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Development of Cu(II)-specific peptide shuttles capable of preventing Cu–amyloid beta toxicity and importing bioavailable Cu into cells†

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Copper (Cu) in its ionic forms is an essential element for mammals and its homeostasis is tightly controlled. Accordingly, Cu-dyshomeostasis can be lethal as is the case in the well-established genetic Wilson's and Menkes diseases. In Alzheimer's disease (AD), Cu-accumulation occurs in amyloid plaques, where it is bound to the amyloid-beta peptide (Aβ). *In vitro*, Cu–Aβ is competent to catalyze the production of reactive oxygen species (ROS) in the presence of ascorbate under aerobic conditions, and hence Cu–Aβ is believed to contribute to the oxidative stress in AD. Several molecules that can recover extracellular Cu from Aβ and transport it back into cells with beneficial effects in cell culture and transgenic AD models were identified. However, all the Cu-shuttles currently available are not satisfactory due to various potential limitations including ion selectivity and toxicity. Hence, we designed a novel peptide-based Cu shuttle with the following properties: (i) it contains a Cu(II)-binding motif that is very selective to Cu(II) over all other essential metal ions; (ii) it is tagged with a fluorophore sensitive to Cu(II)-binding and release; (iii) it is made of a peptide platform, which is very versatile to add new functions. The work presented here reports on the characterization of AKH-αR5W4^{NBD}, which is able to transport Cu ions selectively into PC12 cells and the imported Cu appeared bioavailable, likely *via* reductive release induced by glutathione. Moreover, AKH-αR5W4^{NBD} was able to withdraw Cu from the Aβ_{1–16} peptide and consequently inhibited the Cu–Aβ based reactive oxygen species production and related cell toxicity. Hence, AKH-αR5W4^{NBD} could be a valuable new tool for Cu-transport into cells and suitable for mechanistic studies in cell culture, with potential applications in restoring Cu-homeostasis in Cu-related diseases such as AD.

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Introduction

Copper (Cu), being an essential transition metal in biology, is very important for an array of biological functions in the body and to a large extent in the brain. Cu is present as ions (Cu(I) and Cu(II)) and is absorbed by cells of the choroid plexus at the blood–brain barrier into the cerebrospinal fluid (CSF),¹ from where it is transported into brain cells by the transmembrane

Cu-transporter, CTR1, in its reduced Cu(I) form.² The entering Cu(I) is transferred to Cu-chaperons like CCS (copper chaperone for superoxide dismutase), responsible for the delivery of Cu(I) to superoxide dismutase (SOD), an antioxidant enzyme.^{3,4} Another Cu-chaperon, COX17, transports Cu(I) to cytochrome C oxidase in the mitochondria,⁵ whereas ATOX1 (ref. 6 and 7) delivers Cu(I) to the Trans Golgi Network (TGN), where Cu is incorporated into cupro-proteins mainly destined to the extracellular space. These cupro-proteins play important roles in maintaining physiological activities. For example, the cupro-protein ceruloplasmin is primordial for Fe homeostasis and transport, while another cupro-protein, dopamine-β-hydroxylase, is needed for the conversion of dopamine to norepinephrine.⁷ Despite the importance of Cu in biology, excess Cu can provoke an array of dysfunctions, and therefore systems are in place in the advent of excess intracellular Cu. The Cu-transporting P-type ATPase, ATP7A, which is normally located to the Trans-Golgi Network (TGN), relocates to vesicles in proximity to the plasma membrane during excessive intracellular Cu levels and pumps Cu into these vesicles which are then

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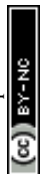
excreted from the cell.^{8,9} This pathway of Cu excretion is also used to regulate synaptic signals, as Cu has been shown to play a role in regulating NMDA receptor activity by binding to the post-synaptic NMDA receptors.¹⁰ Cu has also been shown to bind to other cell surface proteins such as PrP, AMPA, and GABA receptors amongst others.¹

Given the crucial role of Cu in biological systems, the deregulation of Cu homeostasis by mutations in the ATP7 transporters provoke Menkes and Wilson's diseases, both leading to neurodegeneration in patients.^{11–13} Moreover, Cu dyshomeostasis has also been linked to other neurodegenerative diseases such as amyotrophic lateral sclerosis (ALS), Parkinson's, Huntington's and Alzheimer's diseases (AD).¹³ The role of Cu dyshomeostasis in AD is one of the most studied in the literature with the main hallmark for AD being the appearance of neurofibrillary tangles (NFTs) and amyloid plaques in brain regions affected.¹⁴ Recent models actually suggest that these alterations may be cumulative and have demonstrated a connection between the amyloid hypothesis (amyloid plaques) and tau pathology (NFTs).^{14–16} Additionally, there has also been an increasing number of studies on other causative factors including mitochondrial dysfunction and increased reactive oxygen species (ROS) levels.^{17–20} Equally, emerging studies have also demonstrated the important role of amyloid species in the production of ROS.²¹ These studies show that amyloid-beta (A β) the main component of amyloid plaques is capable of binding metal ions at a relatively high affinity. Interestingly, high concentrations of Fe(III) \approx 1 mM, Zn(II) \approx 1 mM and Cu(II) \approx 0.4 mM have been found in amyloid plaques, with A β having the highest affinity for Cu(II) with a conditional affinity value at pH 7.4 of 10^{10} M^{-1} .²² In the reducing intracellular environment of cells, Cu is mainly found in its reduced form, whereas in the less reductive extracellular environment, Cu is found mainly in its oxidized form. However, Cu is never free, as it is bound to chaperones or cupro-proteins or other molecular ligands such as A β , where Cu can switch between its oxidized and reduced forms as a result of reductants such as ascorbate found in abundance in living organisms.^{23,24} These reductants reduce Cu(II) to Cu(I), which can be oxidized back to Cu(II) by oxidizing agents. The reaction of Cu(II) with dioxygen produce ROS, which can cause damage to proteins, lipids (including cell membranes), and DNA.^{21,25} Under physiological conditions, brain cells are protected from overproduction and accumulation of ROS by the production of antioxidants such as glutathione and SOD and by the cleavage of A $\beta_{1-40/42}$ to A $\beta_{4-40/42}$.^{26,27} In the A $\beta_{4-40/42}$ form, A β possesses an Amino Terminal Cu(II)- and Ni(II)-Binding (ATCUN) motif, which is composed of three amino acids with histidine (His) at position 3 (Xxx-Zzz-His). ATCUN motifs are highly selective for Cu(II) and possess a high affinity ranging from 10^{12} to 10^{15} M^{-1} .²⁸ The first two positions in ATCUN motifs can be composed of any amino acid (but Pro in position 2) with the motif in A $\beta_{4-40/42}$ being Phe-Arg-His.^{27,29} They have been shown to bind Cu(II) in a coordination unsuitable for Cu(I), thereby preventing its transition from the oxidized to the reduced form of Cu with a physiological reductant.³⁰ These ATCUN motifs are found in numerous endogenous proteins, with the most

common being Asp-Ala-His (DAH) found in Human Serum Albumin (HSA).³¹

Cu binding to A β has been reported to increase toxicity to cells, with proposed mechanisms such as Cu-catalyzed ROS production and/or its ability to change the A β conformation and hence the promotion of more toxic aggregates of A β . Both mechanisms could also occur together in the case of ROS induced tyrosine bridges between two A β , which reduces clearance efficiency by degrading the A β oligomers.³² Additionally, the binding of Cu to amyloid plaques increases the dyshomeostasis between intracellular and extracellular Cu. There have been studies and clinical trials with the main objective to correct this Cu dyshomeostasis found in AD, which led to studies on Clioquinol, PBT2 and GSTM.^{33–35} These molecules, termed ionophores or shuttles, are synthetic chemical molecules capable of binding Cu extracellularly such as retrieving Cu from extracellular plaques, entering the cell and releasing the Cu intracellularly.^{33,34} These compounds are not specific to Cu(II) as they have been shown to bind to other metals such as Zn. However, in AD there is also accumulation of Zn, Fe as well as Cu. Hence from a therapeutic point of view it is not clear which is the best strategy, either targeting all metal ions at the same time and place or to use selective ionophores targeting one defined metal ion. The latter would allow the targeting of each type of metal ion in a time and may be space specific manner. However, as a tool to better understand the role of Cu in AD, a selective shuttle would be very useful. Another potential problem of ionophores such as PBT-2 is linked to their toxicity and recently the term cuproptosis was coined. Elesclomol, a Cu ionophore, has been shown to induce the so-called cuproptosis at low nanomolar concentrations, which was attributed the Cu induced aggregation of lipoylated mitochondrial proteins in the TCA cycle.³⁶ This may partially explain why Cu ionophores in clinical trials, such as Clioquinol, were unsuccessful.³³ On the other hand, the same compound Cu-elesclomol could alleviate the Menkes pathology, a genetic disease of Cu-deficiency.³⁷ This indicates that the amount, kinetics, and route of Cu-supply to cells are very critical for the restoration of Cu-homeostasis. Hence, new shuttles that are specific to Cu(II) and presenting less toxicity are attractive alternatives.

