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Misfolded amyloid-β-42 impairs the endosomal–lysosomal pathway

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Abstract
Misfolding and aggregation of proteins is strongly linked to several neurodegenerative diseases, but how such species bring about their cytotoxic actions remains poorly understood. Here we used specifically-designed optical reporter probes and live fluorescence imaging in primary hippocampal neurons to characterise the mechanism by which prefibrillar, oligomeric forms of the Alzheimer’s-associated peptide, Aβ42, exert their detrimental effects. We used a pH-sensitive reporter, Aβ42-CypHer, to track Aβ internalisation in real-time, demonstrating that oligomers are rapidly taken up into cells in a dynamin-dependent manner, and trafficked via the endo-lysosomal pathway resulting in accumulation in lysosomes. In contrast, a non-assembling variant of Aβ42 (vAβ42) assayed in the same way is not internalised. Tracking ovalbumin uptake into cells using CypHer or Alexa Fluor tags shows that preincubation with Aβ42 disrupts protein uptake. Our results identify a potential mechanism by which amyloidogenic aggregates impair cellular function through disruption of the endosomal–lysosomal pathway.

Keywords Amyloid-beta · Self-assembly · Oligomer · Toxicity · Lysosomes · Endocytosis

Introduction
Neurodegeneration occurs in several diseases that are characterised by protein aggregation, including Alzheimer’s, Parkinson’s and Huntington’s disease. In each, there is a strong association between the self-assembly of a particular protein to form amyloid fibrils and the development or progression of the disease [1–3]. However, although this ordered self-assembly is tightly linked with cell death, precisely what constitutes a “toxic” structure, and which molecular mechanisms drive detrimental effects, are still unresolved issues.

In Alzheimer’s disease (AD), the pathological state is characterised both by the deposition of tau in neurofibrillary tangles, and Amyloid-β (Aβ) into amyloid plaques.

The self-assembly and accumulation of Aβ, in particular the 42-residue form [4, 5], is thought to be a very early event in the progression of Alzheimer’s disease, occurring before changes in tau take place or the expression of cognitive impairment [6, 7]. In support of this, compelling genetic evidence links Aβ aggregation to AD [8], and prefibrillar oligomers of Aβ42 are implicated as critical triggers of neuronal cell death [9–11]. Additionally, previous studies have shown that synthetic oligomers of Aβ42 cause synaptic dysfunction and neuronal death in a range of model systems both in vitro and in vivo [1, 12–19].

What is lacking in the current models, however, is a clear understanding of the pathway that links Aβ42 aggregation with the compromised cell health associated with the disease state. Recent work offers an important insight; specifically the observation that proteostasis mechanisms, normally operating in low pH lysosomes to degrade macromolecules and maintain cellular health, appear to be defective in AD and other disease states [20–30]. This raises the possibility that Aβ might be specifically exerting its toxicity through its ability to internalise and affect lysosomal pathways.

Here we investigate this key idea directly, using live fluorescence imaging methods both to characterise the uptake and localisation of Aβ42 oligomers in primary hippocampal neurons and to assess their impact on the endosomal–lysosomal pathway. In particular, we exploit a new pH-sensitive
optical reporter to show that Aβ oligomers rapidly internalise into cells in a dynamin-dependent manner, and then traffic directly to lysosomes where they remain for several days. Protein misfolding is critical for this outcome since a non-assembling Aβ variant undergoes minimal uptake and accumulation with the same assay. By imaging events occurring in real-time using fluorescent probes, we also demonstrate that the presence of Aβ42 disrupts endocytosis of other proteins. Our findings provide a key link between Aβ42 aggregation and deficits in the endosomal–lysosomal pathway, which may account for early events in disease-related neuronal dysfunction.

Results

Toxic Aβ42 oligomers are internalised in defined intracellular compartments

Self-assembly of the Aβ42 peptide into amyloid fibrils is strongly linked with neurodegeneration and the subsequent symptoms observed in AD [1]. As a starting point, we evaluated the time and dose dependent effect of oligomeric Aβ42 [15] on cell viability in primary hippocampal neurons using the Readyprobes assay (Fig. 1a, b). Compared to buffer control, significant differences were observed after incubation with 10 μM Aβ oligomers at all time points, with an increase in cell death from 26.0 ± 3.7% (n total = 1005) to 65.2 ± 8.2% (n total = 568) (Fig. 1b) from 24 h to 14 days incubation respectively. Previous work using an Aβ control peptide at the same concentration showed that toxic effects are specific to wild type Aβ42 [15]. To examine the pathway of action that leads to neuronal cell death, we selected one of the intermediate timepoints (3 days, 44% cell death) for further experiments.

