's thiol functions have been oxidized into disulphide bonds, so that Cu(I) chelation would be activated only once inside the cells. Indeed, intracellular glutathione is millimolar and ensures the reduction of the disulphide bonds. ^{14,17} Finally, because Wilson's disease starts in the hepatocytes, the so-designed pro-chelators were grafted with ligands that specifically recognize the asialoglycoprotein receptors (ASGPR) at the hepatocyte sinusoidal membrane. ¹⁸ The design and the efficiency of the final molecules, i.e. the hepatocyte targeting pro-chelators, have been described earlier. ¹⁹ These studies, performed on hepatocyte-derived cell lines, show that both ASGPR-targeting pro-chelators release an efficient Cu chelator inside the cells, suggesting these compounds were models for the development of new hepatocyte- and Cu(I)- specific chelators. However, the intracellular mechanism of action of the chelators and the fate of the Cu(I) chelator complexes are still to be determined.

The work reported here aims at understanding the mechanism of action of Chel2 (Scheme 1), the hepatocyte-directed pro-chelator releasing a tristhiolate chelator in the cells.¹⁷ a sulphocyanin5 labelled version of Chel2 and on the effects of Chel2 on the intracellular distribution of metal ions, following their X-ray fluorescence at the sub-micron scale (nano XRF) thanks to the ESRF synchrotron beam in Grenoble. Based on these studies, we propose a mechanism of action for the pro-chelator

which would result in the transfer of the Cu(I)-chelator complex into the bile. This mechanism was further tested on *ATP7b*-/- mice, an animal model of Wilson's disease which develops the hepatic form of the disease.²⁰ To this end, *ATP7b*-/- mice were administrated ⁶⁵Cu which accumulated in their liver. We found that a treatment with Chel2 increased the ⁶⁵Cu content of the mice faeces, in agreement with partial restoration of the physiological pathway for Cu excretion through the bile.



Scheme 1 Structures of the glycoconjugates Chel2 and Chel2*and the released chelators Chel(SH)₃ and Chel*(SH)₃.

Results.

Chel2 has been shown to release in WIF-B9 cells the high affinity Cu(I) chelator, Chel(SH)₃ (scheme 1).¹⁷ The latter is a tripodal sulphur chelator based on a nitrilotriacetic core grafted with three cysteine moieties that chelate Cu(I) with properties reminiscent of metallothioneins. The complex, denoted ChelS₃Cu, exhibits a CuS₃ coordination and an affinity constant as large as 10¹⁹.^{16,21} In the hepatocytetargeting pro-chelator Chel2, the three thiol functions are hidden in disulphide bridges carrying the targeting units, namely three polyethylene glycol chains grafted with N-acetyl galactosamine moieties to get an architecture perfectly adapted for the interaction with the ASGP-R.¹⁹

The hepatocyte-derived WIF-B9 cells are able to reconstitute remarkably stable and polarized epithelia with functional bile canaliculi, indicated by asterisks in Fig. 1.²²

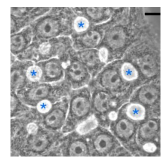


Fig. 1 Phase contrast image of living WIF-B9 cells forming bile canaliculi (phase lucent structures *), scale bar $10 \mu m$.

WIF-B9 cells provide a means to follow the detoxification function of the Wilson protein in case of excess copper. Indeed, the latter protein, localized at the trans-Golgi network membrane under basal condition, migrates to the canalicular membrane in case of excess copper.²³ Hence, the localization of the Wilson protein, as detected by immunofluorescence, can be used to evaluate whether the intracellular Cu concentration is basal or in excess.²³ Under conditions where ATP7B colocalized with an apical membrane marker (*i.e.* the WIF-B9 canalicular membrane), addition of Chel2 into the culture medium actually suppressed the need for detoxification, as illustrated by the re-localization of ATP7B back to the trans-Golgi network.^{14,17} These experiments demonstrate that the ASGPR-targeting prochelators are indeed doing what they were designed for, namely intracellular copper chelation. While performing these experiments, we found that 1 μM copper added to the WIF-B9 culture medium for 2 h is enough to induce the change in the ATP7B localization. Therefore, we could avoid the high concentrations used in previous studies on the localization of the ATP7B protein in WIF-B9 cells and other cell lines such as HepG2, HuH-7, CHO or HEK, where 20 to 200 μM Cu was used.²³⁻²⁷ In the experiments reported hereafter, added Cu was either 1 or 15 μM. They were designed to study the localization of the Cu(I) chelator in hepatic cells and its effect on intracellular Cu distribution.

