

Elucidating the Oligomerization and Cellular Interactions of a Trimer Derived from A β through Fluorescence and Mass Spectrometric Studies

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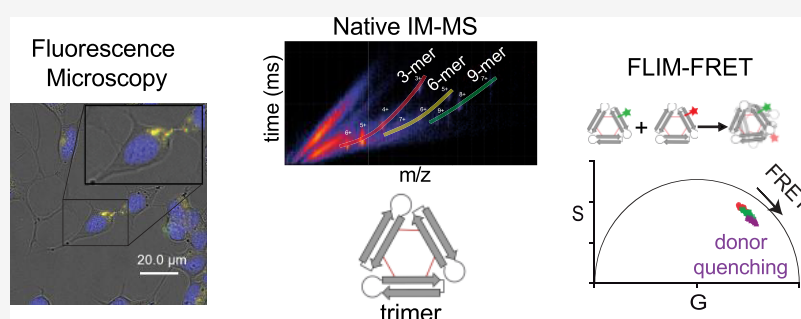
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ABSTRACT: A β oligomers play a central role in the neurodegeneration observed with Alzheimer's disease. Our laboratory has developed covalently stabilized trimers derived from residues 17–36 of A β as model systems for studying A β oligomers. In the current study, we apply the emerging techniques of fluorescence lifetime imaging microscopy (FLIM) and native mass spectrometry (native MS) to better understand the assembly and interactions of the oligomer model system 2AT-L in aqueous solutions and with cells. 2AT-L and fluorescently labeled 2AT-L analogues assemble in the membrane-like environment of SDS-PAGE, showing diffuse bands of oligomers in equilibrium. Native ion mobility-mass spectrometry (native IM-MS) of 2AT-L allows for the identification of discrete oligomers in solution and shows similar patterns of oligomer formation between 2AT-L and fluorescently labeled analogues. Fluorescence microscopy with SH-SY5Y cells reveals that fluorescently labeled 2AT-L analogues colocalize within lysosomes. FLIM studies with phasor analysis further elucidate the assembly of 2AT-L within cells and establish the occurrence of FRET, indicating the presence of oligomers within cells. Collectively, these multiple complementary techniques help better understand the complex behavior of the 2AT-L model system.

KEYWORDS: Alzheimer's disease, amyloid β , oligomer, FRET, ion mobility-mass spectrometry, fluorescence lifetime imaging microscopy

INTRODUCTION

Elucidating the biological properties of A β oligomers and understanding their solution-phase behavior are essential to better understanding Alzheimer's disease.¹ Accumulation of the β -amyloid peptide (A β) in the brain is a hallmark of Alzheimer's disease and is a key contributor to neurodegeneration.² A β aggregates to form toxic oligomers and the fibrils that make up the characteristic plaques observed in the brains from those with Alzheimer's disease.³ These oligomers are thought to be the main synaptotoxic species but have proven difficult to study as they are inherently heterogeneous and metastable with a high propensity to form fibrils.^{4–9} Trimers of A β_{42} are among the most toxic oligomers and are implicated in neuronal cell death.^{7,10–13}

Model systems of stabilized A β -derived oligomers have emerged as tools to better understand endogenous oligomers and provide further insight into the molecular basis of

Alzheimer's disease.^{13–15} Our laboratory has developed a series of covalently stabilized trimers derived from residues 17–36 of A β as chemical models of toxic amyloid oligomers associated with neurodegeneration in Alzheimer's disease.^{16–21} One of the trimers, termed 4AT-L, is composed of three β -hairpins formed by an A $\beta_{17–36}$ peptide, with molecular templates designed to induce β -hairpin folding, block uncontrolled aggregation, and allow disulfide crosslinking of three β -hairpins to form a covalent trimer.²¹ Trimer 4AT-L is toxic toward the neuronal cell line SH-SY5Y, assembles to form ball-shaped dodecamers composed of four trimers in the crystal state, and forms dodecamers in SDS-PAGE.