Initially, we monitored real-time uptake and trafficking of Alexa Fluor-conjugated Aβ42 oligomers [15, 16]. Importantly, our previous work has demonstrated that Alexa Fluor labelling does not alter the self-assembly of Aβ42 into oligomers or amyloid fibrils [16] and that the fluorescently labelled particles are detected by anti-Aβ antibodies [15]. Figure 2a and supplementary movie 1 show the uptake of Alexa Fluor 488 labelled Aβ42 oligomers into primary neurons over 15 h. Small fluorescent puncta could be readily observed at the 1 h timepoint (Fig. 2a, arrows) and by 5 h these were clearly visible in the cytoplasm suggesting that oligomers were being internalised via the plasma membrane into intracellular compartments. This continued in a time-dependent manner (Fig. 2a–c) and at the 3-day incubation timepoint, there was extensive accumulation (Fig. 2b, arrows) with 100% of cells (n = 72) containing visible Aβ42-488 puncta in the cell cytoplasm. Additionally, we saw evidence of fluorescence labelling along the cell boundaries (Fig. 2b, orange arrowheads), indicative of binding of Aβ42-488 oligomers to the extracellular membrane.

Aβ42 internalises in a dynamin-dependent manner into lysosomes

To determine the mechanism of Aβ42-488 oligomer uptake, we tested whether the observed internalisation is sensitive to endocytic blockers using sucrose and chlorpromazine as green, i.e. dead cells can be seen in oligomer treated cultures (white arrows).
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generic, non-specific inhibitors, and dynasore as a specific inhibitor of the GTP-ase, dynamin. It is well-established that the protein transferrin endocytoses via clathrin-coated pits [31, 32] and this was used as a control protein. Serum-starved cells incubated for 15 min with Aβ42-488 oligomers alone showed bright punctate and diffuse labelling indicative of their association with neurons and internalisation (Fig. 2d, left panel). However, pre-treatment with any of the endocytic inhibitors yielded a striking reduction in fluorescence (Fig. 2d, f) similar to the pattern of distribution in the positive control, transferrin-488 (Fig. 2e,g). These results suggest that Aβ42 oligomers, like transferrin, are endocytosed by a dynamin-dependent mechanism.

Next, we investigated the nature of the cellular structures that accumulate Aβ42 oligomers. To confirm the destination for Aβ42 oligomers following endocytosis, neurons were treated with Alexa Fluor 488-labelled Aβ42 oligomers and co-labelled with Lysotracker red (Fig. 3a), an established marker of lysosomes. As a control, we used a fluorescent conjugate of ovalbumin, a protein known to traffic through the endosomal–lysosomal pathway [33] (Fig. 3b). We found that both protein signals co-localised to lysosomes labelled with Texas Red-dextran using an Aβ oligomer specific antibody (NU1) (Supplementary Fig. 1). As further support for this conclusion, we also examined neurons treated with Aβ42 oligomers using transmission electron microscopy (TEM). Aligned with our fluorescence data we observe the presence of large (> 1 μm diameter) round electron-dense vesicular structures.
structures in the cell cytoplasm (Fig. 3c, arrows). These structures, which were not seen in control neurons (Supplementary Fig. 2), were consistent in appearance with enlarged lysosomal structures reported previously in dystrophic AD neurons [22, 34–36]. Taken together, these abnormalities, specific to oligomer-treated cells, support the conclusion that Aβ42 oligomers have a detrimental effect on the cellular pathways that end with lysosomal degradation.

**A novel pH-dependent probe provides a real-time assay of Aβ42 uptake into lysosomes**

Next, to provide detailed temporal information on the lysosomal fate of oligomers, we developed a novel optical reporter that provides a real-time readout of Aβ internalisation into acidic compartments. Specifically, we used CypHer5E, a commercially available pH-sensitive fluorescent probe, which exhibits maximal emission when present in an acidic environment [37]. In previous studies this fluorescent moiety has been used to label antibodies for reporting receptor internalisation [38] and to tag vesicle proteins for assaying exo-endocytic recycling at synaptic terminals [39–41].