Chel2* localization in WIF-B9 cells.

Endogenous asialoglycoproteins such as asialo-ceruloplasmin, -fetuin or -orosomucoid, are known to be cleared from the circulation after binding to ASGPR,²⁸ a receptor of the hepatocyte sinusoidal membrane. In the rat liver, half a million ligand-binding sites per cell allow asialoglycoprotein-ASGPR complexes to undergo endocytosis.^{29,30} Internalization is rapid (4-5 min at 37 °C)³¹ and followed by a separation of ASGPR-containing vesicles that are found at the Golgi complex and ligand-containing vesicles that prefigure lysosomes.³²⁻³⁴ The receptor is recycled by returning to the sinusoidal membrane, whereas the majority of the asialoglycoproteins is degraded in lysosomes. Yet, 1-4% of the asialoglycoproteins is kept intact and secreted into the bile across the canalicular membrane, a phenomenon called transcytosis that primarily depends on the protein.^{31,35,36}

To get further insights into the fate of Chel2, we designed Chel2*, an analogue compound bearing a red fluorescent tag.¹⁹ In Chel2* the sulphocyanin5 red fluorescent tag is tethered to the nitrilotriacetic acid scaffold through an amide bond.¹⁹ When the disulphide bridges are cleaved by the reducing medium of the cell, namely glutathione, the red-fluorescent tag is still attached to the released chelator Chel*(SH)₃ (see Scheme 1). Because we do not know exactly at which stage the chelator is released, the red fluorescence in confocal microscopy was assigned either to the whole molecule (Chel2*), the free chelator Chel*(SH)₃ or to the copper-chelator complex Chel*S₃Cu, depending on the redox state of the molecule and the presence of Cu.

After 2 h of incubation with 1 μ M Cu, 1 μ M Chel2* was added to the culture medium for 30 min, 1 h or 2 h. Various compartments identified during endocytosis were marked by green fluorescence and co-localization was analysed on confocal laser scanning microscopy images, assigning a yellow colour to the co-localization pixels (Fig. 2). After 30 min (Fig. 2A), Chel2* was found in early endosomes (EEA1), close to the nuclei and in late endosomes or lysosomes (LysoTrackerTM and Lamp1), close to

the bile canaliculi, but not in the trans-Golgi network (P58K). The latter finding confirms that Chel2* and ASGPR were segregated at that time.³⁷ There was still some co-localization with late endosomes after 1 h (Rab7, Fig. 2B), and with lysosomes/late endosomes after 2 h (LysoTrackerTM and Lamp1, Fig. 2C).

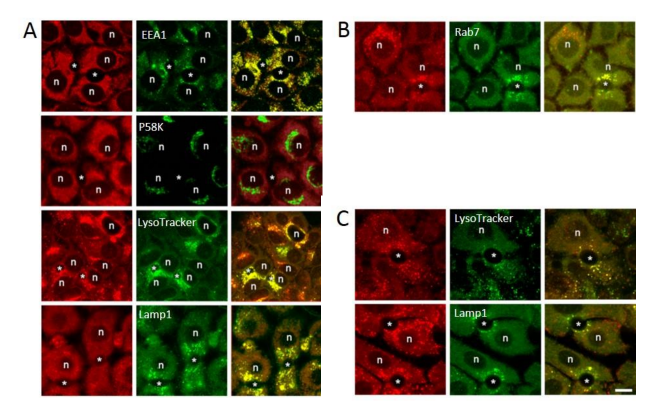


Fig. 2 Fluorescence images obtained by confocal microscopy of WIF-B9 epithelia forming canaliculi (*) and incubated for 2 h with 1 μM Cu and for (A) 30 min, (B) 1 h, (C) 2 h with 1 μM Chel2*. Left panels, red fluorescence of Chel2*; middle panels, green immunofluorescence of various protein markers or fluorescence of LysoTrackerTM added 1 h before Chel2*; right panels, colocalization analysis, the yellow false colour is assigned to the colocalized pixels. EEA1: early endosomes, P58K: trans-Golgi network, LysoTrackerTM: lysosomes, Lamp1: late endosome/ lysosome, Rab7: late endosomes. (n) nucleus, scale bar 10 μm.