To this end, we have developed Cu(II) selective shuttles capable of importing Cu into cells thanks to a cell penetrating peptide (CPP) domain. CPPs are composed of amino acids forming usually an aliphatic or cationic chain.^{38,39} These peptides are capable of forming secondary structures when in contact with the cell membrane and driving cellular entry usually through direct penetration and/or endocytosis.^{38,40} Among the most studied CPPs are TAT,^{41–43} penetratin,^{44–46} and pVEC.^{47–49} For this study, we identified CPPs capable of penetrating PC12 cells used as a model of neuronal cells. These CPPs were coupled with the high Cu(II) affinity ATCUN motif AKH and a NBD (4-chloro-7-nitrobenzofurazan) molecule to permit tracking of the molecule in cells. The novel Cu(II) specific shuttles were tested for their ability to retrieve Cu(II) from A β_{1-16} , to transport Cu into PC12 cells and release the Cu intracellularly.



Results and discussion

Characterization of a Cu(II)-binding module grafted to CPP (AKH-CPP^{NBD})

Design of peptide Cu-shuttles. To obtain a peptide Cu-transporter for cell uptake, four candidate CPPs were tested for cellular uptake efficiency. Hence, the known CPPs: SG3,⁵⁰ and α R5W5,³⁹ and the antimicrobial peptide LAH4,⁵¹ along with DSIP (delta sleep inducing peptide), were synthesized by solid-phase peptide synthesis and tagged on the orthogonal sidechain of a lysine with a fluorophore (NBD, λ_{ex} 478 nm, and λ_{em} 545 nm). PC12 cells were exposed to these four peptides and their uptake was qualitatively evaluated by confocal microscopy after 1 h incubation (data not shown). The two CPP sequences with the best cell penetration ability, α R5W4 and LAH4, were selected for further studies. The α R5W4 and LAH4 CPP sequences were elongated with the well-known ATCUN motif (Xxx-Zzz-His).⁵² For the Xxx and Zzz amino acids in the motif, Ala and Lys were used. Thus, two peptides, AKH-LAH4^{NBD} and AKH- α R5W4^{NBD} containing the Cu(II)-binding motif AKH and a CPP, were synthesized and were subsequently tagged with the fluorophore NBD on a Lys sidechain at position 2 or 4, respectively (see the Materials and methods). As a control, a peptide without any CPP sequence was synthesized, AKHK^{NBD}, consisting of the Cu(II)-binding site with an NBD fluorophore on the Lys in position 4. An additional peptide, DAHK, comprising the ATCUN motif found on human serum albumin (HSA) was optionally used as an additional control (Table 1). These last two peptides were expected to bind Cu(II) as the parent CPP-containing peptides (AKH-LAH4^{NBD} and AKH- α R5W4^{NBD}), but not to penetrate the cell membrane.

Cu(II) quenching of NBD fluorescence. We next tested the ability of the fluorophore (NBD) linked to these peptides to sense Cu(II) binding. As expected in the absence of Cu(II), we observed the characteristic NBD absorption spectrum with a maximum between 460 nm and 490 nm and the appearance of a second peak at 350 nm.⁵³ Indeed, upon excitation at 478 nm, emission with a maximum at 545 nm was observed for AKHK^{NBD}, AKH- α R5W4^{NBD} and AKH-LAH4^{NBD} (Fig. 1).

Upon the addition of Cu(II), all three peptides showed a steady decrease in fluorescence emission. AKHK^{NBD} and AKH- α R5W4^{NBD} had a clear break point at 1 equivalent with an ensuing plateau. The total quenching was about 90% in HEPES solution. This is in line with the strong affinity of the Xxx-Zzz-His motif for Cu(II) and the well-known quenching effect of Cu(II).⁵⁴ For AKH-LAH4^{NBD}, a similar behavior was observed, but the break point was less marked (absence of a sharp break point). This could in principle be due to a lower affinity, but further experiments suggest rather a time-dependent

fluorescence change of NBD emission due to peptide aggregation. Indeed, AKH-LAH4^{NBD} fluorescence evolved with time at pH 7.4, but not in the presence of 6 M GndHCl, pH 7.4 (or at pH 2) (Fig. S3†). This observation shows the importance of prudence in the interpretation of fluorescence data relating to AKH-LAH4^{NBD} (Fig. S3†). For further tests, only data with AKH- α R5W4^{NBD} will be shown, but the results of AKH-LAH4^{NBD} are available in the ESI (Fig. S1–S11†).

Specific quenching of AKH- α R5W4^{NBD} by Cu(II). Given the efficient quenching of AKH-CPP^{NBD} by the binding of Cu(II) to the AKH ATCUN motif, we evaluated the selectivity of this quenching response on towards other essential biological d-block metals. In particular, we tested the potentially most competitive biologically essential ion Zn(II) as well as Fe(III) and Mn(II) (Fig. 2a). None of Zn(II), Fe(III) and Mn(II) induced a significant quenching in the fluorescence of AKH- α R5W4^{NBD} in 100 mM HEPES at pH 7.4. This showed that only Cu(II) was able to quench NBD fluorescence in AKH- α R5W4^{NBD}. Importantly, even in the presence of Zn(II), Fe(III) and Mn(II), Cu(II) was still able to quench the fluorescence to a similar degree to that in the absence of these metal ions. This supports the selectivity of Ala-Lys-His in AKHK^{NBD}, AKH- α R5W4^{NBD} and AKH-LAH4^{NBD} for Cu(II)-binding and is in line with the general feature of the Xxx-Zzz-His motif.⁵²

Release of Cu(II) from the AKH-CPP^{NBD} peptide with glutathione. The design of an effective Cu importer also requires the intracellular release of Cu(II). Indeed, the Xxx-Zzz-His motif is a strong Cu(II)-binder but has much weaker affinity for Cu(I). Thiols such as glutathione can reduce Cu(II) in this motif and then sequester Cu(I).³⁰ Hence, GSH present in high concentration in the cell could potentially release the Cu from AKH-CPP^{NBD}. To evaluate this aspect, AKHK^{NBD} and AKH- α R5W4^{NBD} pre-complexed to Cu(II) were exposed to 5 mM GSH at 37 °C. The recovery of the NBD fluorescence was monitored, in line with the reduction and withdrawal of Cu(II) from the peptides.

As expected, there was a recovery in fluorescence emission levels after incubation with GSH. Unexpectedly the release of Cu from AKH- α R5W4^{NBD} was 3 times faster than that from AKHK^{NBD}, indicating that the additional amino acids in AKH- α R5W4^{NBD} favor the reduction of Cu(II) and its removal by GSH. These data support a model in which most of the Cu(II) can be released from our shuttle intracellularly when triggered by physiological millimolar GSH concentrations.