The endo-lysosomal pathway is characterised by a shift in pH, reducing from ~6–7 in early endosomes to ~5 in the lysosome [42, 43]. Therefore, we hypothesised that a protein tagged to CypHer5E that experienced this environmental change would yield important temporal information on internalisation seen as an increase in fluorescence emission, and thus allow us to dynamically monitor uptake.

To validate this approach, we conjugated CypHer5E with ovalbumin (referred to as Ova-Cy) (Fig. 4a). When Ova-Cy was applied to neurons, we observed a time-dependent increase in punctate fluorescence localised to cell bodies, consistent with the presence of this reporter in a cellular compartment that undergoes a drop in pH (Fig. 4b, d). To corroborate this finding, we also tested the effect of pre-treating with NH₄Cl, a weak base that increases the pH in the lysosome, prior to the addition of Ova-Cy. In this case, a fluorescence increase was not observed, confirming that CypHer5E is an effective monitor of basal lysosomal acidification state (Fig. 4c, d).
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Moreover, the fluorescence was reduced when NH₄Cl or bafilomycin, an ATPase blocker, were added to cells following addition of Ova-Cy to primary hippocampal neurons. No fluorescence increase is observed in cells pre-treated with NH₄Cl, a weak base that raises the pH of acidic compartments. One 1.0-μm slice is shown. Scale bar is 25 μm. Quantification of mean fluorescence intensity from cells shown in b and c. Each trace represents an individual cell body either untreated (grey, n = 34) or pre-treated with 10 mM NH₄Cl for 30 min (purple, n = 28).

Data pooled from three independent experiments, 3–4 FOV per experiment. Colour bar for look-up table indicates mean fluorescence intensity. Fluorescence arising from Ova-Cy diluted into buffer of pH 5.4 (purple) or 7.4 (turquoise). Ova-Cy diluted into buffer containing NH₄Cl, pH 7.4 is also shown (orange). The absorbance maximum for CypHer5E at physiological pH 7.4 is 500 nm (left plot) and at pH 5.4 (similar to the pH in a lysosome) is approximately 645 nm. Ova-Cy in acidic (pH 5.4) buffer has increased intensity over neutral pH when excited with 644 nm light. The presence of NH₄Cl has no effect on the spectral properties of the dye.

Next, we used the same tagging method to examine the fate of internalised Aβ (Fig. 5). Aβ42 was conjugated to CypHer5E (Aβ42-Cy, Fig. 5a and Supplementary movie 2) and applied to neurons, resulting in an increase in fluorescence intensity over a 150-min period, comparable to our findings with Ova-Cy (Fig. 5a, e). Notably, the distribution of fluorescent puncta in the cytoplasm was consistent with our observations for both Ova-Cy (Fig. 4b) and Aβ42-488 (Figs. 2b, 3a) suggesting that all were localised in the same way. CypHer tagged Aβ42 oligomers lead to a comparable level of cell death (51%) to untagged oligomers (see Fig. 1, 44.8%) and in our previous report ([15], 46.9%) (Supplementary Fig. 4).
To examine whether uptake is specific to prefibrillar misfolded forms of Aβ42, or alternatively a generic outcome with Aβ proteins, we took advantage of a non-assembling form of Aβ42 (variant Aβ42 or vAβ42), which we have recently developed as a control peptide [15]. Importantly, we have previously demonstrated that this variant structure has none of the detrimental effects of Aβ42 with regard to neuronal biology or memory formation [15]. As above, CypHer5E was conjugated to vAβ42 and applied to cells and monitored over 150 min. Compared to cells treated with Aβ42-Cy, only a negligible fluorescence increase was observed in cultures treated with vAβ42-Cy and no change was observed over time (Fig. 5d, e). The toxicity of vAβ42-Cy was also examined (25.4%, Supplementary Fig. 4) and found to be similar to levels induced by the untagged peptide we previously reported (21.7%, [15]).