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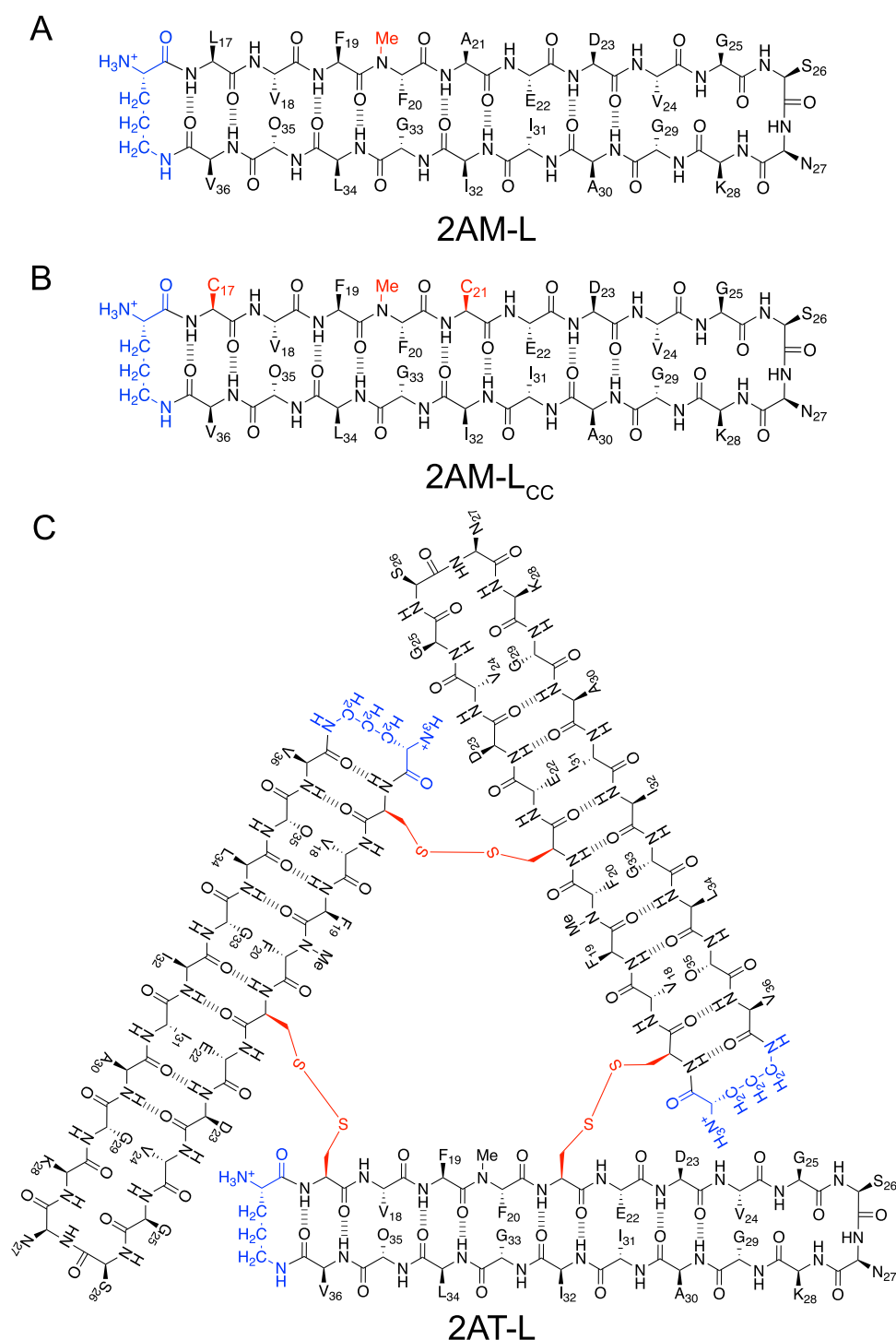


Figure 1. Chemical structures of β -hairpin peptides and a trimer derived from $A\beta_{17-36}$. (A) Macrocytic β -hairpin 2AM-L incorporating an N-methyl residue to prevent uncontrolled aggregation and a δ Orn turn (blue) mimic to constrain the N- and C-termini. (B) β -Hairpin 2AM-L_{CC} incorporates cysteine mutations at residues 17 and 21. (C) Three 2AM-L_{CC} β -hairpins form covalently stabilized trimer 2AT-L through disulfide bonds.

Fluorescence lifetime imaging microscopy (FLIM) and native mass spectrometry (native MS) have emerged as important new tools to probe the assembly and interactions of peptides and proteins.^{22–26} Native MS provides information about the stoichiometry of oligomeric assemblies present in solution and can thus complement the structural information elucidated from X-ray crystallography and SDS-PAGE to build a broader understanding of the solution-phase assembly of $A\beta$ -

derived oligomers.^{27,28} Fluorescence microscopy to visualize $A\beta$ -derived oligomers in the presence of cells and the application of FLIM to determine the occurrence of Förster resonance energy transfer (FRET) can complement the previously observed cellular cytotoxicity.^{29–35} In the current study, we set out to apply these and other techniques to better understand the assembly and interactions of a covalently

stabilized trimer derived from residues 17–36 of A β in aqueous solutions and with cells.

RESULTS AND DISCUSSION

Design and Synthesis of 2AT-L. In trimer 4AT-L, the native phenylalanine at position 20 is replaced with cyclohexylalanine. For the current study, we prepared homologue 2AT-L, which behaves similarly but has the native phenylalanine at position 20. Trimer 2AT-L is composed of three crosslinked A β _{17–36} β -hairpins. In each β -hairpin, a δ -linked ornithine (δ Orn) turn unit between residues 17 and 36 helps enforce a folded β -hairpin conformation, and an *N*-methyl group on phenylalanine 20 helps block uncontrolled aggregation (Figure 1A). In 2AT-L, residues 17 and 21 are replaced by cysteines, which provide covalent crosslinks that hold the trimer together by connecting the monomer subunits at the vertices (Figure 1B,C). The crosslinked trimers are homogeneous, stable, and mimic some of the biological properties of A β oligomers.^{19,21}

We synthesized trimer 2AT-L by the same procedures that we previously used to prepare trimer 4AT-L. We first prepared macrocyclic peptide 2AM-L_{CC} by solid-phase peptide synthesis of the corresponding linear peptide on 2-chlorotrityl resin, followed by cleavage of the protected peptide from the resin, solution-phase macrocyclization, deprotection, and purification by RP-HPLC. 2AM-L_{CC} was oxidized at 6 mM in 20% aqueous DMSO with 60 mM triethylamine for 48 h. 2AT-L was isolated from the oxidation reaction by RP-HPLC. Pure fractions were lyophilized, affording >95% pure 2AT-L as the trifluoroacetate (TFA) salt.