AKH-CPP^{NBD} peptide as a bioavailable Cu importer

After showing *in vitro* that the two AKH-CPP^{NBD} peptides can bind Cu(II) and that 5 mM GSH can reduce Cu(II) and that this

Table 1 List of peptides selected comprising an ATCUN motif and the NBD fluorophore

Peptide	ATCUN motif	Fluorophore (AA position)	Linker + CPP sequence	C-ter
AKH- α R5W4 ^{NBD}	AKH-	NBD (4)	-KGRRWRRWR	Amide
AKH-LAH4 ^{NBD}	AKH-	NBD (2)	-KKALLALALHHLAHLALALALKKA	Amide
AKHK ^{NBD}	AKH-	NBD (4)	-K	Amide
DAHK	DAH-	NBD (4)	-K	Amide



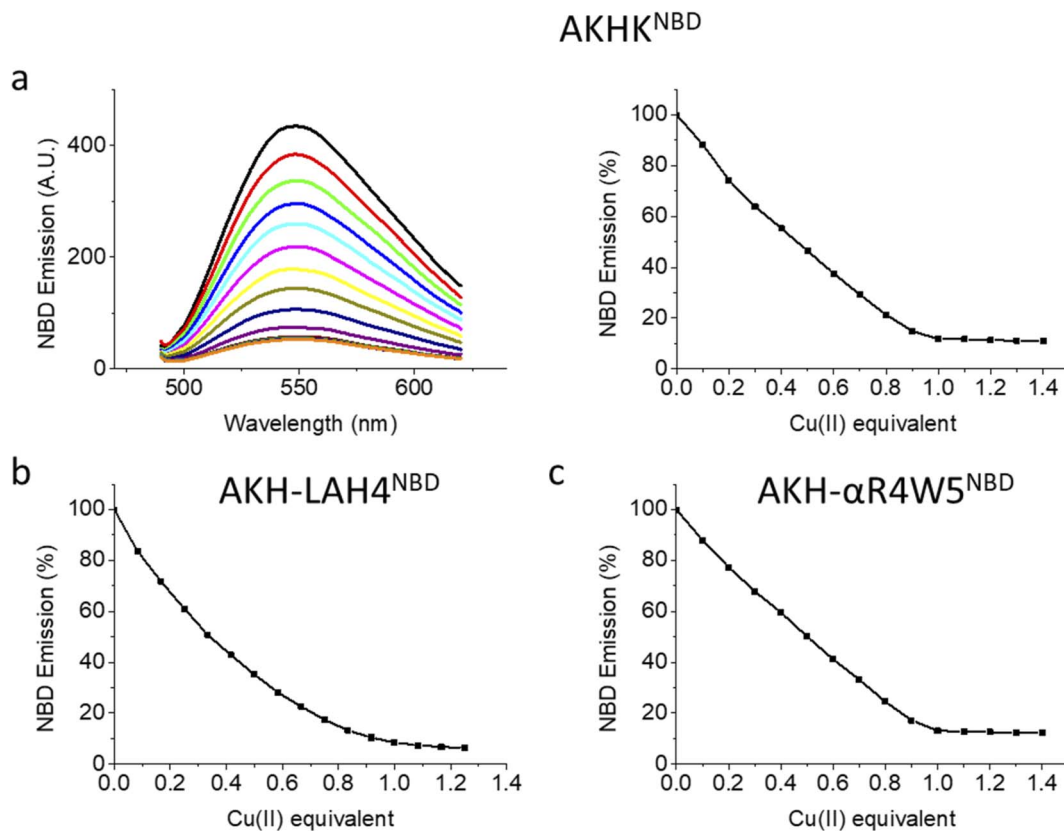


Fig. 1 Fluorescence spectra and Cu(II) dependent quenching: (a) AKH- α R5W4^{NBD}: left: spectra taken after each addition of 0.1 equivalents of Cu(II), and titration from black to brown; right: fluorescence intensity at 545 nm vs. added Cu(II); (b) AKH-LAH4^{NBD} and (c) AKHK^{NBD}. Conditions: 4 μ M peptides, addition of Cu(II); 0.1 equivalent = 0.4 μ M; HEPES buffer: 100 mM, pH 7.4. Excitation wavelength: 478 nm; emission wavelength: 545 nm.

release can be followed by fluorescence, we aimed to measure their Cu-importer activity on cultured cells. We used three different approaches for this evaluation, including intracellular Cu-concentration measurements *via* ICP-MS, confocal microscopy to follow expected fluorescence increase upon Cu(II)-

release from AKH-CPP^{NBD} in the cell and the translocation of the Cu(I) extruder, ATP7A, used as a sensor for bioavailable intracellular Cu.

Cu-concentrations *via* ICP-MS. The Cu(II) transport efficacy of the AKH-CPP^{NBD} peptides was evaluated and compared to

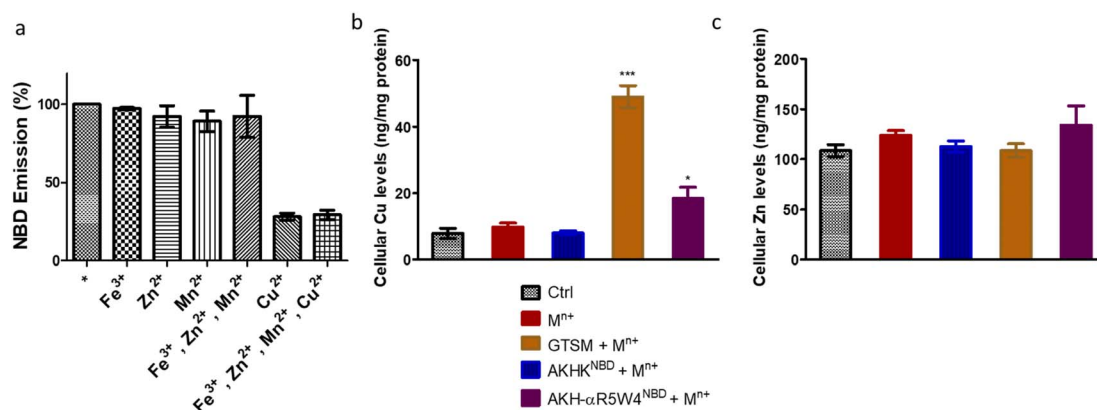


Fig. 2 Specific binding of AKH-CPP^{NBD} to Cu(II) in the presence of Fe(III), Zn(II) and Mn(II). (a) The emission spectrum of AKH- α R5W4^{NBD} was recorded before (*) and after the addition of Cu(II), Zn(II), Fe(III), Mn(II) or all of the above. Conditions: AKH- α R5W4^{NBD}: 7 μ M; Cu(II), Zn(II), Fe(III), and Mn(II): 5 μ M; HEPES 100 mM, pH 7.4. (b) Import of Cu ions or (c) Zn ions in PC12 by AKH- α R5W4^{NBD}. The peptides were pre-complexed with an equimolar concentration of Mⁿ⁺ (Cu(II), Zn(II), Fe(III), and Mn(II)) in 100 mM HEPES, and then cells were treated for 1 h at 37 °C in DMEM. Conditions: AKH- α R5W4^{NBD}: 5 μ M; Cu(II), Zn(II), Fe(III), and Mn(II). Each condition was done in duplicate; $n = 3$ independent experiments with 4×10^6 PC12 cells. A parametric ANOVA test was carried out with Dunnett's multiple comparison test, * $p < 0.01$ and *** $p < 0.0001$.



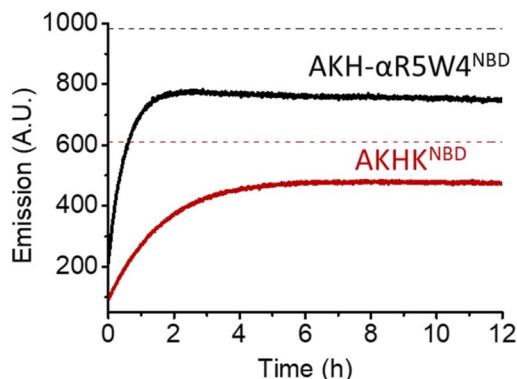


Fig. 3 Reduction of Cu(II) bound to AKH- α R5W4^{NBD} and AKHK^{NBD} by glutathione monitored by the increase in NBD fluorescence at 545 nm over time. Black and red dashes mark the expected emission endpoint (emission in the absence of Cu(II)) of AKH- α R5W4^{NBD} and AKHK^{NBD}, respectively. Conditions: 4 μ M Cu(II)-AKH-CPP^{NBD}, 5 mM GSH, HEPES 100 mM, pH 7.4 and 37 °C; representative traces of $n = 2$ independent experiments are shown.

that of the well-established glyoxal-bis N4-methylthiosemicarbazone (GTSM) and to that of the control peptides AKHK^{NBD} and DAHK^{NBD} (lacking the CPP moiety). As control experiments, we performed spiking experiments, where known amounts of Cu were added to samples just prior to lysis and ICP-AES/MS (see Fig. S5[†]). Indeed, the measured Cu levels increased in a proportional manner to the amount of Cu added. ICP-AES showed a 40% loss in Cu compared to the known Cu added amount, which we attribute to the matrix effect brought about by the cell pellet and the limit of the technique. However, a lower amount of Cu (25%) was lost while analyzing the samples with ICP-MS, indicating that ICP-MS is the best-suited approach to measure relative changes in the Cu concentrations of the cells under our conditions, though absolute values might be underestimated.