**Aβ42 oligomers inhibit uptake of ovalbumin in primary neurons**

We have shown that Ova-Cy fluorescence effectively reports the pH drop experienced by the protein trafficking through the endosomal system to lysosomes. This provides us with a powerful assay for monitoring the potential impact of Aβ42 oligomers on endosomal–lysosomal pathway function. To investigate this, cells were pre-treated with Aβ42 oligomers before the addition of Ova-Cy. Strikingly, following Aβ42 exposure, the fluorescence intensity of Ova-Cy was clearly lower than cells pre-treated with buffer alone (Fig. 6a–c). To ensure these effects were specific to Aβ42 oligomers and not just due to general overloading with high concentrations of protein, we also performed an additional control experiment (Supplementary Fig. 5). Neurons were pre-incubated with ovalbumin in place of Aβ42 at the same molar concentration (10 μM) for the same duration (72 h). Ova-Cy uptake was monitored and no difference observed in the internalised Cypher fluorescence intensity compared with buffer, suggesting that it is the Aβ42 oligomers that specifically slow the internalisation of Ova-Cy. These results provide strong support for a mechanism in which Aβ42 oligomers first traffic to lysosomes and then influence their function.

To examine whether the reduced Ova-Cy fluorescence intensity is due to an altered pH or the amount of Ova internalised, we also carried out a complementary experiment in which we tagged ovalbumin with Alexa Fluor 488 (Fig. 6d–f). Analogous to the Ova-Cy experiments, we
observed a robust reduction in fluorescence intensity in cells pre-treated with Aβ42 oligomers compared to control, showing that the uptake and trafficking of Ova is impaired. Taken together, our findings suggest that the presence of intracellular Aβ42 causes a slowing of the endosomal–lysosomal system, which may be important for the mechanism of Aβ42 toxic action on living cell function.

Discussion

The ordered assembly of peptides and proteins into amyloid is strongly linked with neurodegeneration in vivo and in vitro [44, 45]. Understanding the cause-and-effect relationship between protein self-assembly and neuronal dysfunction is challenging since early events are difficult to capture or recapitulate. Nevertheless, the case for protein misfolding as a cause of neuronal loss is compelling [11, 46–48]. On the strongly evidenced premise that misfolded proteins are able to damage neurons, we developed an experimental model using oligomers of the Alzheimer’s-associated Aβ42 peptide in conjunction with novel applications for fluorescent probes to explore the link between oligomer exposure and degeneration of hippocampal neurons. Benefiting from time-lapse imaging of specially engineered optical reporters in living neurons, we characterised structural and functional properties of cells in real-time as a powerful strategy to elucidate the underlying mechanisms of aggregated peptide-induced neuronal cell death. Initially, we confirmed that the synthetic, pre-fibrillar Aβ42 oligomers were toxic to primary hippocampal neurons, consistent with previous findings [1, 2, 49]. Using fluorescence imaging, we then examined the endosomal–lysosomal physiology of individual live cells that contained endocytosed Aβ42 oligomers. This real-time readout allowed us to reveal dysfunctional mechanisms prior to cell death, providing important insights into early events that can lead to neurodegeneration. Specifically, we found that oligomers are internalised into neurons predominantly by a dynamin-dependent mechanism and that these are then trafficked to lysosomes and remain intracellular for several days.

It is well established that the acidification of lysosomes is critical to their function [50]. In order to understand the potentially direct effects of Aβ42 oligomers on lysosomal function, we developed a method to tag Aβ42 and ovalbumin with a pH-sensitive probe (CypHer, Roche) that selectively fluoresces in acidic compartments. Using this approach, the uptake and localisation of Aβ42-Cy peptides within the cell could be monitored in real-time and compared with an assembly-incompetent variant of Aβ42. In particular, our findings showed that this uptake is specific to self-assembling, wild-type Aβ42.

To explore the downstream effects of accumulated Aβ42 on the endo-lysosomal pathway, we utilised tagged versions of ovalbumin to assay uptake and acidification events in cells pre-treated with Aβ42 oligomers. We found that the levels of Ova-488 and Ova-Cy fluorescence were markedly reduced in Aβ42 treated cells compared to controls, implying that the accumulated Aβ oligomers are affecting the endo-lysosomal pathway. Because we only considered live cells in this analysis (those with an intact and smooth plasma membrane and a clearly defined nucleus, where often nucleoli were visible), we can be confident in concluding that the impairment of the endosomal–lysosomal system occurs prior to cell death. This implicates a potentially catastrophic and ultimately fatal effect of Aβ42 on neuronal cells.

