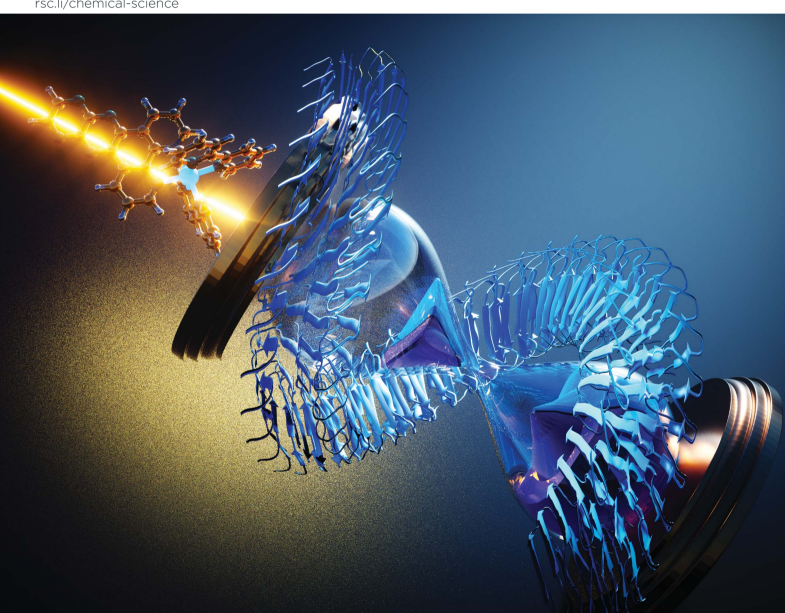
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Deconvoluting binding sites in amyloid nanofibrils using time-resolved spectroscopy†

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Steady-state fluorescence spectroscopy has a central role not only for sensing applications, but also in biophysics and imaging. Light switching probes, such as ruthenium dipyridophenazine complexes, have been used to study complex systems such as DNA, RNA, and amyloid fibrils. Nonetheless, steady-state spectroscopy is limited in the kind of information it can provide. In this paper, we use time-resolved spectroscopy for studying binding interactions between amyloid-β fibrillar structures and photoluminescent ligands. Using time-resolved spectroscopy, we demonstrate that ruthenium complexes with a pyrazino phenanthroline derivative can bind to two distinct binding sites on the surface of fibrillar amyloid-β, in contrast with previous studies using steady-state photoluminescence spectroscopy, which only identified one binding site for similar compounds. The second elusive binding site is revealed when deconvoluting the signals from the time-resolved decay traces, allowing the determination of dissociation constants of 3 and 2.2 µM. Molecular dynamic simulations agree with two binding sites on the surface of amyloid-β fibrils. Time-resolved spectroscopy was also used to monitor the aggregation of amyloid- β in real-time. In addition, we show that common polypyridine complexes can bind to amyloid-\$\beta\$ also at two different binding sites. Information on how molecules bind to amyloid proteins is important to understand their toxicity and to design potential drugs that bind and guench their deleterious effects. The additional information contained in time-resolved spectroscopy provides a powerful tool not only for studying excited state dynamics but also for sensing and revealing important information about the system including hidden binding sites.

Introduction

Alzheimer's disease (AD) is a degenerative disease that affects the human brain, particularly in elderly people, with symptoms that include memory loss, confusion, and personality changes. The amyloid cascade hypothesis proposes the accumulation of amyloid- β (A β) aggregates as a mechanism in the progression of Alzheimer's disease. While the relation between amyloid aggregates (soluble and insoluble) and the onset of AD has been known for more than 100 years, it was only last year when the first amyloid-targeting drug (an antibody called Aducanumab⁴) was approved by the FDA, with many other medications of the

same kind on the horizon.⁵ This has prompted a resurgence of interest in small molecule-based treatments targeting amyloids. Understanding how small molecules bind and interact with amyloid aggregates is, therefore, essential to developing appropriate treatments for AD. The study of binding sites in amyloid aggregates has greatly benefited from the use of photoluminescence titration experiments. This method provides the dissociation constants of ligands by simply measuring the steady-state photoluminescence intensities over a series of samples. However, standard steady-state titration methodologies have the drawback that they assume the target molecule has a single unique binding site. Therefore, the presence of multiple binding sites needs to be confirmed through other analytical methods.⁶