Basal Cu concentrations in PC12 cells are similar to what was obtained by Ogra *et al.*, 2016, *i.e.* about 10 ng mg⁻¹ protein.⁵⁵ For the positive control, PC12 cells were treated with 1 μ M GTSM, after treatment for 1 h, and we detected a 3–4-fold increase in intracellular Cu levels, lower than the increase previously reported in NGF-treated PC12 cells.⁵⁶ It is of note that NGF differentiated PC12 cells have an increased Cu(II) intake capacity, also verified by a higher basal Cu(II) content in these cells.⁵⁵

After confirming ICP-MS as a reliable tool for detecting cellular Cu uptake, 1 μ M Cu(II)GTSM, 5 μ M Cu(II)-AKH- α R5W4^{NBD} and 1 μ M Cu(II)-AKH-LAH4^{NBD} were tested. These concentrations did not induce significant toxicity on PC12 cells, as determined by a MTT assay (Fig. S5[†]). The addition of Cu(II)-AKH- α R5W4^{NBD} and Cu(II)-AKH-LAH4^{NBD} yielded an increase in Cu concentration in PC12 cells (Fig. 4 and S7[†]). 5 μ M Cu(II)-AKH- α R5W4^{NBD} was about as efficient as 1 μ M GTSM (3–4 times increase), whereas 1 μ M Cu(II)-AKH-LAH4^{NBD} was less efficient (2–3 times increase). From these experiments, the AKH- α R5W4^{NBD} peptide proved to be a reliable candidate for a Cu(II) shuttle. Subsequently we analyzed its capacity to specifically bind Cu(II) in the presence of other bioavailable transition

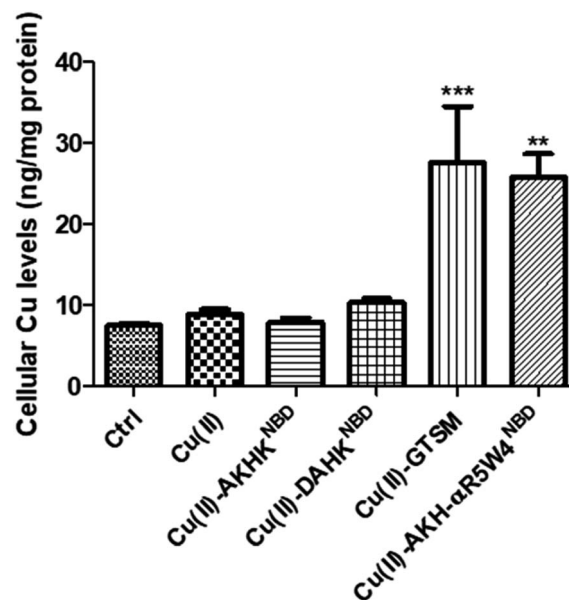


Fig. 4 AKH- α R5W4^{NBD} imports Cu into PC12 cells after 1 h incubation at 37 °C, measured by ICP-MS. Each condition was done in duplicate; $n = 3$ independent experiments with 2×10^6 PC12 cells. A parametric ANOVA test was carried out with a Dunnett's multiple comparison test, ** $p < 0.001$ and *** $p < 0.0001$.

metals Mⁿ⁺ (Zn(II), Fe(III) and Mn(II)). For this, we mixed the AKH- α R5W4^{NBD} peptide with Mⁿ⁺ and Cu(II) in HEPES before addition to PC12 cells in DMEM (Dulbecco's modified Eagle's medium) media for 1 h. ICP-MS data (Fig. 2b) showed that the AKH- α R5W4^{NBD} peptide is still capable of importing Cu(II) ions but did not import Zn(II) (Fig. 2c), the most competitive ion.

Fluorescence microscopy. The tagging of the peptides with NBD allowed the detection of the peptides *via* confocal microscopy and the selective fluorescence quenching by Cu(II) binding makes the fluorescence intensity sensitive to Cu(II) binding. Hence, intracellular fluorescence should reveal peptide uptake, whereas a subsequent increase in fluorescence intensity should indicate Cu(II) release or peptide accumulation. Hence, PC12 cells were treated with the control peptide Cu(II)-AKHK^{NBD} and the peptide shuttles, Cu(II)-AKH- α R5W4^{NBD} and Cu(II)-AKH-LAH4^{NBD} (Fig. 5 and S8[†]).

A clear entry of Cu(II)-AKH- α R5W4^{NBD} (Fig. 5) and Cu(II)-AKH-LAH4^{NBD} (Fig. S8[†]) into PC12 cells was seen as early as t0 (0 min) after the cells were incubated with the peptide complexes. Both Cu(II)-AKH- α R5W4^{NBD} and Cu(II)-AKH-LAH4^{NBD} seem to accumulate in vesicular compartments with their fluorescence increasing with time, which could suggest the reduction and release of Cu in these compartments (Fig. 3). For the case of Cu(II)-AKHK^{NBD} (absence of CPP), no detectable cell penetration is detected within an incubation period of 30 minutes. In conclusion, AKH- α R5W4^{NBD} and AKH-LAH4^{NBD} proved to be good Cu(II) shuttles as observed using fluorescence microscopy as they seem to import Cu and possibly release Cu in PC12 cells.

Translocation of ATP7A. Using both fluorescence microscopy and ICP-MS we have shown that Cu(II)-AKH- α R5W4^{NBD} and Cu(II)-AKH-LAH4^{NBD} can penetrate PC12 cells and import Cu into



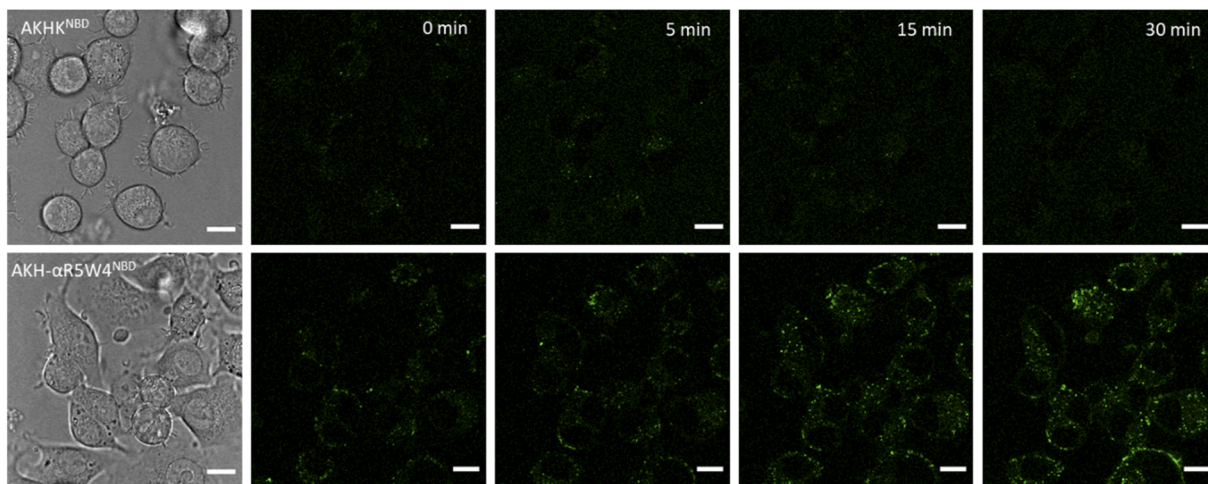


Fig. 5 Penetration of Cu(II)-AKH- α R5W4^{NBD} into PC12 cells. Representative images at the indicated time obtained from live imaging performed on 5×10^4 PC12 cells incubated with Cu(II)-AKHK^{NBD} or Cu(II)-AKH- α R5W4^{NBD} = 5 μ M, at 37 °C using a Leica TCS SP5 II confocal microscope, with an excitation wavelength at 478 nm for 30 min. Time point zero is about 1 minute after incubation with the peptides. Bars = 10 μ m. Similar observations were obtained with 5 different cell cultures.

PC12 cells; however it was unknown if the imported Cu becomes bioavailable (available for cellular processes). To indirectly monitor the bioavailability of the imported Cu, we used the Cu-sensitive delocalization of the Cu(I) exporter, ATP7A.