How does internalised Aβ42 impair the endosomal–lysosomal pathway? We propose that a number of mechanisms could conceivably be involved. These include a direct effect of Aβ42 on acidification through action on the ATPase [26, 27], through lysosomal leakage associated with membrane interactions [30, 51, 52], or simply through its accumulation and overloading of lysosomal proteolytic capacity [53–55]. In our study, we clearly demonstrate that a consequence of Aβ42 internalisation is an inhibition of further endocytosis resulting in a reduction in ovalbumin internalisation. This result implies that the oligomers impair the endo-lysosomal pathway and thus stall a process critical for cellular health, although our experiments do not allow us to definitively establish the mechanism by which this outcome arises. Incubating the cells with another, non-toxic, protein did not lead to any detrimental effects on endocytosis providing strong evidence that it is Aβ42 oligomers specifically that are causing damage, and not just general endocytic or lysosomal burden. It is interesting to note the presence of Aβ42 on the extracellular side of the cell membrane, which could provide a site for oligomeric binding, or further aggregation [56]. In either case, it is conceivable that the accumulation of aggregated proteins along the plasma membrane could perturb endocytosis and might be another potential mechanism for our observations.

Collectively these results outline a mechanism of Aβ42-induced cell death that leads to the disrupted endocytosis of essential proteins and nutrients. Indeed, they offer a mechanistic basis for previous in vivo studies that have reported defects in lysosomal function in neurodegenerative diseases, including the appearance of abnormal lysosomes and autophagosomes within dystrophic neurites in human AD and mouse model brains [21, 26, 34, 35, 57]. Abnormalities to the endocytic pathways in neurons from AD brain tissue have also been observed [58]. Here we have gained key information on the specific pathway of action by which misfolded proteins exert toxic action on cells whereby uptake of Aβ42 oligomers into neurons renders them unable to carry out further endocytosis. We suggest that the accumulation
Fig. 6 Pre-treating neurons with Aβ42 oligomers inhibits uptake of ovalbumin. Cells were treated either with 10 μM Aβ42 oligomers or equivalent volume of buffer for 72 h, followed by addition of 25 μg/mL Ova-Cy (a) or Ova-488 (d). Quantification of mean fluorescence intensity from Ova-Cy (b) or Ova-488 (e) in the cell body (one 0.5 μm slice) over 150 min from cells either treated with buffer (blue) or pre-treated with 10 μM Aβ42 oligomers for 72 h (red). Each trace arises from the same individual cell measured over time. Data pooled from three separate experiments 4–9 FOV per experiment, n (cells) buffer = 84, Aβ42 = 19 (b), 4–5 FOV per experiment, n (cells) buffer = 53, Aβ42 = 33 (e). Example images of live cells treated with Aβ42 oligomers (top) or equivalent volume of buffer (bottom) followed by Ova-Cy (c) or Ova-488 (f), taken at increasing time points. Fluorescence intensity is only reduced in cells treated with oligomers. Arrows point to neuronal cell bodies. One 0.5-μm slice is shown. Scale bar is 25 μm.
of aggregated Aβ42 within lysosomes stalls the endosomal system and that this is an early event in oligomer-induced neurodegeneration and collectively provides an explanation for cytotoxicity caused by misfolded proteins.

**Methods**

**Peptide preparation**

**Aβ42 oligomers and vAβ42**

Hexafluoroisopropanol (HFIP) films of recombinant Aβ1-42 were purchased from rPeptide and vAβ1-42 was purchased from JPT Peptide Technologies. Peptides were prepared using an adapted protocol [59], which ensures the removal of potentially contaminating trace solvents. HEPES buffer (10 mM HEPES, 50 mM NaCl, 1.6 mM KCl, 2 mM MgCl₂, 3.5 mM CaCl₂) was used to mimic the culture media as previously described [16, 59, 60]. All preparation was conducted using protein LoBind Eppendorfs and tips (Alpha Laboratories). Briefly, 0.2 mg Aβ1-42 (rPeptide) was solubilised in 200 μL HFIP (Sigma-Aldrich) to disaggregate the peptide. The solution was then vortexed on high for one minute and sonicated in a 50/60 Hz bath sonicator for five minutes. HFIP was dried completely using a low stream of nitrogen gas. Once completely dried, 200 μL dry dimethylsulfoxide (DMSO) (Sigma-Aldrich) was added, vortexed for one minute, and sonicated for one minute. Solutions were added to a 7 K MWCO Zeba buffer-exchange column (Thermo Scientific) equilibrated with HEPES buffer with 40 μL HEPES as a stacking buffer. The protein solution was kept on ice and the absorbance at 280 nm measured with a NanoDrop spectrophotometer using a molar absorption coefficient of 1490 M⁻¹ cm⁻¹. Solutions were immediately diluted to 50 μM with HEPES buffer and incubated where indicated or used immediately in further experiments.