Preparation of Fluorescently Labeled 2AT-L. Substoichiometric labeling of 2AT-L with fluorophore NHS esters permits the isolation of singly labeled 2AT-L.³⁶ We developed a labeling procedure in which 2AT-L is treated with 0.05 molar equivalents of the commercially available sulfo-cyanine3 (sCy3) or sulfo-cyanine5 (sCy5) NHS esters, and the singly labeled trimer is isolated by RP-HPLC, and the unlabeled trimer is recycled. We found that heating the HPLC column greatly facilitates the separation of the labeled trimer from the unlabeled trimer and also permits the removal of the small amounts of doubly labeled trimer that form. We selected the bis-sulfonated analogues of the commonly used FRET partners Cy3 and Cy5 as fluorophores because they provide enhanced aqueous solubility and reduced aggregation. Labeling 3 mg of trimer by this procedure typically permits isolation of ca. 150 μ g of singly labeled trimer 2AT-L-sCy3 or 2AT-L-sCy5 as the TFA salt, which was sufficient for many of the experiments described below.

Oligomerization of 2AT-L and Fluorescently Labeled 2AT-L Analogues by SDS-PAGE. We used SDS-PAGE to initially assess the effect of the sCy3 and sCy5 fluorophores on the assembly of 2AT-L into higher-order oligomers. In SDS-PAGE, trimer 2AT-L assembles to form higher-order oligomers. When 2AT-L (6.6 kDa) is run in SDS-PAGE and visualized by silver staining, it forms a downward-streaking band from ca. 32 kDa to ca. 13 kDa, suggesting assembly into oligomers ca. 2–5 trimers in size (Figure 2A). In contrast, the monomer 2AM-L (2.2 kDa) migrates at ca. 3 kDa, indicating the absence of assembly into oligomers. 2AT-L-sCy3 (7.2 kDa) migrates as a compact band at ca. 14 kDa, suggesting hexamer formation (2 trimers), while 2AT-L-sCy5 (7.2 kDa) migrates as a more diffuse band, from ca. 21 kDa to ca. 13 kDa,

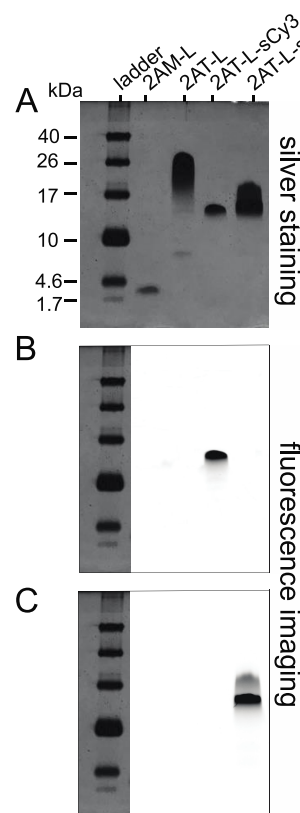


Figure 2. SDS-PAGE of peptides 2AM-L, 2AT-L, 2AT-L-sCy3, and 2AT-L-sCy5. (A) Silver-stained image. (B) Fluorescence image in the Cy3 channel. (C) Fluorescence image in the Cy5 channel. SDS-PAGE was performed in Tris buffer at pH 6.8 with 2% (w/v) SDS on a 16% polyacrylamide gel with 50 μ M solutions of peptide in each lane. Fluorescence imaging was performed before silver staining.

suggesting the formation of hexamers or perhaps hexamers and nonamers (2–3 trimers).

Fluorescence imaging provides additional insights into the bands formed by the trimers (Figure 2B,C). Notably, the diffuse band formed by 2AT-L-sCy5 shows an intense component at ca. 14 kDa and a weaker component at ca. 21 kDa. The difference in shape between the bands formed by 2AT-L-sCy5 and 2AT-L-sCy3 may reflect the greater hydrophobicity and flat hydrophobic surface area provided by the larger sCy5 fluorophore. Both the sCy3 and sCy5 labels appear to impede the assembly, with hexamers as the main species observed, while the unlabeled 2AT-L forms predominantly larger oligomers.³⁷

Oligomerization of 2AT-L and Fluorescently Labeled 2AT-L Analogues by Native IM-MS. We used native MS coupled with ion mobility spectrometry (IM) to investigate the oligomeric assembly of the trimers to complement the SDS-PAGE studies and avoid SDS-induced oligomer formation.³⁸ In native IM-MS, oligomers and noncovalent assemblies are ionized and separated without dissociation.³⁹ IM separates ionized oligomers of different sizes, shapes, and charges in the gas phase as they travel through the ion mobility device at different rates, and their arrival times are measured.⁴⁰ The ions then are analyzed by a mass analyzer to determine their mass-to-charge ratio, m/z (Figure 3A). Trimer 2AT-L and fluorescently labeled 2AT-L analogues show a charge state distribution in the mass spectrum, leading to many overlapping m/z peaks that arise from multiple assemblies of trimers. The