Under basal conditions, a good colocalization is observed in the PC12 cell between ATP7A and giantin, a marker of the TGN. However, upon increase in Cu(I) levels, ATP7A is known to relocate from the TGN to vesicles located in proximity to the plasma membrane.^{57,58} Accordingly, upon treatment with Cu(II)-AKH- α R5W4^{NBD} and Cu(II)GTSM, the staining of ATP7A delocalized from the TGN to vesicles in proximity to the plasma membrane (Fig. 6), suggesting an increase in Cu levels in the cells. As negative controls, Cu(II)-AKHK^{NBD}, Cu(II)EDTA and free Cu(II) were used. Given the negative charge of EDTA, it is unexpected to penetrate the cell membrane and accordingly, no delocalization of ATP7A was observed under these conditions after treatment. The peptide, AKHK^{NBD}, not having a CPP did not provoke a significant delocalization of ATP7A, neither did free Cu(II) which could enter through the Cu transporter, CTR1.

In summary, all three methods support the notion that Cu(II)-AKH- α R5W4^{NBD} is able to import Cu into cells, which becomes bioavailable as revealed by the ATP7A delocalization from the TGN. Bioavailable intracellular Cu is likely to be Cu(I), in line with the reduction of Cu(II) peptide complexes by intracellular reductants leading to the subsequent release of Cu(I).

Inhibition of ROS production *in vitro* and *in vivo* by the AKH-CPP^{NBD} peptide

Retrieval of Cu(II) from A β ₁₋₁₆. The A β peptide is the major component of amyloid plaques. These plaques accumulate Cu, which is bound to the metal-binding domain (N-terminal amino acids 1-16) of A β _{1-40/42}. Cu(I/II)A β _{1-40/42} and Cu(I/II)A β ₁₋₁₆ catalyze quite efficiently ROS production in the presence of dioxygen and the physiological reducing agent, ascorbate. Ascorbate is present extracellularly around neurons with a concentration

around 0.1 mM.^{59,60} Thus, the ability of AKH- α R5W4^{NBD} and AKHK^{NBD} to retrieve Cu(II) from pre-complexed A β ₁₋₁₆ was investigated in HEPES buffer. A solution of pre-complexed 10 μ M Cu(II)-A β ₁₋₁₆ 0.5 : 1 was incubated with 0.5 equivalents (with respect to A β ₁₋₁₆) of AKH- α R5W4^{NBD} or AKHK^{NBD} and the transfer of Cu(II) from A β ₁₋₁₆ to our Cu(II) shuttles was followed using the emission quenching of NBD when Cu(II) is coordinated to the ATCUN motif (AKH) in AKH- α R5W4^{NBD} or AKHK^{NBD}.

In 10% DMEM buffer, the addition of AKH- α R5W4^{NBD} and AKHK^{NBD} to Cu(II)-A β ₁₋₁₆ resulted in fluorescence quenching in line with their high Cu(II)-binding affinity. The intensity of the quenching after 90 min is similar to the quenching of their respective pre-formed complexes (dashed lines), indicating that most of the Cu(II) was transferred. Given the aggregative propensity of AKH-LAH4^{NBD} (Fig. S3†), the transfer of Cu from A β ₁₋₁₆ was not studied. However, it is expected to be similar to AKH- α R5W4^{NBD} and AKHK^{NBD}, since all peptides have the same ATCUN motifs.

Ascorbate oxidation and ROS production. We further investigated the ability of AKH- α R5W4^{NBD} and AKHK^{NBD} to prevent ROS production caused by the cycling of Cu-bound A β ₁₋₁₆ (Scheme 1).

In agreement with the literature, Cu(II)-A β ₁₋₁₆ induced a fast decrease in the absorption of ascorbate at 265 nm (Fig. 8, black) due to the oxidation of ascorbate and the concomitant ROS production.^{61,62} Then Cu(II)-A β ₁₋₁₆ was added at the same time without pre-incubation (co-addition) with AKH- α R5W4^{NBD} or AKHK^{NBD}. In this case, AKH-LAH4^{NBD} (Fig. S9†), AKH- α R5W4^{NBD} or AKHK^{NBD} (Fig. 8a and b) was able to decrease ascorbate oxidation, albeit not being immediate. At the beginning of the recordings, most of the Cu is still bound to A β ₁₋₁₆. Nevertheless, with time ascorbate oxidation slowed down and a plateau is reached, higher than for Cu(II)-A β ₁₋₁₆ alone, indicating that more and more Cu is transferred to AKH-LAH4^{NBD}, AKH- α R5W4^{NBD} or AKHK^{NBD}. This is indeed in accordance with



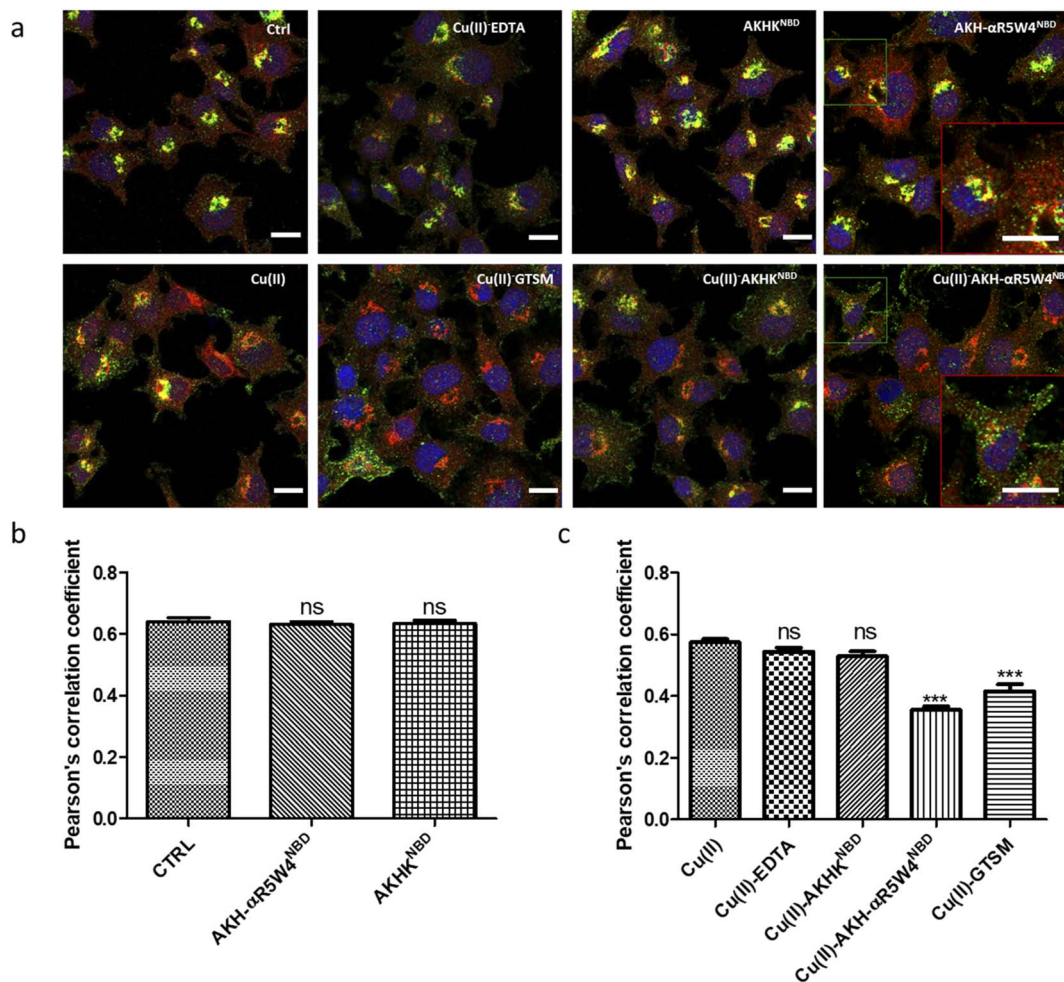
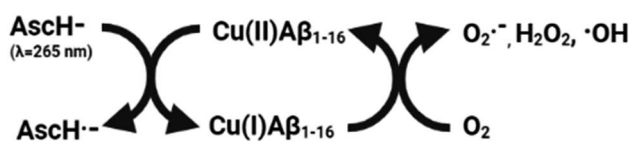


Fig. 6 Cu import by AKH- α R5W4^{NBD} induced delocalization of ATP7A from the TGN to vesicular structures. (a) PC12 cells were incubated for 1 h in DMEM media alone (control = Ctrl) or 5 μ M Cu(II), 5 μ M Cu(II)-EDTA, 5 μ M AKHK^{NBD}, 5 μ M Cu(II)-AKHK^{NBD}, 5 μ M AKH- α R5W4^{NBD}, 5 μ M Cu(II)-AKH- α R5W4^{NBD} or 1 μ M Cu(II)-GTSM, before fixation and processing for immunostaining; blue: Hoechst (nucleus marker); green: ATP7A; red: Giantin (Golgi marker). Representative images are shown. Bars = 10 μ m. $n = 3$ independent experiments. A zoomed inset is displayed for AKH- α R5W4^{NBD} and Cu(II)-AKH- α R5W4^{NBD} to illustrate the vesicular subplasmalemmal redistribution of ATP7A staining. (b and c) Quantification of the colocalization between ATP7A and Giantin staining measured as Pearson's correlation coefficient. Number of cells analyzed ≥ 27 for each condition. A parametric ANOVA test was carried out with Dunnett's multiple comparison test *** $p < 0.0001$.