**Alexa Fluor conjugated peptides**

Peptides were treated as outlined above up to the addition of 200 μL dry DMSO. Alexa Fluor tags were prepared as per the manufacturer’s instructions (Life Technologies). Briefly, 10 μL dH₂O was added to the Alexa Fluor TFP ester and kept on ice. This was added to the peptide in 200 μL DMSO along with 20 μL 1 M sodium bicarbonate pH 8.3 to increase the reaction efficiency, mixed and incubated for 15 min at room temperature protected from light. Following this, the procedure resumed as above, and the solution was then added to the Zeba buffer exchange column. The calculations for the concentration were adjusted to take into account the absorbance of the dye.

**Cell culture**

Rats were housed within a specialised facility under Home Office UK guidelines and killed using procedures in accordance with Animals (Scientific Procedures) Act 1986, Amendment Regulations 2012 and approved by a local ethical review panel. Primary neurons were prepared from the hippocampus of P0–P1 rats as described previously [61] by initially dissecting the tissue into ice cold HBSS containing 0.1 M HEPES. Following washes in pre-warmed basal Eagle’s Minimum (BME) (Gibco) containing 0.5% glucose, 2% FCS, 1 mM Na-Pyruvate, 0.01 M HEPES pH 7.35, 1% Penicillin–Streptomycin, 1% B27 supplement and 1% Glutamax, tissues were triturated using a 1 mL pipette until fully dissociated. The cell suspension was diluted further with complete BME media and approximately 40,000 cells plated into 2 cm² wells containing a coverslip coated in 20 μg/mL Poly-D-Lysine with a layer of hippocampal astrocytes that had been growing for 4–5 days. After 2–3 days, cells were treated with 3.25 μM cytosine arabinoside to curb further proliferation of astrocytes. Experiments were carried out on cells 10–14 DIV (days in vitro).

**Cell viability assay**

After the desired incubation time, one drop of each ReadyProbes reagent (Life Technologies) was added to cells in 500 μL of media. The kit contains a blue stain to label all cells, and a green stain to label dead cells only. Cells were incubated at 37 °C for 15 min and then imaged using widefield fluorescence microscopy with a Zeiss Cell Observer widefield microscope fitted with a Hamamatsu ORCA camera and a Nikon Plan Fluor 10X NA 0.3 air objective. DAPI fluorescence was captured using a G 365 excitation filter and a LP 420 emission filter with a FT 395 dichroic. Green fluorescence was captured using a FITC filter set (BP 450–490 excitation filter, BP 515–565 emission filter and FT 510 dichroic). Identical acquisition settings were used for all experimental and control samples. Images were analysed using FIJI. Neuronal cell bodies were identified using DIC and DAPI channels (astrocytes were excluded), which indicated total cell number (live and dead). Those that were also fluorescent in the FITC channel were defined as dead. The numbers of dead cells were expressed as a percentage of the total number of cells. At least three fields of view were imaged per sample in each experiment. Each experiment i.e. using newly prepared peptide added to different cultures, was performed at least three times.

**Aβ42 oligomer uptake and blocking**

All imaging experiments outlined below were carried out using a Leica SP8 confocal microscope and image
processing using FIJI. For overnight imaging, the Adaptive Focus Control feature was used to maintain constant focal planes throughout the course of the experiment. Where multiple fluorescent probes were imaged, samples were scanned sequentially to prevent spectral bleedthrough.

To monitor live uptake, neurons were plated on top of a layer of astrocytes in poly-D-Lysine coated 35 mm dishes with 1.5 coverslips (Mattek) and cultured for 10–14 days. Fluorescent-tagged oligomers were added and cells imaged live using a 63× numerical aperture (NA) 1.2 oil objective. The environment was maintained at 37 °C with humidified CO₂.

For blocking experiments, cells plated on coverslips in 24-well plates were washed with warm external bath solution (EBS; 137 mM NaCl, 5 mM KCl, 3 mM CaCl₂, 1 mM MgCl₂, 10 mM D-Glucose and 5 mM HEPES pH 7.3) then serum starved for 1 h in 500 μL warm EBS. To block endocytosis, cells were treated with one of the following for 30 min: 0.5 M sucrose, 10 μM Dynasore (Sigma), 25 μg/mL chlorpromazine or left untreated. After blocking, fluorescently labelled Aβ42 oligomers were added, or 100 μg/mL transferrin-488 as a control to confirm blocking effects. After incubation for 30 min, cells were washed 3–4 times quickly with warm EBS, fixed using 2% paraformaldehyde (PFA) in EBS for 15 min, rinsed with PBS then the coverslip mounted onto a glass slide using Prolong Gold with DAPI (Life Technologies). Fixed cells were imaged with a 40×NA 1.1 water objective.