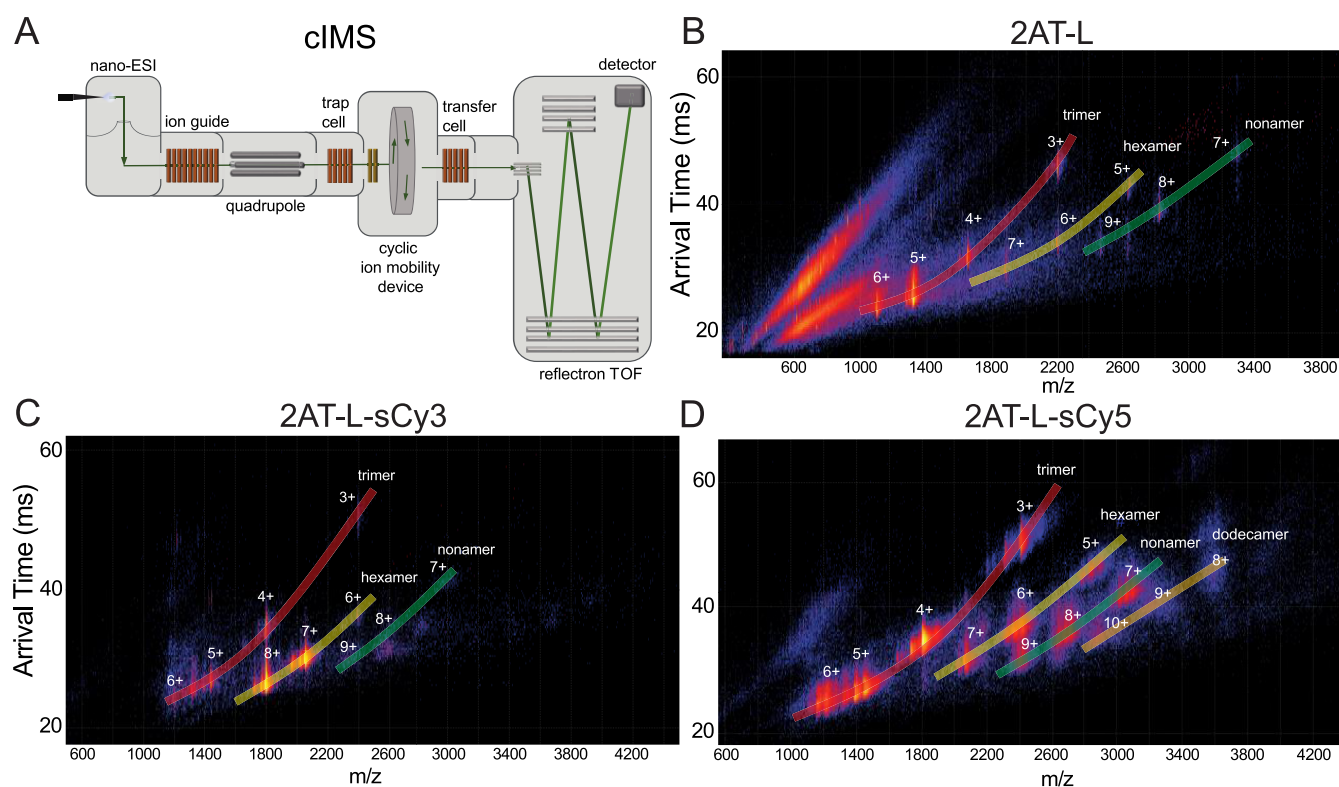


Figure 3. Native IM-MS of 2AT-L and labeled analogues. (A) Diagram of the cIMS instrument used to run native IM-MS experiments. Molecules are ionized through nano-electrospray ionization (nano-ESI) and then travel through an ion guide and quadrupole. Ions reach a trap cell before entering the cyclic ion mobility device in which species are separated by size, shape, and charge. Ions exit the mobility device and proceed through a reflectron time-of-flight (TOF) mass analyzer to separate ions by mass-to-charge ratio. The time for an ion to travel through the mobility cell to detection is recorded as the arrival time. (B–D) Mobiligrams of 2AT-L, 2AT-L-sCy3, and 2AT-L-sCy5. Native IM-MS was performed on a SELECT SERIES Cyclic IMS Q-cIMS-TOF (cIMS) instrument (Waters Corporation). Numeric labels indicate the charge state of the corresponding oligomeric species. Samples of 2AT-L and fluorescently labeled 2AT-L analogues were prepared at a concentration of 40 μ M in 400 mM ammonium acetate.

separation by arrival times that occurs in the ion mobility device permits the separation of species with identical mass-to-charge ratios into their component oligomers.

Native IM-MS reveals that 2AT-L and fluorescently labeled 2AT-L analogues also assemble to form oligomers in aqueous solution. Each IM-MS experiment generates a mobiligram, in which the arrival time is plotted against mass-to-charge ratio, with the relative intensity displayed as a heat map. The IM-MS mobiligram for trimer 2AT-L shows that 2AT-L forms hexamers and nonamers, in addition to the trimer (Figure 3B). IM-MS shows that 2AT-L-sCy3 and 2AT-L-sCy5 also undergo assembly. The mobiligram for 2AT-L-sCy3 shows the formation of hexamers, as well as a low abundance of nonamers and dodecamers, in addition to the trimer (Figure 3C). The mobiligram for 2AT-L-sCy5 also shows the formation of hexamers, nonamers, and dodecamers, in addition to the trimer (Figure 3D).⁴¹

Together, the SDS-PAGE and native IM-MS experiments reveal that oligomers of 2AT-L and the fluorescently labeled 2AT-L analogues can form in either the presence or absence of SDS. Although the fluorophore labels appear to perturb the formation of higher-order oligomers in SDS-PAGE, a similar perturbation is not seen in native IM-MS.

Co-Oligomerization of 2AT-L-sCy3 and 2AT-L-sCy5 by FRET. We performed steady-state FRET experiments with mixtures of 2AT-L-sCy3 (donor) and 2AT-L-sCy5 (acceptor) to further assess the assembly of 2AT-L into higher-order

oligomers in solution. The occurrence of FRET can be determined by selectively exciting at a wavelength absorbed by the donor and observing fluorescence at a wavelength emitted by the acceptor. In practice, it is often difficult to selectively excite only the donor. Thus, we used an excitation wavelength of 490 nm—which is well below the 548 nm λ_{max} of sCy3—to minimize direct excitation of sCy5.⁴² We monitored fluorescence emission at 662 nm—the emission maximum of sCy5—and varied the ratio of 2AT-L-sCy3 and 2AT-L-sCy5 from 100:0 to 0:100 while maintaining a total concentration of 5 μ M.