Scheme 1 Cycling of Cu between its oxidized and reduced forms on $A\beta_{1-16}$ in the presence of ascorbate.

the time dependent transfer of Cu from $A\beta_{1-16}$ to AKH-LAH4^{NBD}, AKH- α R5W4^{NBD} or AKHK^{NBD} as seen in Fig. 7.

Next, we tested the reaction with a 1 h pre-incubation of AKH-LAH4^{NBD}, AKH- α R5W4^{NBD} or AKHK^{NBD} with Cu(II)- $A\beta_{1-16}$, to allow Cu(II) transfer from $A\beta_{1-16}$ to the ATCUN motif. The ascorbate consumption was markedly insignificant (Fig. 8a and b). This is in line with the quasi-inexistent redox-activity of Cu(II) bound in the Xxx-Zzz-His motif.³⁰ When Cu(II)- $A\beta_{1-16}$ is pre-

incubated with AKH-CPP^{NBD} for 1 h, there is complete transfer of Cu(II) from $A\beta_{1-16}$ to the AKH Cu(II)-binding site, and therefore there is no efficient redox cycling of Cu and thus no reduction in ascorbate absorption. Hence, once Cu(II) is bound to the AKH motif in AKH-LAH4^{NBD}, AKH- α R5W4^{NBD} or AKHK^{NBD}, it is almost completely redox silenced and incapable of catalyzing deleterious amounts of ROS.

Intriguingly, the higher ascorbate oxidation rate observed in the case of AKH- α R5W4^{NBD} was unexpected, as it has the same Cu(II)-binding motif as AKHK^{NBD}. This higher ROS production rate (Fig. S9d[†]) in the presence of AKH- α R5W4^{NBD} was also seen to the same extent in the presence of pre-complexed Cu(II)-AKH- α R5W4^{NBD} (Fig. 8c). Further experiments were able to attribute this higher ascorbate oxidation to Fe(III) contamination in the AKH- α R5W4^{NBD} sample, which is markedly reduced by the addition of a strong Fe(III) chelator, deferoxamine (DFO) (Fig. 8c). Such Fe(III) contamination in the batch could



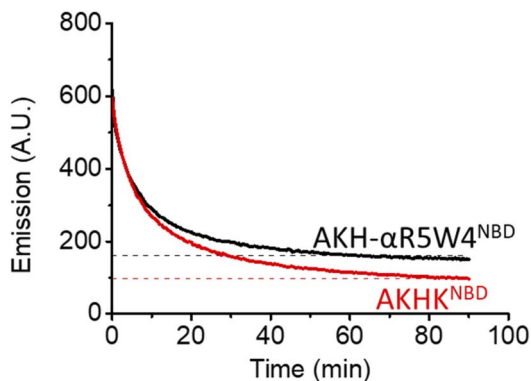


Fig. 7 Retrieval of Cu(II) from $A\beta_{1-16}$ by AKH- α R5W4^{NBD} and AKHK^{NBD}. Dashed lines indicate the expected emission endpoint of AKH- α R5W4^{NBD} (black) and AKHK^{NBD} (red), respectively, in the presence of Cu(II) in a 1 : 1 complex. Conditions: AKH- α R5W4^{NBD} and AKHK^{NBD}: 5 μ M; $A\beta_{1-16}$: 10 μ M; Cu: 5 μ M; 25 °C; $n = 3$ independent experiments. Experiments were carried out in DMEM diluted to 10% (10% DMEM) with salts: 0.2 g L⁻¹ CaCl₂, 0.0001 g L⁻¹ Fe(NO₃)₃, 0.098 g L⁻¹ MgSO₄, 0.4 g L⁻¹ KCl, 3.7 g L⁻¹ NaHCO₃, 6.4 g L⁻¹ NaCl, 0.11 g L⁻¹ NaH₂PO₄, and 4.5 g L⁻¹ D-glucose.

contribute to the ROS-based toxicity to cells as studied below. However, it did not hamper the protecting effect of AKH- α R5W4^{NBD} against Cu- $A\beta_{1-16}$ generated ROS, possible due to the much higher affinity to Cu(II) compared to Fe(III) of AKH- α R5W4^{NBD} (Fig. 2).

Altogether, these results indicate that both in the absence or presence of pre-incubation, AKH-LAH4^{NBD}, AKH- α R5W4^{NBD} or AKHK^{NBD} was able to retrieve Cu from $A\beta_{1-16}$ and lower ROS

production. As a control, we carried out the same experiment using Cu(II)- $A\beta_{1-42}$, which is the main toxic element of $A\beta$ and showed that AKH- α R5W4^{NBD} or AKHK^{NBD} was equally able to lessen ROS production *in vitro* (Fig. S9b and c†).

Protection of PC12 cells against Cu(II) induced toxicity. We further tested the ability of our peptides to protect PC12 cells from Cu(II)- $A\beta_{1-16}$ or Cu(II)- $A\beta_{1-42}$ toxicity in the presence of ascorbate. We pre-incubated in 10% DMEM 10 μ M Cu(II)- $A\beta_{1-16}$ /42 0.5 : 1 with 5 μ M AKH-CPP^{NBD} for 1 h, after which 300 μ M ascorbate was added to the mixture prior to incubation with PC12 cells for 24 h, with cell toxicity being evaluated using a MTT assay.

Treatment with AKH- α R5W4^{NBD} or AKHK^{NBD} effectively prevented Cu(II)- $A\beta_{1-16}$ toxicity, in line with Cu-binding and redox silencing. Of note AKH-LAH4^{NBD} seemed toxic to PC12 cells at 5 μ M when present in 10% DMEM (Fig. S11†), which was unexpected given that no toxicity was observed in 100% DMEM at that concentration (Fig. S6†). Recapitulating the *in vitro* data, there were almost no rescue effects from AKH-LAH4^{NBD}, AKH- α R5W4^{NBD} or AKHK^{NBD} on Cu(II)- $A\beta_{1-16}$ -induced cell toxicity without 1 h preincubation (data not shown). Indeed *in vitro* experiments above showed that the transfer of Cu from Cu(II)- $A\beta_{1-16}$ to AKH-CPP^{NBD} (Fig. 7) takes time, thereby allowing the production of a significant amount of ROS catalyzed by Cu(II)- $A\beta_{1-16}$ before complete transfer occurred.