**Lysosome labelling**

Ovalbumin-488 was purchased from Life Technologies and reconstituted in PBS to a stock concentration of 5 mg/mL. Hippocampal neurons plated on Mattek dishes were treated with Ova-488 to a final concentration of 100 μg/mL for 3 h or Aβ42-488 oligomers to 10 μM for 24 h, then 100 nM Lysotracker red (Life Technologies) was added, incubated for 1.5 h and imaged using a 100×oil objective.

For experiments using NU1 and Texas Red-Dextran, neurons were incubated with Aβ42 oligomers for 24 h then with 0.5 mg/mL 10,000 MW Texas Red-Dextran (ThermoFisher) for 1 h. Cells were washed twice with fresh media and incubated with neuron-conditioned media for a further 3 h to chase into lysosomes. Cells were then fixed with 0.3% PFA, permeabilised with 0.025% saponin, blocked with Superblock (ThermoFisher) containing 0.025% saponin and then labelled with NU1 (courtesy of the William Klein laboratory) and goat anti-Mouse Alexa Fluor 488 conjugated secondary (Invitrogen) both diluted in wash buffer (25% Superblock in PBS) at 1:500. Coverslips were mounted in Prolong Gold containing DAPI and imaged using a Leica confocal SP8 microscope.

**Transmission electron microscopy**

Cells grown on coverslips were washed once with EBS then fixed in 2% paraformaldehyde, 2% glutaraldehyde in PBS for 15 min at room temperature. The fixative was replaced with 100 mM glycine and incubated for 1 h followed by a 1 min wash in 100 mM NH₄Cl. Cells were then washed twice with 0.1 M sodium cacodylate buffer before a second fixation step in 1% osmium tetroxide with 1.5% potassium ferrocyanide in sodium cacodylate for 1 h. Following repeated rinses with 0.1 M sodium cacodylate, 10 mg/mL tannic acid was added (45 min) to stabilize and enhance contrast, before a further incubation in 10 mg/mL anhydrous sodium sulphate (5 min). The cells were rinsed with water (5 repeats) and dehydrated using increasing concentrations of ethanol (50%, twice with 70%, twice with 90% then three times with 100%, 5 min each). The coverslips were then transferred into a 1:1 mixture of polypropylene oxide and EPON resin for 1 h, then moved into pure EPON resin (Electron Microscopy Sciences) (exchanged after 2 h) and left overnight. The following day, the coverslip was placed cell-side down onto an EPON block and dried at 60 °C overnight until polymerised. The coverslip was removed by submerging in liquid nitrogen and the block sectioned (70 nm thickness). Sections were viewed on a JEOL transmission electron microscope (100 kV).

**Conjugation of CypHer5E to ovalbumin and Aβ peptides**

CypHer5E Mono NHS Ester (Roche) was prepared as per the manufacturer’s instructions. Briefly, 100 μL of dimethyl sulphoxide (DMSO) was added to 1 mg CypHer5E and the solution sonicated in a 50/60 Hz bath sonicator for 15 s. 5 μL of dye was diluted into 4 mL PBS-sodium bicarbonate (0.5 M) pH 8.3 and the absorbance at 500 nm measured. The concentration of dye was calculated based on a molar extinction coefficient of 40,000. Chicken egg white albumin was purchased from Sigma and dissolved to a concentration of 10 mg/mL in water. The concentration was confirmed using absorbance readings at 280 nm and an extinction coefficient of 32,176. The solution was diluted to 1 mg/mL in PBS-sodium bicarbonate (0.5 M) pH 8.3 and a 20-M excess of CypHer5E dye added. The solution was protected from light and gently agitated on a rocker for 1 h. Following the labeling reaction, unconjugated dye was separated and removed from the solution using a purification column packed with BioGel P-30 fine size exclusion purification resin (BioRad), which is designed to separate free dye from proteins with a molecular weight of greater than 40,000 kDa (MW ovalbumin = 45,000 kDa). Elution buffer (10 mM potassium phosphate, 150 M sodium chloride pH 7.2) was run through the column and once excess buffer had drained into the column
bed the reaction mixture was added, allowed to enter into the resin, then eluted using elution buffer. The labelled protein (Ova-Cy) was collected and the concentration of protein and dye measured using a Nanodrop spectrophotometer. The stock solution was kept at 4 °C for a maximum of 3 months. For Aβ42 and vAβ42, labelling with CypHer5E was performed using a protocol similar to the Alexa Fluor labelling method described previously. CypHer5E dye was prepared as described above and a 20 M excess added to the peptide dissolved in 200 μL DMSO along with 20 μL 1 M sodium bicarbonate pH 8.3. Briefly, the reaction mixture was incubated for 15 min, then applied to a Zeba buffer-exchange column equilibrated with HEPES to remove the DMSO and free dye. The protein and dye concentrations of the eluted solutions were measured, diluted to 50 μM (protein), incubated for 2 h to form oligomers (for WT Aβ42) and then diluted to working concentration.