A remarkable feature of Chel2* is that we did not observe a uniform distribution of red fluorescence in the cytosol. The red fluorescence always appeared as spots of about 200 nm, suggesting it was trapped in vesicles on the endocytosis/transcytosis pathway.

Chel2* reaches bile canaliculi.

At this point, special attention was paid to the search for Chel2* in the canaliculi after 1 to 3 h. To this end, we looked for canaliculi surrounded by Chel2* and analysed the fluorescence intensity profile across these canaliculi. Examples of such images are shown in Fig. 3, where the canaliculus is defined thanks to the interferential profile in panels B and D. After 1 h incubation, Chel2* detected from the red fluorescence profile is found at the border of the canaliculus close to the cell apical membrane (Fig. 3 A-B) and after 3 h incubation, Chel2* is found in the middle of the canaliculus (Fig. 3 C-D).

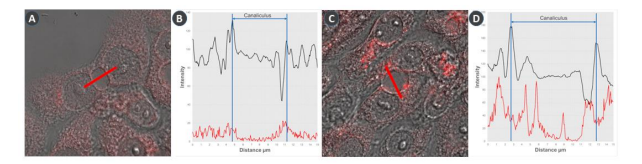


Fig. 3 Detailed analysis of Chel2* position in the region of the canaliculus. Analysis from interferential contrast and fluorescence images obtained by confocal microscopy of WIF-B9 epithelia forming canaliculi and incubated with 1 μ M Chel2* for (A-B) 1 h or (C-D) 3 h. (A and C) merged images of interferential contrast and red fluorescence. The canaliculi are crossed by a 15- μ m red line along which interferential contrast and fluorescence intensities are analysed. (B and D) interferential profiles in black and red fluorescence profiles along the line crossing the canaliculus.

Copper intracellular distribution by nano XRF.

To get further insights into the fate of the chelator and the Cu(I)-chelator complexes, the WIF-B9 cells were analysed for their intracellular metal distribution, paying special attention to the bile canaliculi and the cytoplasmic region close to the canalicular membrane.

Nano XRF maps were performed on WIF-B9 cells forming a polarized epithelium and submitted to different conditions, starting with the basal culture medium, adding Cu for 2 h and then adding Chel2 for 3 h (Fig. 4). The nano XRF images shown in Fig. 4 are representative of the 2-5 canalicular regions scanned under each condition (Fig. S1 in the Electronic Supplementary Information), bile canaliculi are indicated by asterisks. The P distribution covers the nuclei and the Zn distribution indicates the nuclei showing nucleoli as holes. Comparison with the phase image (Fig. 1) shows that the WIF-B9 organization and polarization were preserved by the sample preparation and the scans. The image obtained under the basal condition (Fig. 4A) shows two cells sharing a bile canaliculus and the Cu distribution is homogeneous across the cytoplasm with only scarce hot spots. After 2 h in the presence of 1 μ M Cu, the Cu distribution shows a lot more hot spots in the three cells forming a canaliculus (Fig. 4B). Under the next condition, 5 h in 1 μ M Cu including 50 μ M Chel2 for the last 3 h, the Cu distribution appears concentrated along the apical membranes forming the canaliculus (Fig. 4C).

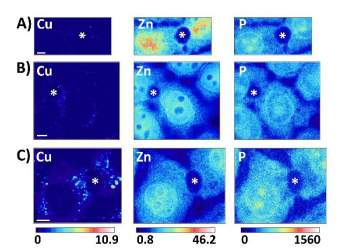


Fig. 4 Cell content analysed by nano XRF in WIF-B9 epithelia forming canaliculi under 3 different conditions: (A) basal Cu, (B) after 2 h with 1 μ M Cu and (C) after 2 h with 1 μ M Cu followed by addition of 50 μ M Chel2 for 3 h. Under each condition, the Cu, Zn and P maps are shown. Scale bar 5 μ m. The pixel size is 150x150 nm². Intensity scales are given in ng/cm².

All the scanned regions, whatever the condition, showed empty canaliculi, although their structure, and thus the cell organization, was preserved. The explanation for this observation was found by checking the distribution of abundant and loosely bound metal ions such as potassium or calcium. Actually, none of them was detectable, because the cryofixation step did permeabilize the cell membranes. Indeed, despite our efforts we did not succeed in finding a cryofixation method of WIF-B9 epithelia that would neither modify their membrane permeability nor break the canaliculi structure.