When 2AT-L-sCy3 alone (100:0) is irradiated at 490 nm, emission (0.57 rfu) occurs at 662 nm (Figure 4). This emission results from the sCy3 fluorophore, which has an emission maximum of 556 nm, but which extends weakly to 662 nm and beyond. When a 50:50 mixture of 2AT-L-sCy3 and 2AT-L-sCy5 is irradiated, the emission at 662 nm increases (1.00 rfu). The enhanced fluorescence reflects the occurrence of FRET and thus demonstrates the co-oligomerization of 2AT-L-sCy3 and 2AT-L-sCy5. Only modest emission (0.29 rfu) occurs from excitation of 2AT-L-sCy5 alone (0:100) at 490 nm, providing further evidence that the enhanced fluorescence in the 50:50 mixture results from FRET. Additional mixtures of 2AT-L-sCy3 and 2AT-L-sCy5 (80:20, 60:40, 40:60, 20:80) show intermediate levels of emission, providing further evidence for co-oligomerization and FRET.

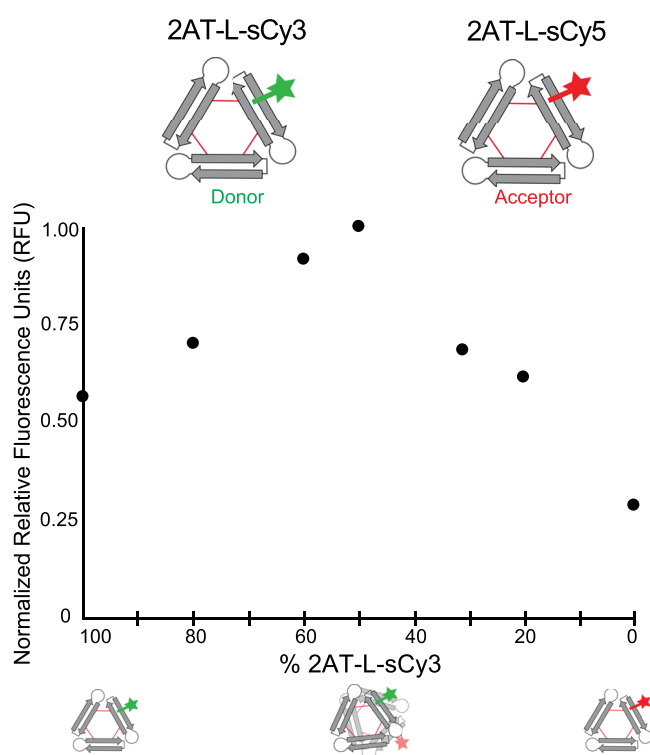


Figure 4. Fluorescence at the emission maximum of 2AT-L-sCy5 as a function of the percentage of 2AT-L-sCy3. Fluorescence spectra of mixtures of varying ratios of 2AT-L-sCy3 and 2AT-L-sCy5 were acquired with excitation at 490 nm and observation at 662 nm. All spectra were collected on aqueous solutions of 2AT-L-sCy3 and 2AT-L-sCy5 with a total concentration of 5 μ M.

Fluorescence Microscopy of 2AT-L-sCy3 and 2AT-L-sCy5. To gain insights into the biological roles of the oligomers observed by SDS-PAGE, native IM-MS, and steady-state FRET experiments, we performed fluorescence microscopy with SH-SY5Y cells. Trimer 4AT-L is toxic toward SH-SY5Y cells, and trimer 2AT-L exhibits similar toxicity.²¹ To further explore the interaction of A β -derived trimers with mammalian cells, we used fluorescence microscopy. SH-SY5Y cells were incubated with fluorescent analogues of 2AT-L at a total concentration of 5 μ M. Cells were then counterstained with Hoechst 33342 nuclear stain (blue), washed with phenol red-free DMEM:F12 media, and imaged. Fluorescence micrographs were collected with emissions observed in the respective Cyanine3 and Cyanine5 channels.

Fluorescence microscopy reveals that both 2AT-L-sCy3 and 2AT-L-sCy5 are internalized by cells. Incubation of SH-SY5Y cells with 2AT-L-sCy3 resulted in defined intracellular puncta and some small fluorescent features bound to the cell membrane (Figure S2). Incubation of SH-SY5Y cells with 2AT-L-sCy5 resulted in similar features (Figure S3). Incubation of SH-SY5Y cells with both 2AT-L-sCy3 and 2AT-L-sCy5 also resulted in similar features (Figure 5). Merged images, showing both the green (2AT-L-sCy3) and red (2AT-L-sCy5) channels, show a yellow hue, indicating co-localization of the fluorescently labeled trimers. Punctate features were observed as early as 5 h after treatment and at concentrations of fluorescently labeled 2AT-L as low as 8 nM (Figure S4).

The punctate intracellular features observed for the labeled 2AT-L are similar to those reported for fluorescently labeled A β ₄₂, where uptake occurred primarily through endocytosis.^{43–50} These studies have shown that A β localizes in lysosomes. To assess whether 2AT-L also localizes into lysosomes, we performed further experiments using the lysosomal marker LysoTracker Green. Treatment of SH-SY5Y cells with LysoTracker Green and either 2AT-L-sCy3 or 2AT-L-sCy5 showed co-localization, indicating sequestration in the lysosomes (Figures S5 and 6). Thus, it appears that the uptake of 2AT-L occurs through endocytosis, in a fashion similar to that of A β .