A different method to study the photoluminescence of a molecular probe is time-resolved spectroscopy. Lincoln *et al.* reported that they observed two different lifetimes while studying the emission of ruthenium(II) complex bound to DNA, which were yielded by two different geometries of binding to DNA.^{7–9} More relevant to amyloid proteins, fluorescence lifetime imaging microscopy (FLIM) has been used to image and study amyloid fibrils based on changes in lifetime.¹⁰ These studies

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have used lifetime in qualitative ways, not fully capitalizing on the reach of time-resolved spectroscopy. We have expanded the application of time-resolved spectroscopy to determine quantitative binding information on fibrillar amyloids. This also compensates for the shortfalls of steady-state photoluminescence titrations, particularly when photoluminescent metal complexes are used as probes.

We have previously reported the use of photoluminescent metal complexes of ruthenium11-15 and rhenium16,17 to study amyloid aggregates. 18,19 Particularly, we investigated the binding of light-switching ruthenium and rhenium dipyridophenazine complexes to Aβ fibrils using steady-state photoluminescence molecular titrations, dynamics (MD) simulations,11 and photochemical footprinting.16 These studies have provided important information on the binding equilibrium of small molecules to Aβ, but the values were obtained under the assumption that AB has only one binding site. In another work, we used a ruthenium polypyridyl complex $([Ru(bpy)_2(dpqp)]^{2+}, bpy = 2,2'-bipyridine, dpqp = pyrazino$ [2',3':5,6]pyrazino[2,3-f][1,10]phenanthroline) to monitor the formation of AB oligomers using photoluminescence anisotropy.15 This metal complex was selected because it lacks a strong light-switching response in the presence of Aβ aggregates,20 which enable us to observe the anisotropy change of the complex. A molecule without the light-switching response is impractical for determining the binding information using steady-state photoluminescence titration; however, it is ideal for monitoring all the species present in a system (e.g. bound to $A\beta$ fibrils and free in solution) using time-resolved spectroscopy.

Here, we study binding sites on A β using $[Ru(bpy)_2(dpqp)]^{2+}$ as a probe and time-resolved photoluminescence spectroscopy to evaluate the concentration of probes bound to Aβ. When performing titrations to elucidate the binding equilibrium, we found that two different binding sites are present, one that promotes a change in the lifetime of the ruthenium complex and the other that does not affect the lifetime. MD simulations were carried out to shed light on the nature of the interaction between $[Ru(bpy)_2(dpqp)]^{2+}$ and A β fibrils. By selectively deconvoluting the emission from [Ru(bpy)2(dpqp)]2+ bound to A β , we were able to monitor the aggregation of A β in real-time. The formation of AB fibrils was confirmed by atomic force microscopy (AFM). In addition, we demonstrated the binding of common ruthenium and iridium polypyridine complexes $([Ru(bpy)_3]^{2+}$ and $[Ir(ppy)_2(bpy)]^+$, ppy = 2-phenylpyridine) to two different binding sites on AB fibrils. Interestingly, both $[Ru(bpy)_3]^{2+}$ and $[Ir(ppy)_2(bpy)]^+$ showed binding to A β , even when they do not have traditional Aβ recognition (or binding) elements. Due to the relevance of AB aggregation on the development of AD, elucidating new ways to study Aβ aggregates, their binding sites, and the process of aggregation is of great importance.

Experimental section

Purification of Aβ₁₋₄₀

Synthetic $A\beta_{1-40}$ was purchased from 21st Century Biochemicals. For HPLC purification, desalted $A\beta_{1-40}$ was dissolved in

0.1% TFA in water by vortexing for 1 min. HPLC purification was performed using a reverse-phase C-18 peptide column with a gradient elution of water with 0.1% TFA-acetonitrile changing from 70:30 to 10:90 over 25 min with a flow rate of 1.5 mL min⁻¹ at 60 °C. Purified Aβ was lyophilized and stored at -20 °C.