The scans were next used to quantify the amount of Cu per surface unit. There was no significant change in whole scan Cu areal density (Fig. 5A) going from the basal condition towards 1 μ M Cu for 2 h; however, a clear-cut increase was observed in the presence of 15 μ M Cu for 2 h (p=0.002). There was no further change in whole scan areal density when the cells were left for 3 more hours with 15 μ M Cu. Some nano XRF images acquired with 15 μ M Cu incubation are shown in Fig. S2 in the Electronic Supplementary Information. These observations confirm that in the preparations, Cu homeostasis was active, the Wilson protein pumping Cu towards the bile canaliculus to protect the cell from extra Cu accumulation. A 50-fold excess of Chel2 in the presence of 1 μ M Cu induced a significant increase in whole scan areal density of Cu (p<0.001). Bearing in mind that under these conditions, the Wilson protein is back at the trans-Golgi network as if the condition was basal, ¹⁷ the extra Cu measured must be chelated or hidden from the intracellular medium. This shows that Chel2 has released the chelator in the cells (Chel(SH)₃, Scheme 1) and that the latter efficiently binds the excess Cu, becoming ChelS₃Cu and thereby allowing more Cu to enter the cells from the culture medium.

The Cu areal density was also evaluated for regions of interest in the cells treated with 1 μ M Cu, the nucleus and the canaliculus vicinity, denoted 'at bc'. The latter was defined as a crown around the canalicular membrane that was counted separately from the rest of the cytoplasm. The 'at bc' area accounts for 10.8 \pm 1.1% of the cytoplasm area or 5.8 \pm 0.4% of the whole scan area (mean \pm SEM, n=16). Analysis of the Cu distribution in the cells (Fig. 5B) shows that after 2 h with 1 μ M Cu, only the 'at bc' region shows a significant increase (p=0.033). In the presence of Chel2 however, the cytoplasm has also an increased areal density. Interestingly, the most important Cu increase is found in the canaliculus region (p<0.001), demonstrating a Cu relocalization upon Chel2 treatment.

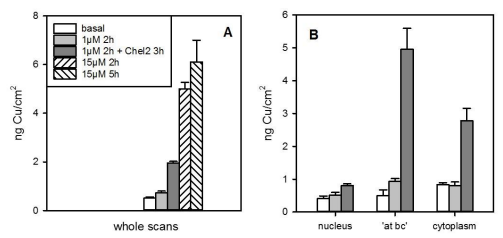


Fig. 5: Effects of Cu and Chel2 on the Cu areal density in WIF-B9 cells (mean and SE, standard error of the mean). In absence of Chel2, Cu was added to the culture medium at the concentration indicated in A. 50 μ M Chel2 was added 2 h after Cu and left for 3 h. (A) Cu areal density of the whole scans; (B) Cu areal densities in the 3 subcellular regions, as indicated

Faecal copper excretion in ATP7b^{-/-} mice.

Assuming that the relocalization of Cu seen in the nano XRF experiments reflects Cu excretion in the bile canaliculi due to Chel2 transcytosis, we next designed an experiment on *ATP7b*-/- mice to determine whether excess Cu could be found in their faeces after a treatment with Chel2. This section describes experiments performed on the *ATP7b*-/- mice kindly provided by Professor Svetlana Lutsenko.²⁰

Tracing Cu excretion is feasible using Cu stable isotopes measured by multi-collector inductively coupled plasma mass spectrometry (MC-ICPMS). The MC-ICPMS technique gives access to natural variations of the ⁶⁵Cu/⁶³Cu ratio including biological samples such as cells, animal fluids and organs.³⁸⁻⁴⁰ Hence, we designed the following experiment. We reasoned that if the hepatic copper content could be enriched in ⁶⁵Cu by injecting a ⁶⁵Cu solution to the *Atp7b*-/- mice, we could collect their faeces during the following days and find out whether any ⁶⁵Cu-enrichment could be measured. Assuming that Chel2 can restore Cu excretion through the bile, *i.e.* the physiological pathway, then treating the mice by injecting Chel2 after the ⁶⁵Cu injection should induce an increase in the ⁶⁵Cu/⁶³Cu ratio in the faeces.