Previous reports have suggested that oligomerization is a prerequisite for the cellular uptake of A β .⁵¹ The covalently stabilized oligomer model 2AT-L is readily taken up by cells and may thus serve as a model to further study the cellular uptake of A β oligomers. To continue to probe the assembly state of 2AT-L-sCy3 and 2AT-L-sCy5 and assess for higher-order oligomer formation, we turned to FRET microscopy.

FLIM-FRET Studies of 2AT-L-sCy3 and 2AT-L-sCy5. We used fluorescence lifetime imaging microscopy (FLIM) to further investigate the intracellular puncta observed when fluorescently labeled 2AT-L analogues are incubated with cells. FLIM allows the detection of FRET and can thus reveal molecular co-localization at a resolution higher than can be achieved through confocal microscopy. If 2AT-L molecules bearing FRET partners co-assemble or are otherwise in close proximity (<10 nm), FRET can occur and will manifest as a reduction in the fluorescence lifetime of the FRET donor through fluorescence quenching.⁵² Measurement of fluorescence lifetimes through FLIM-FRET offers advantages over intensity-based FRET measurements because the measurement of the lifetime is generally independent of the concentration of

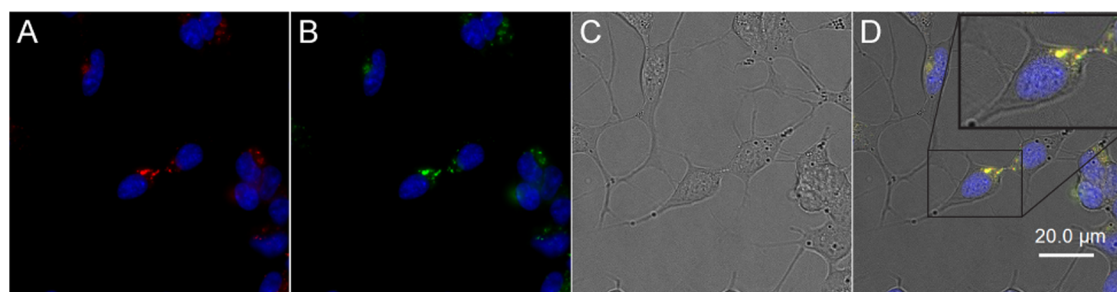


Figure 5. Micrographs illustrating intracellular co-localization of fluorescently labeled 2AT-L analogues in SH-SY5Y cells. (A) 2AT-L-sCy5 (red) and Hoechst 33342 nuclear stain (blue). (B) 2AT-L-sCy3 (green) and Hoechst 33342 nuclear stain (blue). (C) Brightfield image. (D) Merged fluorescent images with brightfield images. Cells were incubated with 5 μ M 2AT-L-sCy3 and 5 μ M 2AT-L-sCy5 for 6 h at 37°C, counterstained with Hoechst 33342 nuclear stain, and imaged.

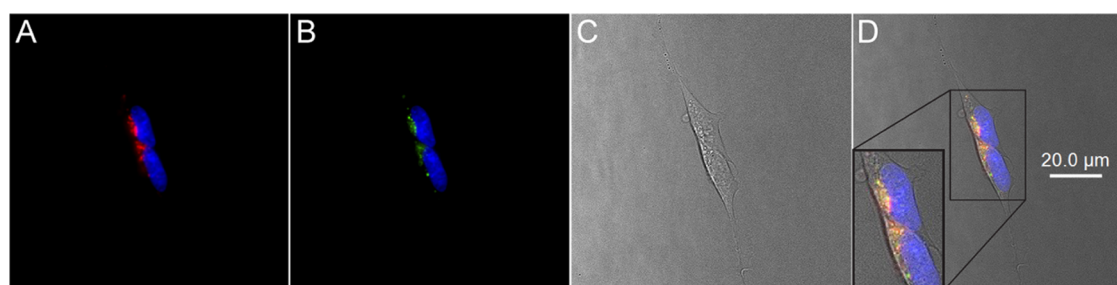


Figure 6. Micrographs illustrating intracellular co-localization of 2AT-L-sCy5 and Lysotracker Green in SH-SY5Y cells. (A) 2AT-L-sCy5 (red) and Hoechst 33342 nuclear stain (blue). (B) Lysotracker Green (green) and Hoechst 33342 nuclear stain (blue). (C) Brightfield image. (D) Merged fluorescent images with brightfield images. Cells were incubated with 5 μ M 2AT-L-sCy5 and 100 nM Lysotracker Green for 6 h at 37°C, counterstained with Hoechst 33342 nuclear stain, and imaged.

the fluorophores and does not require correction for spectral crosstalk.⁵³

To assess FRET inside cells, we treated SH-SY5Y cells with either 2AT-L-sCy3 (donor) alone or with mixtures of 2AT-L-sCy3 (acceptor) and 2AT-L-sCy5 and looked for decreases in the fluorescence lifetime of sCy3 with increasing mole fraction of sCy5. Although 2AT-L-sCy3 oligomers or 2AT-L-sCy5 oligomers can form, only oligomers containing both 2AT-L-sCy3 and 2AT-L-sCy5 are expected to show decreased fluorescence lifetime (Figure 7A).⁵⁴ We used the phasor approach to analyze changes in the fluorescence lifetime of sCy3. In this method, each pixel of a FLIM image is transformed to a point on a phasor plot, allowing for simple visualization of changes in the donor lifetime in a micrograph and thus facilitating the assessment of FRET.^{55–58} Changes in fluorescence lifetime with increases in mole fraction of sCy5 manifest as changes in the positions of points on the phasor plot (Figure 7B), with greater changes corresponding to a greater fraction of molecules undergoing FRET.^{55,59,60}