We tested two different batches $A\beta_{1-42}$ to evaluate their toxicity on PC12 cells (individual batch data in Fig. S11b†). The incubation of $A\beta_{1-42}$ was toxic to PC12 cells at 10 μ M and the complexation of $A\beta_{1-42}$ with Cu(II) did not change the level of toxicity. However, upon addition of ascorbate, the latter showed a significant increase in toxicity on PC12 cells ($p < 0.0001$). An

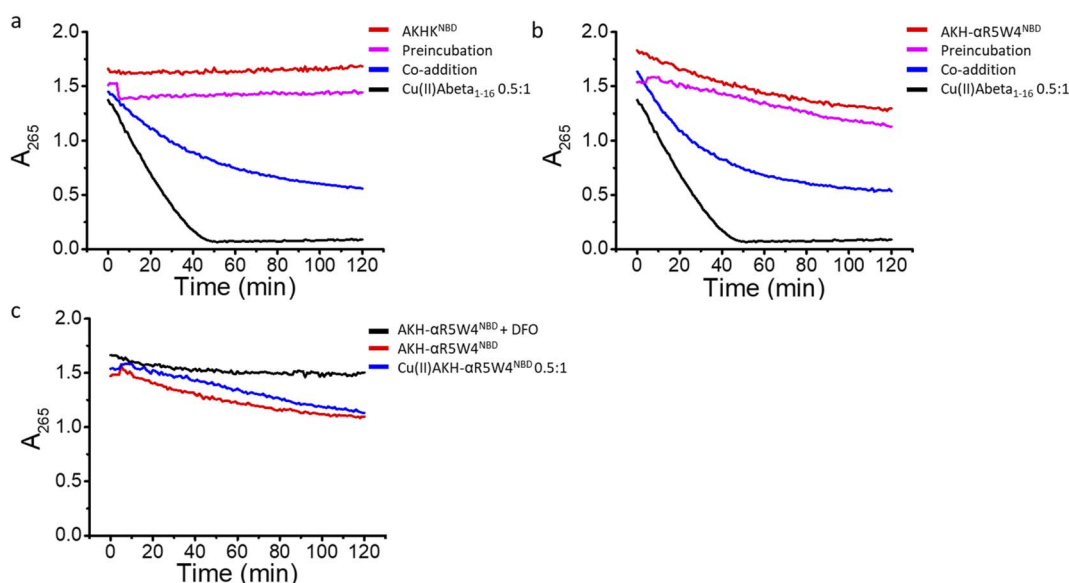


Fig. 8 Retrieval of Cu(II) from $A\beta_{1-16}$ by AKH- α R5W4^{NBD} or AKHK^{NBD} reduces ROS production. Inhibition of ROS production by (a) AKH- α R5W4^{NBD} or AKHK^{NBD} (b) monitored by the absorbance at 265 nm of ascorbate. Preincubation: 10 μ M Cu(II)- $A\beta_{1-16}$ 0.5 : 1 and 5 μ M AKH- α R5W4^{NBD} or AKHK^{NBD} was incubated for 1 h before the addition of ascorbate followed by absorbance measurement; co-addition: 10 μ M Cu(II)- $A\beta_{1-16}$ 0.5 : 1, 5 μ M AKH- α R5W4^{NBD} or AKHK^{NBD} is added together with ascorbate followed by absorbance measurement. (c) Comparison of ascorbate absorption in the presence of 5 μ M AKH- α R5W4^{NBD} with or without 10 μ M deferoxamine (DFO). Conditions: Asc: 100 μ M, AKH- α R5W4^{NBD} = AKHK^{NBD} = $A\beta_{1-16}$: 10 μ M, Cu: 5 μ M, DFO: 10 μ M; HEPES 100 mM, pH 7.4. $n = 2$ independent experiments.



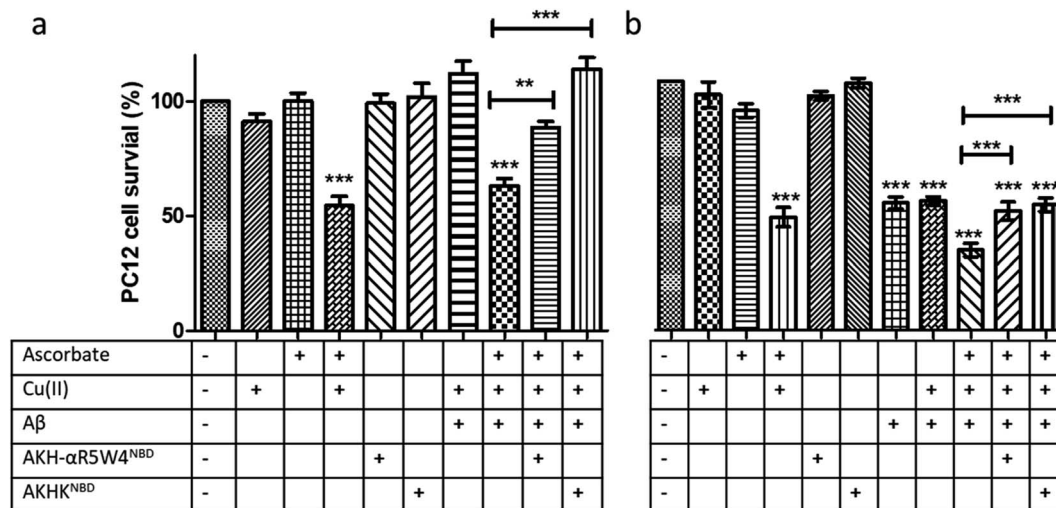


Fig. 9 Transfer of Cu(II) from Aβ to the ATCUN motif prevents Cu-induced ROS production and toxicity to PC12 cells. (a) Aβ₁₋₁₆ and (b) Aβ₁₋₄₂ (compilation of two batches, see Fig. S11†). 5 μM AKH-αR5W4^{NBD} or AKHK^{NBD} was pre-incubated with 10 μM Cu(II)-Aβ_{1-16/42} 0.5 : 1 in 10% DMEM in a test tube for 1 h before the addition of ascorbate and immediate administration on the PC12 cells. Experiments were done in triplicate, *n* = 3. A parametric ANOVA test was carried out with a Tukey post-test, ***p* < 0.001 and ****p* < 0.0001. Experiments were carried out in DMEM diluted to 10% (10% DMEM) with salts: 0.2 g L⁻¹ CaCl₂, 0.0001 g L⁻¹ Fe(NO₃)₃, 0.098 g L⁻¹ MgSO₄, 0.4 g L⁻¹ KCl, 3.7 g L⁻¹ NaHCO₃, 6.4 g L⁻¹ NaCl, 0.11 g L⁻¹ NaH₂PO₄, and 4.5 g L⁻¹ D-glucose.

Table 2 Operational conditions of ICP-MS analysis

ICP-MS	Agilent 7700
RF power	1550 W
Sampling and skimmer cones	Nickel
Nebulizer	MicroMist
Spray chamber	Scott double-pass, quartz
Plasma gas	15.0 L min ⁻¹
Nebulizer gas	1.03 L min ⁻¹
Makeup gas	0.00 L min ⁻¹
Auxiliary gas	0.9 L min ⁻¹
Spray chamber temp	2 °C
Sampling depth	8 mm
Dwell time	0.003 s
Isotopes	⁶³ Cu ⁶⁵ Cu ¹¹⁵ In
ORS mode	He collision ORS3
Nebulizer pump	0.10 rps

hour pre-incubation of Cu(II)-Aβ₁₋₄₂ with either AKH-αR5W4^{NBD} or AKHK^{NBD} rescued the toxicity induced by the Cu, although the toxicity due to Aβ₁₋₄₂ alone was not remediated as expected (Fig. 9c).

Conclusions

Re-equilibrating Cu homeostasis in neurodegenerative pathologies such as AD is a complex task, but it deserves to be addressed based on the potentially important outcomes, such as reducing oxidative stress and Aβ toxicity. This involves the design of a molecule that is highly selective to Cu(II) in comparison to other more concentrated bioavailable metals in the brain such as Zn(II) and Fe(III). Equally the molecule has to have a high enough affinity for Cu(II), to withdraw the latter

from the Cu(II)-binding domain of Aβ (Aβ₁₋₁₆) in amyloid plaques in the brain but not from other essential Cu(II)-proteins, such as ceruloplasmin. Finally, the molecule should be capable of penetrating the plasma membrane of neuronal cells as well as releasing imported Cu into neurons or glia cells.

Probing here the potential of the AKH-CPP^{NBD} shuttle, we have been able to synthesize, for the first time, a Cu(II) peptide shuttle capable of binding specifically Cu(II) in biological media in the presence of Zn(II), Mn(II) and Fe(III). This shuttle is selective for Cu(II) since it is based on the ATCUN motif that binds very weakly to other essential metal ions under extracellular relevant conditions. This differs from CQ and PBT2, used in clinical trials in the treatment of AD, which are able to shuttle Zn as well. The absence of selectivity of these shuttles for Cu makes it difficult to study the effect on Cu dyshomeostasis singularly in several pathologies including AD.