We first evaluated how much excess 65 Cu could be found in the liver from $Atp7b^{-/-}$ mice during the days following an intravenous (IV) injection of 65 Cu. Preliminary experiments on $5 Atp7b^{-/-}$ mice showed that after injecting ≈ 700 ng 65 Cu, their liver contained ≈ 500 ng of excess 65 Cu one or two days later. The 24h-faeces collected in the meantime also contained excess 65 Cu (11-22 ng/g faeces) accounting for less than 5% of that found in the liver (487-660 ng/g liver). Thanks to these results, we could envisaged to treat the mice with Chel2 for a week-long experiment.

Eight adult $Atp7b^{-/-}$ mice received an IV injection of 1-1.5 µg ⁶⁵Cu at day -4 and from day 0 on, 8 IV injections of Chel2 (30 mg/kg, once a day for 4 days in a row, repeated the following week), as described in the upper panel of Fig. 6. The experiment was stopped on day 11. Another eight adult $Atp7b^{-/-}$ mice received Chel2 by subcutaneous (SC) injection instead of the IV route. The latter series

lasted only one week and was stopped at day 4. Two other groups of 4 *Atp7b*-/-mice were treated for control experiments that were stopped at day 4. They received either one dose of ⁶⁵Cu at day -4 and 4 injections of buffer instead of Chel2, or one injection of vehicle at day -4 and 4 injections of Chel2. The faeces of all mice were collected every other day during treatment and at euthanasia, their liver was removed for analysis.

The excess of 65 Cu, denoted 65 Cu_{xs} from now on, reflects in all samples the effect of the 65 Cu spike on the isotopic Cu distribution. It is calculated by the difference between 65 Cu_{spl}, the 65 Cu amount measured in the sample and the natural 65 Cu amount in the sample. The latter is calculated from 63 Cu_{spl}, the 63 Cu amount measured in the sample and r_{ref} , the natural 65 Cu/ 63 Cu ratio measured in the isotopic standard NIST SRM 976 solution that always brackets the samples during the analysis.

Denoting $r_{ref} = {}^{65}\text{Cu}_{ref}$ / ${}^{63}\text{Cu}_{ref}$ and $r_{spl} = {}^{65}\text{Cu}_{spl}$ / ${}^{63}\text{Cu}_{spl}$, the isotopic ratio measured in the reference solution and the sample, respectively and designating by $\text{Cu}_t = {}^{65}\text{Cu}_{spl} + {}^{63}\text{Cu}_{spl}$, the total Cu measured in the sample, ${}^{65}\text{Cu}_{xs}$ is calculated as follows:

65
Cu_{xs} = (r_{spl} - r_{ref}) × 63 Cu_{spl},
with 63 Cu_{spl} × (r_{spl} + 1) = Cu_t,
and finally 65 Cu_{xs} = (r_{spl} - r_{ref}) / (r_{spl} + 1) × Cu_t

The results are then expressed per unit of wet weight, ranging from 5 to 117 ng/g for the faeces and from 0.1 to 1.4 μ g/g for the liver. In the control experiment where the mice did not receive 65 Cu, the faeces contained 1.6 ng/g excess 65 Cu (mean from 12 samples) and the 4 livers, -0.012 μ g/g, showing that r_{spl} was not changed by the Chel2 treatment. In the other experiments where 65 Cu was injected to the mice, dividing 65 Cu_{xs} obtained in the faeces by 65 Cu_{xs} obtained in the liver allows to evaluate 65 Cu excreted in the faeces, independently of individual variability. All the data is shown in Fig. 6 in permil.

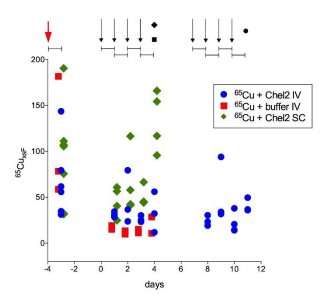


Fig. 6 Effect of Chel2 injections (starting at day 0, black arrows on top) on the fraction of hepatic ⁶⁵Cu excreted in the faeces of mice (⁶⁵Cu_{xsF} in permil) following an IV injection of ⁶⁵Cu at day -4 (red arrow). (■) 4 mice received 4 IV injections of buffer from day 0 to day 4; (●) 7 mice received 8 IV injections of Chel2 (30 mg/kg) from day 0 to day 10; (◆) 8 mice received 4 SC injections of Chel2 (50 mg/kg) from day 0 to day 4. On top, segments figure the 24-h faeces collections and black symbols, euthanasia.