We performed three sets of experiments, in which SH-SY5Y cells were treated with 2AT-L-sCy3 and 2AT-L-sCy5 in 100:0, 50:50, and 25:75 ratios while maintaining a total concentration of 5 μ M. For each ratio, we collected 25–26 images of individual cells. We observed the photon intensity in the donor channel decrease as the fraction of 2AT-L-sCy5 increased (Figure 7C). The fluorescence decay from each pixel was determined, and the average from each image was plotted onto a phasor plot, with different colors (red, green, and violet) representing the three sets of ratios studied (Figure 7D,E). The resulting phasor plot shows significant decreases in fluorescence lifetime with an increasing fraction of sCy5 and thus provides compelling evidence for FRET and hence intimate proximity of 2AT-L-sCy3 and 2AT-L-sCy5 in the cells.

The fluorescence lifetimes measured in FLIM-FRET can also be represented as a heat map, with each pixel of an image colored to reflect its relative fluorescence lifetime. We have thus represented the distribution of fluorescence lifetimes on three representative images at the 100:0, 50:50, and 25:75 ratios, with red representing the longest lifetimes and violet representing the shortest lifetimes, and colors ranging from orange to blue representing intermediate lifetimes (Figure 7F). The shift from red toward violet across the series of images reflects the decrease in fluorescence lifetime with an increasing fraction of 2AT-L-sCy5 and further demonstrates that FRET increases with the addition of sCy5. Collectively, these FLIM-FRET experiments provide further evidence that 2AT-L-sCy3 and 2AT-L-sCy5 co-oligomerize in cells.

SUMMARY AND CONCLUSIONS

Our covalently stabilized A β -derived trimers mimic some of the assembly and biological properties of A β oligomers. In the crystal state, 2AT-L and homologue 4AT-L assemble to form ball-shaped dodecamers. In the membrane-like environment of SDS-PAGE, 2AT-L dodecamers are also observed but appear to be in equilibrium with oligomers ca. 2–5 trimers in size. Labeling 2AT-L with sCy3 and sCy5 appears to attenuate assembly in SDS-PAGE, with 2AT-L-sCy3 and 2AT-L-sCy5 migrating mainly as hexamers. Native IM-MS of 2AT-L identifies discrete hexamers and nonamers in solution. Native IM-MS of 2AT-L-sCy3 and 2AT-L-sCy5 shows similar oligomer formation to 2AT-L.⁶¹

Steady-state FRET experiments also show that 2AT-L-sCy3 and 2AT-L-sCy5 form oligomers in solution. Fluorescence microscopy shows that 2AT-L-sCy3 and 2AT-L-sCy5 form intracellular puncta in SH-SY5Y cells, which co-localize with a lysosomal marker, suggesting an uptake mechanism through endocytotic vesicles that is consistent with previous studies of A β .⁴⁹ FLIM studies and phasor analysis establish the occurrence of FRET within cells treated with both 2AT-L-sCy3 and 2AT-L-sCy5, suggesting oligomerization within cellular vesicles.

The aggregation of A β to form oligomers is difficult to study in detail due to the complexity of the different species that form and the propensity of A β to ultimately form insoluble fibrils. The constraint of an A β -derived peptide into a stable trimer, 2AT-L, provides a model system that is easier to study and can be explored by a variety of biophysical and biological techniques. While X-ray crystallography provides a simple answer that assemblies of dodecamers form in the crystal state, the behavior in the solution state is more complex. SDS-PAGE shows multiple oligomeric states in equilibrium but does not allow the identification of discrete oligomers. Native IM-MS, on the other hand, allows the identification of individual oligomers. The addition of fluorescent labels is required to visualize the interactions of 2AT-L in cells, although SDS-PAGE shows that these labels can alter the oligomerization, at least somewhat. FLIM-FRET experiments with phasor analysis allow the observation of oligomerization of the labeled 2AT-L in cells. Collectively, the application of multiple complementary techniques helps better elucidate the complex behavior of the 2AT-L model system. We anticipate that these techniques may also be useful for understanding the yet more complex oligomerization of A β .