**Live-cell imaging of neurons treated with CypHer5E conjugated proteins**

Neurons were plated on 35 mm Mattek dishes as described previously. After 10–14 DIV, cells were either treated with Aβ42 oligomers, Ova-Cy, Aβ42-Cy, vAβ42-Cy or an equivalent volume of HEPES buffer. All imaging was carried out live in media at 37 °C with humidified CO₂ using a 1.4 NA HC PL APO CS2 63× oil objective with a 633-nm laser line and fluorescence emission collected between 650 and 700 nm with a PMT detector. For experiments assessing pH dependence of CypHer, Ova-Cy was added to a final concentration of 10 μg/mL with or without 50 mM NH₄Cl and imaging started. For experiments monitoring a decrease in fluorescence, NH₄Cl was added to a final concentration of 50 mM or bafilomycin to 1 μM and imaging continued. For experiments where cells were pre-incubated with Aβ42 or vAβ42, Ova-Cy was added to a final concentration of 25 μg/mL. For all CypHer analyses, where one 0.5 μM z-slice has been analysed, a focal plane in the middle of the cell that contained none or a negligible proportion of saturating pixels within the cell body was selected from the image that had maximal fluorescence. Focal planes in images at other time points were matched to this slice to ensure the same focal plane of the cell was being monitored and mean fluorescence intensity calculated. A region of interest was drawn around the cell body excluding the nucleus. The mean fluorescence intensity in either one z-slice or a maximum projection image of several slices at each time point was measured in each cell body. All cells within a given FOV were analysed. Where images are displayed, LUTs were matched for each image within an experiment. All analysis was carried out using FIJI. Graphs and statistical analysis were prepared in GraphPad Prism.

**Fluorimetry**

19 mM Ova-Cy stock solution was diluted tenfold into EBS that had a pH of either 5.4 or 7.4 when it contained 10 mM NH₄Cl. Scans were collected on a Varian Cary Eclipse fluorimeter. Samples were excited at either 500 nm or 644 nm and emission collected from 650 to 800 nm. Excitation and emission slit widths were set to 5 nm, and the scan rate was 600 nm/min with 1 nm data intervals and an averaging time of 0.1 s.

**Significance statement**

The misfolding and aggregation of proteins is intimately linked with many neurodegenerative diseases. This manuscript reports the use of a new optical probe to establish how prefibrillar, oligomeric forms of the Alzheimer’s associated peptide, Aβ42, bring about damaging effects in primary neurons. We demonstrate that CypHer-Aβ42 oligomers are rapidly taken up into cells in a dynamin-dependent manner and trafficked via the endosomal–lysosomal pathway resulting in accumulation in lysosomes. Preincubation with Aβ42 influences the subsequent uptake of other proteins, indicative of impairments in endo-lysosomal mechanisms. Here we dynamically monitor the cellular response to the presence of Aβ42 oligomers in primary neurons and establish a new link between Aβ42 aggregation, uptake and deficits in the endosomal–lysosomal pathway.

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**Author contributions**

KM and DV conducted the toxicity and uptake experiments. KM conducted remaining experiments, managed the work and wrote the manuscript. KM, LS and KS planned the experiments. LS and KS directed the work and edited the manuscript and figures.

**Data availability**

The datasets supporting the conclusions of this article are included within the article and its supporting information.

**Compliance with ethical standards**

**Conflict of interest**

The authors declare no competing financial interests.

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