Before any treatment, *i.e.* during the 24h-period following the 65 Cu injection, excess of 65 Cu was evidenced in the faeces (see 65 Cu_{xsF} at day -3 in Fig. 6). In the control group, after the first buffer injection and thereafter, there was almost no more excess of 65 Cu in each 24h-faeces sample and 65 Cu_{xsF} value was around 15‰. Bearing in mind that these mice do not excrete Cu from their liver, the measured 65 Cu_{xs} value in the faeces at day -3 suggests the existence of an excretion pathway for Cu from blood to faeces, crossing the intestinal epithelium. The existence of such a Cu-excretion pathway has not been documented up to now. As a secondary excretion pathway appearing in the $Atp7b^{-/-}$ mice, it could also reflect the role of non absorptive cells in the intestine.

In the groups that received Chel2, either by IV or by SC injections, the fraction of hepatic ⁶⁵Cu excreted in the faeces was significantly higher than in the control group. The ⁶⁵Cu_{xsF} values for each group are pooled per day in Fig. S3 (mean and SEM). The IV-treated mice excretion was stable for the two weeks of treatment with a value around 33%, twice as much as the control mice. In the Chel2 SC-treated animals, the excretion reached 45% at day 1 and tended to increase during the week of treatment. The data collected from the different mice during the 4 days were pooled and submitted to statistical analysis, which demonstrates that both 4-day treatments were more effective than the control treatment, with a larger efficacy of the SC treatment with respect to the IV one: Chel2 IV-injections 33±4% (n=14) or Chel2 SC-injections 73±11% (n=16) vs buffer 15±2% (n=8) (p<0.001, t-test). When Chel2 was given through IV injections for a second week, the effect on ⁶⁵Cu_{xsF} was the same as during the first week, suggesting that it was limited by the amount of Chel2 reaching the liver and the bile. As a matter of fact, Chel2 is a highly hydrophilic compound, therefore expected to be rapidly cleared from the serum. Altogether, the IV injections were probably limiting the amount of Chel2 reaching the liver and crossing the hepatocytes. In contrast with the IV treatment, the SC injections were more effective in excreting excess ⁶⁵Cu from the liver, suggesting a more progressive delivery of Chel2 to the liver.

Discussion.

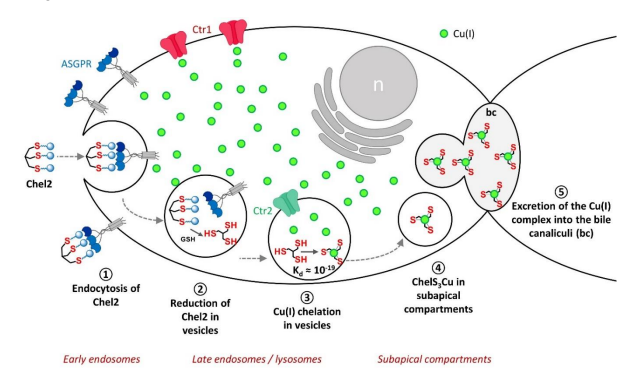
The mechanism of action of Chel2, the liver-targeting Cu(I) chelator, was studied in WIF-B9 cells that reconstitute polarized epithelia with functional bile canaliculi. The relationship between Chel2 and Cu localization and distribution in these cells could be inferred from fluorescence confocal microscopy using the fluorescent derivative Chel2* and nano XRF, respectively.

For further discussion, we need to bear in mind that upon endocytosis in WIF-B9 cells, Chel2 releases Chel(SH)₃, the high affinity Cu(I) chelator. This was shown before, following the effect of Chel2 addition to the culture medium in the presence of 1 μ M Cu on the Wilson protein localization and it provided the conditions used here for chelation (50 μ M Chel2 + 1 μ M Cu).¹⁷ We can therefore expect the same for Chel2* (1 μ M Chel2* + 1 μ M Cu) which will also release the high affinity chelator. Therefore, we assign to Chel*(SH)₃ and Chel*S₃Cu(I) the red spots in Fig. 2 and 3.

Asialoglycoprotein endocytosis in polarized WIF-B9 cells has been previously studied and a remarkable feature is that the proteins pass through a subapical compartment that sorts them: some are sent to the lysosomes for degradation and others are sent to the apical membrane for transcytosis. Using the same route with Chel2* instead of a protein, however, shows that there is colocalization with lysosomes and that red vesicles also concentrate around the canaliculi (Fig. 2). This suggests that after reduction of Chel2*, some Chel*(SH)₃ and Chel*S₃Cu are trapped in non-lysosomal vesicles on their way to exocytosis. Indeed, red fluorescence is seen in the middle of the canaliculus after 3 h (Fig. 3). Interestingly enough, these subapical vesicles could also be the hot spots of Cu seen by XRF, which would be full of ChelS₃Cu, under the chelation conditions (50 µM Chel2, Fig. 4C).