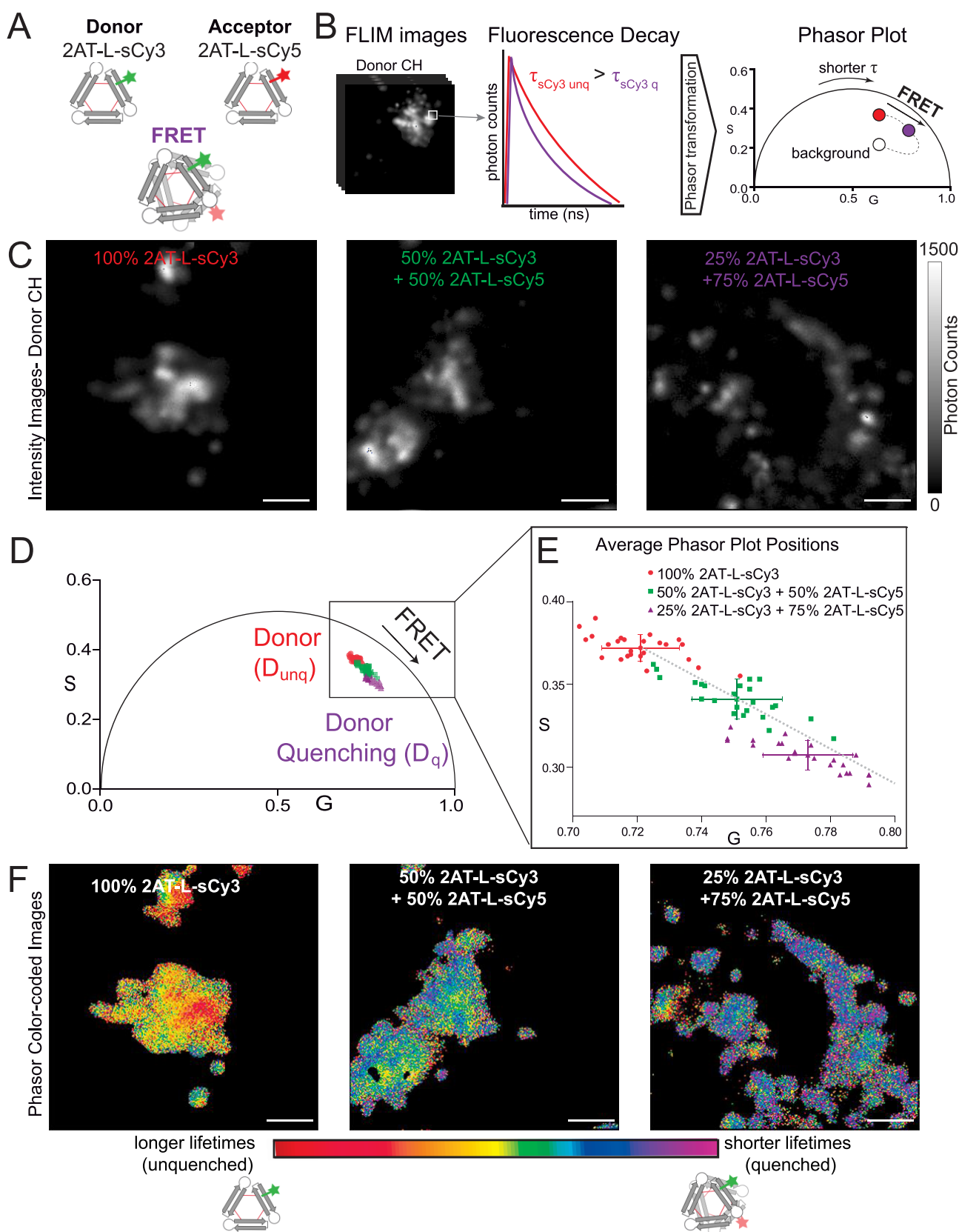


Figure 7. Phasor analysis of FLIM micrographs of SH-SY5Y cells treated with 2AT-L-sCy3 and 2AT-L-sCy5. Oligomeric assemblies of 2AT-L-sCy3 and 2AT-L-sCy5 within cells were analyzed by FLIM-FRET and the phasor approach. (A) Representation of the co-assembly of 2AT-L-sCy3 (donor) and 2AT-L-sCy5 (acceptor) and the occurrence of FRET. (B) Representation of the phasor transformation of FLIM images. Fluorescence lifetime of each pixel of each image is analyzed and plotted on a phasor plot. When FRET occurs, the excited state lifetime of the donor decreases and the phasor position shifts toward shorter lifetimes. The curved FRET trajectory (dotted line) follows the classical definition of FRET efficiency and is influenced by the background fluorescence lifetime. (C) Representative images showing photon intensity of SH-SY5Y cells treated with 2AT-

Figure 7. continued

L-sCy3 and 2AT-L-sCy5 in 100:0, 50:50, and 25:75 ratios. Images were acquired in the donor (sCy3) channel. Scale bar = 5 μm . (D) Phasor plot illustrating the average phasor position in each image. Images collected for the 100:0 ratio are represented by red points and correspond to unquenched donor (D_{unq}). Images collected for the 50:50 ratio are represented by green points. Images collected for the 25:75 ratio are represented by violet points and correspond to increased donor quenching (D_q). (E) Details of phasor plot in panel (D). Average phasor plot positions ($\pm\text{SD}$) from each image. (F) Representative images of SH-SY5Y cells with false color illustrating fluorescence lifetime. Longer fluorescence lifetimes are represented in red and orange, while shorter fluorescence lifetimes reflecting increased FRET are represented progressively in yellow, green, blue, and violet.

MATERIALS AND METHODS

Peptides 2AM-L, 2AM-L_{CC}, and 2AT-L were synthesized by procedures analogous to those described previously.¹⁹ SDS-PAGE and silver staining were performed as described previously.²¹ Procedures detailing the preparation of 2AT-L-sCy3 and 2AT-L-sCy5, native mass spectrometry studies, steady-state FRET experiments, fluorescence microscopy, and FLIM-FRET can be found in the Supporting Information.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at <https://pubs.acs.org/doi/10.1021/acschemneuro.2c00313>.

Procedures detailing the preparation of 2AT-L-sCy3 and 2AT-L-sCy5, native mass spectrometry studies, steady-state FRET experiments, fluorescence microscopy, and FLIM-FRET (PDF)

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Author Contributions

G.G. and J.S.N. designed the research; G.G. synthesized the compounds, performed SDS-PAGE, steady-state FRET, cell culture, and fluorescence microscopy; M.W. synthesized the compounds; G.G. and B.T. performed FLIM experiments; Y-F. L. performed native IM-MS experiments; and G.G. and J.S.N. wrote the paper.

Notes

The authors declare no competing financial interest.

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