Preparation of Aβ solutions

A basic stock solution of monomeric AB was prepared by dissolving one tube of purified Aβ in 200 µL of 26.5 mM NaOH solution. The solution was diluted to 400 µL, and the concentration was measured using a UV-Visible spectrometer (Shimadzu 2450) at 292 nm ($\varepsilon = 2132 \text{ M}^{-1} \text{ cm}^{-1}$). The stock A β solution was diluted to a final concentration of 100 µM by adding phosphate buffer (PB). The final concentration of PB was 25 mM in the Aβ solution, and the pH was set at 7.4. For fibrils formation, the solution was incubated at 37 °C, with orbital stirring at 600 rpm. To quantify the amount of non-fibrillar species left after incubation, the incubated mixture was passed through a 10 kDa filter (Amicon Ultra-0.5 Centrifugal Filter Devices) at 14 000 g for 15 minutes. The absorbance of the filtered solution was compared to the absorbance of a control sample of a non-incubated buffered solution of AB, both measured at 280 nm. The non-fibril components of the final incubated solution was determined to be ca. 6% of the starting concentration of AB monomers. Therefore, all calculations of Aβ fibril concentrations were adjusted by a correction factor of 94%.

Synthesis of [Ru(bpy)2(dpqp)]2+

The synthesis of this compound is reported elsewhere. 20 Briefly, cis-Ru(bpy)2Cl2 and the dpqp ligand were refluxed for 4 h in a mixture of 1:1 ethanol/water. After the mixture reached room temperature, the volume of the mixture was halved, and then a saturated aqueous KPF₆ solution was added to precipitate the PF₆ complex. The purification of this compound was performed by successive precipitations from acetone/KPF₆ (aq), where the aqueous soluble phases were extracted with dichloromethane (3) imes 10 mL). The solution was evaporated to dryness, and the solid was washed with diethyl ether, yielding a red solid attributed to $[Ru(bpy)_2(dpqp)](PF_6)_2$ (73% yield). To make the complex soluble in water, tetrabutylammonium chloride was mixed with the PF₆ complex in acetone, converting the product to the chloride salt.

Binding assay (saturation titration experiments)

A constant 4 µM concentration of the metal complex was mixed with different concentrations of A β fibrils (2, 5, 15, 20, 30, 40, 60, and 80 µM) to prepare a series of solutions. Samples were excited with a picosecond pulse diode laser (200 kHz) at 370 nm. The emission was collected at 620 nm using an Edinburgh Instruments OD470 single-photon counting spectrometer and a high-speed red detector. A long pass filter (570 nm) was added to avoid scattering. Inner filter effect was not corrected since AB does not absorb at 370 nm, and it is known that correction is not necessary for moderate levels of scattering.21 The lifetimes

and preexponential factors for the different samples were determined using global fitting of all the decay curves.

Real-time aggregation assays using lifetime

A β aliquots were taken at different incubation times and mixed with $[Ru(bpy)_2(dpqp)]^{2^+}$ in aqueous solution to a final concentration of 50 μ M and 3 μ M, respectively. Time-resolved decays were collected for 1200 s for all samples to make the total photoluminescence intensity consistent between samples. The lifetime was fitted with a bi-exponential decay where τ_1 was fixed to be the lifetime of free complexes. In addition, 20 μ M ThT was added to each sample after lifetime measurements, allowing the detection of A β fibrils.

Computational simulations

The initial structure of [Ru(bpy)₂(dpqp)]²⁺ was modeled using the [Ru(bpy)₂(dppz)]²⁺ - DNA co-crystal structure (PDB ID: 4E1U).22 The structure was optimized at the B3LYP23/LANLTZ (+)24 level of theory utilizing the Hay-Wadt effective core potential²⁵ for Ru²⁺ using the Gaussian 09 program²⁶. The rest of the atoms (C, N, H) were treated with the 6-31 g(d) basis set. Additionally, dispersion effects were included utilizing the Grimme's function with the Becke-Johnson damping effect (GD3BJ).²⁷ The initial [Ru(bpy)₂(dpqp)]²⁺ - Aβ fibril complex was built through molecular-docking procedures on the two-fold Aβ₁₋₄₀ fibril structure provided by Robert Tycko²⁸ using the Autodock Vina 1.5.6 software²⁹ (Fig. S1 and S2†). In the docking process, the Ru²⁺ was replaced by Fe²⁺ since the parameters for Ru were not available in the program. This approximation was not expected to affect the binding poses, given the [Ru(bpy)2(dpqp)]²⁺ complex interacts with the Aβ fibril only through its dpqp and bpy aromatic ligands.