Going back to the effect of 1 μM Cu for 2 h, we recall that under these conditions, ATP7B is found in the canalicular membrane, where it pumps Cu out of the cell, into the bile canaliculus. 17 According to Nyasae et al.⁴³ whose conclusions were derived from a similar polarized epithelium, we can assume ATP7B to have left the trans-Golgi network in vesicles and trafficked to the canalicular membrane via at first, basolateral endosomes and then, the subapical compartment. The same observation in Can-10 hepatic cells, 44 that are also able to reconstitute a polarized epithelium, suggests that in polarized hepatic cells, proteins undergoing transcytosis and proteins meant to reach the apical membrane use the same route and there is no such a secretory pathway from the Golgi to the apical membrane. Therefore, we can assume that while being transported towards the canalicular membrane, ATP7B has pumped Cu inside the vesicles. Hence, the hot spots of Cu that are seen in the cytoplasm of the cells (Fig. 4B). In the presence of Chel2, endocytosis vesicles, that have a Cu transporter in their membrane,⁴⁵ are also able to accumulate Cu because Chel(SH)₃ binds Cu(I) inside the vesicles, hence creating a gradient for the Cu transporter. This will last until the basal "free" Cu concentration is reached, when ATP7B will return to the trans-Golgi network. At that time, the canalicular region will be full of subapical vesicles entrapping ChelS₃Cu and ready for exocytosis. All these observations led us to propose the mechanism shown in scheme 2 for Chel2 fate in hepatic cells and its relationship with intracellular copper chelation. It can be described as follows: (1) Chel2 binds to ASGPR and enters the hepatocytes by basolateral or sinusoidal endocytosis, (2) Chel(SH)₃, the high affinity chelator, is released from Chel2 by reduction with GSH in these vesicles, (3) ChelS₃Cu, the Cu(I) complex, is formed in sinusoidal endocytic vesicles, (4) some vesicles fuse with lysosomes, while others become the subapical compartment and (5) the latter spills its load into the canalicular lumen leading to Cu excretion into the bile canaliculi.

If this were the mechanism occurring in *ATP7b*^{-/-} hepatocytes, then Chel2 would restore the pathway for Cu excretion out of the liver. Our study with *ATP7b*^{-/-} mice following excess ⁶⁵Cu by MC-ICPMS supports this mechanism. Mice that received ⁶⁵Cu followed by Chel2 treatment showed a significant increase of the ⁶⁵Cu content in their faeces. This demonstrates increased Cu excretion after IV injections and even more efficiently after SC injections of Chel2 and most importantly excretion through the bile.



Scheme 2 Proposed mechanism for Chel2 fate in hepatic cells in the presence of Cu.

Conclusions.

In search of the mechanism of action of Chel2, the liver-targeting Cu(I) chelator, we observed in hepatocytes that among the Chel2 molecules that entered by endocytosis, some were localized in the lysosomes, whereas others were found in vesicles close to the canaliculus membrane, most probably in subapical compartments, and finally in the canaliculus lumen. Interestingly enough, hot spots of Cu were found surrounding the canaliculi in WIF-B9 cells, suggesting that the chelator-containing vesicles were full of Cu. Because in polarized WIF-B9 cells, subapical compartments are known to fuse with the apical membrane, we assumed that the chelator released by Chel2 was expelled in the canaliculus lumen together with Cu. Therefore, intracellular Cu chelation by the high affinity chelator released by Chel2 is expected to occur in vesicles along Chel2 transcytosis pathway, leading ultimately to Cu excretion into the bile canaliculi. Importantly, Chel2 injection either intravenously or subcutaneously to a murine model of Wilson's disease increased excretion of Cu in the faeces, confirming the biliary excretion mechanism at the animal level. As things stand now, Chel2 turns to a possible means to collect and excrete hepatic Cu in the faeces, hence restoring the physiological pathway. It would be interesting to check this on bigger animals whose condition can be monitored daily, to envisage the use of Chel2 for patients who do not tolerate the current treatments of Wilson's disease.

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