Cu(II) bound to the AKH-CPP^{NBD} shuttle is also capable of being reduced to Cu(I) by a physiological concentration of GSH *in vitro*. This seems also to occur *in cellulo*, as suggested by the translocation of ATP7A from the TGN to vesicles in proximity to the plasma membrane. This is in line with a Cu(II) import by AKH-αR5W4^{NBD}, reduction to Cu(I) by GSH and subsequent release in a bioavailable Cu(I) pool. Our construct AKH-αR5W4^{NBD} has several important features needed to re-distribute Cu from extracellular space to an intracellular bioavailable Cu pool, and hence can be an interesting tool for studying dyshomeostasis in Cu-related pathologies. In addition, AKH-αR5W4^{NBD} is capable of retrieving Cu(II) from Aβ₁₋₁₆, the metal-binding domain responsible for Cu complexation by Aβ_{1-40/42} as well as from the full length Aβ₁₋₄₂. In contrast to Aβ, AKH-αR5W4^{NBD} binds Cu(II) and stabilizes it, such that ascorbate oxidation and concomitant ROS production are precluded. In line with this, AKH-αR5W4^{NBD} is able to protect PC12 from



cells from Cu(II)-A β induced damage in the presence of ascorbate. However, the transfer of Cu from A β to AKH- α R5W4^{NBD} is relatively slow, and hence protection is not immediate. In conclusion, the shuttle described here might serve as a seminal scaffold to identify derived shuttles with faster Cu(II) complexation in future studies, as well as in studies with more elaborated cell models related to AD.

Materials and methods

Materials

All compounds used during this study were purchased from accredited merchants unless stated otherwise: Dulbecco's modified Eagle's medium-high glucose (Sigma, D5796), horse serum (Gibco, 26050070), Fetal calf serum (Gibco, 10270-160), penicillin/streptomycin (Sigma, P4458), Trypsin (Gibco, 25300-054), MTT (Fisher Scientific, 10133722), anti-giantin antibodies (Abcam, ab24586), monoclonal anti-ATP7A antibodies (Abcam, ab13995), secondary antibodies (Life sciences) and Hoechst (Thermo Fisher, 62249). Batch 1 A β ₁₋₄₂ (AnaSpec, >95% purity) and Batch 2 A β ₁₋₄₂ (Genecust, >95% purity).

Peptide synthesis

Peptide synthesis was carried out according to a general SPPS protocol.⁶³ The peptide synthesis was carried out using the Biotage® Initiator+ Alstra™. A Fmoc-Rink-Amide Tentax XV RAM resin (charge of 0.24 mmol g⁻¹) was used for the synthesis of all peptides. 0.5 M 1 : 1 ethyl cyano(hydroxyimino)acetate (Oxyma)/diisopropylcarbodiimide (DIC) was used for activation, and 2 M DIEA dissolved in *N*-methyl-2-pyrrolidone (NMP) was used as the base. 5 M anhydride acetic acid was used to cap unreacted free amino groups.

The reaction time was 2 × 30 minutes coupling at ambient temperature and 2 × 5 minutes coupling when using microwaves (μ WF) for compatible amino acids at 75 °C with the exception of cysteine, basic amino acids, and glycine after acidic amino acids. Fmoc deprotection was carried out using μ WF at 75 °C. Lysine was added as the *N* ϵ -allyloxycarbonyl-protected (Fmoc-Lys (Alloc)-OH) form to allow specific sidechain deprotection in order to graft the fluorophore NBD. The Alloc protecting group of the lysine side chain is deprotected using tetrakis(triphenylphosphine)palladium(0), which is a palladium-based mechanism.⁶⁴ The NBD fluorophore is added to the free lysine by nucleophilic substitution of 4 eq. NBD-Cl in the presence of DIEA. Finally, deprotection was carried out using TFA in the presence of scavengers (H₂O and triisopropylsilane).

Preparation of stock solution

All peptide stock solutions were prepared in ultra-pure Milli Q water ($\rho = 18.2 \text{ M}\Omega \text{ cm}^{-1}$) and were stored at -20 °C. The pH of the solutions was maintained at \approx pH 2, except when complexed to Cu(II) in which case they were buffered at pH 7.4 using 100 mM HEPES.

Fluorescence spectroscopy

The emission of NBD on AKH-CPP was taken in the presence and absence of Cu(II) using a Horiba Fluoromax plus fluorimeter using a 500 μ L 1 cm path quartz cuvette. 4 μ M of each peptide was titrated by adding 0.1 equivalent Cu into the cuvette (Fig. 1). AKH- α R5W4^{NBD}; excitation slit: 4 mm; emission slit: 3 mm, AKH-LAH4^{NBD} and AKHK^{NBD}; excitation slit: 2 mm; emission slit: 2 mm.

Ascorbate assay (UV/Vis spectroscopy)

Using a CLARIOstar (BMG LABTECH) plate reader, experiments were carried out in a UV-Star® 96 well microplate (Greiner), Half Area, clear. The absorbance of 100 μ M ascorbate (DO \approx 1.5) at 265 nm was followed under different conditions to indirectly follow ROS production. The final volume per well was 100 μ L and was buffered at pH 7.4 using 100 mM HEPES.

Cell culture

PC12 cells were maintained in culture in a DMEM high glucose medium (4500 mg mL⁻¹) with 10% horse serum, 5% FBS and 1% penicillin/streptomycin as described previously.⁶⁵ The cells were split once a week using trypsin for detachment and replaced by a fresh batch of cells after a maximum of 15 splits.

Immunohistochemistry and live imaging

PC12 cells were cultured on poly-D-lysine-coated glass coverslips in NUNC 4-well plates at a density of 5 × 10⁴ per well, 24 h before experiments. After the experiment, the medium was removed and the wells were washed with 1X PBS, and the cells were fixed with 4% paraformaldehyde (Sigma-Aldrich) for 10 min at room temperature. The cells were permeabilised in 0.1% Triton X-100 (Thermo Fisher Scientific) for 10 min followed by 1 h saturation in 1X PBS containing 0.2% Triton X-100, 10% donkey serum (Sigma-Aldrich) and 3% Bovine Serum Albumin (BSA, Sigma - Aldrich) at 37 °C. The cells were then incubated overnight at 4 °C with primary antibodies, and after washing in PBS 1X were incubated with secondary antibodies and Hoechst for 45 min at 37 °C before being rinsed and mounted.

For live imaging, PC12 was grown in glass bottom Ibidi chamber coverslip cells and maintained in Locke solution (140 mM NaCl, 4.7 mM KCl, 2.5 mM CaCl₂, 1.2 mM KH₂PO₄, 1.2 mM MgSO₄, 11 mM glucose, 0.56 mM ascorbic acid, 0.01 mM EDTA and 15 mM HEPES, pH 7.5) for recording. Fluorescent peptides are added < 30 seconds before recording for a duration of 30 minutes at 10 frames/min.

MTT assay

A MTT stock solution at 5 mg mL⁻¹ was prepared in PBS 1X. The cells were incubated with a final MTT concentration of 0.5 mg mL⁻¹ in DMEM media for 4 h at 37 °C after which the medium was removed, and the formazan precipitate dissolved in DMSO. The absorption was recorded at 595 nm with a plate reader.



ICP-MS

Lysis. The cell pellet was resuspended in 500 μL 1X PBS, from which 10 μL was taken to determine the protein concentration by Bradford analysis. The remaining cells were mixed in 300 μL ultrapure HNO_3 , 70% in glass vials pretreated with 3 mM ultrapure HNO_3 . For complete cell lysis, the mixture was heated for 1 h at 90 $^\circ\text{C}$ and left at room temperature until ICP-MS measurements.

Measurement. Operational conditions for Cu analysis are given in Table 2. The limit of detection is 0.1 $\mu\text{g kg}^{-1}$ for both isotopes ^{63}Cu and ^{65}Cu . A standard addition calibration curve was used to avoid matrix effects observed with an external calibration. Indium 10 ppb was used as the internal standard. The specificity of the measurement was checked using the recovery yields of spiked samples. The recovery values were 95–105% for both isotopes.

Statistical analysis

All analyses were carried out using a GraphPad prism. The data set underwent the Shapiro test for normality. For comparisons of the mean, a parametric test with Dunnett's multiple comparison post-test or non-parametric Kruskal–Wallis with Dunn's multiple comparisons test was performed.

Data availability

We added as much as possible of the experimental data in the ESI.†

Author contributions

Investigation: M. O., P. G., P. R., and I. E.; writing – original draft: M. O.; supervision: A. B., L. R., N. V., and P. F.; initial conceptualization: C. H.; funding acquisition: C. H., N. V., and P. F.; writing – review & editing: L. R., C. H., N. V., and P. F.; methodology: L. R., N. V., P. F., S. C.-G., S. O., and S.G.

Conflicts of interest

The authors declare no conflict of interest.

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