The energetically most stable poses of the $[Ru(bpy)_2(dpqp)]^{2+}$ Aβ complex provided by the docking procedure were equilibrated through 200 ns all-atom MD simulations using the AMBER 03 force field as implemented in the GROMACS program30,31 in explicit aqueous solution. This complex was placed in cubic boxes $(80 \times 80 \times 80 \text{ Å})$ filled with TIP3P³² water molecules. Sodium and chloride ions were added to simulate a physiological concentration of 0.154 mM. A matrix of distance restraints (semi-rigid model) with a high energetic penalty (≥1000 kJ mol⁻¹) was added to the [Ru(bpy)₂(dpqp)]²⁺ complex in order to accurately maintain the geometry as well as the coordination to the metal center. Parameters for ruthenium were obtained from a previous experimental work done with a similar kind of ligand environment.11 The root mean square deviation (RMSD) of the trajectories for sites 1 and 2 are shown in Fig. S3.† Analysis of the trajectories and simulated structures were performed with the inbuilt tools of the GROMACS program. The binding free energies between the $A\beta_{40}$ fibril and $[Ru(bpy)_2(dpqp)]^{2+}$ complexes were calculated using a thermodynamic cycle that defines the bound and unbound states applying the lambda (λ) particle approach.33,34 The dependence of these structures on force field parameters and water models is further assessed by performing MD simulations with AMBER 99-ILDN force field35 and TIP4P-EW water model³⁶. The structures provided by 50 ns production phase of these simulations were subsequently used to compute binding energies using the molecular mechanics/Poisson–Boltzmann surface area (MM/PBSA) method.³⁷

Results and discussion

Photoluminescence lifetime is an intrinsic property of an emissive probe, which represents the average time a molecule spends in the excited state before emitting a photon. The photoluminescence decay can be expressed as:

$$I(t) = \sum_{i=1}^{n} A_i \exp(-t/\tau_i)$$
 (1)

where A_i is the photoluminescence intensity at time zero (also called the preexponential factor or amplitude) and τ_i is the lifetime of the ith component. The excited state decay of a photoluminescent molecule in dilute solution of a pure solvent can be typically described with a single exponential (i = 1 in eqn (1)). However, when more complex environments are present, then multiple exponentials are needed to describe the time-decay curves. The photoluminescence lifetime of d6 metal polypyridyl complexes is generally in the range of tens of nanoseconds to microseconds due to a mixed singlet-triplet excited state.38-46 This photoluminescence is sensitive to the microenvironment around the complex. For [Ru(bpy)₂(dpqp)]²⁺ in aqueous solution, the lifetime of the free complex is ca. 446 ns (shown in Fig. 1 blue curve). The lifetime of [Ru(bpy)₂(dpqp)]²⁺ does not change significantly in the presence of AB monomers (Fig. S4†). We found that in the presence of Aβ fibrillar aggregates, the photoluminescence decay curves show a biexponential decay profile with lifetimes of ca. 446 ns and ca. 1082 ns (Fig. 1, red curve). The fastest component is assigned to $[Ru(bpy)_2(dpqp)]^{2+}$ free in solution, and the slow component is related to $[Ru(bpy)_2(dpqp)]^{2+}$ bound to A β fibrils. This significant increase in photoluminescence lifetime is selective for AB

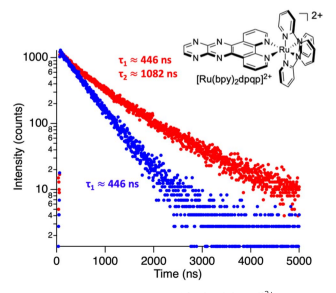


Fig. 1 Photoluminescence decay of $[Ru(bpy)_2(dpqp)]^{2+}$ in aqueous solution (blue) and in an aqueous suspension of $A\beta$ fibrils (red).

fibrils and not observed for oligomers or monomers, allowing the use of $[Ru(bpy)_2(dpqp)]^{2+}$ as a probe for detecting A β fibril formation and binding sites. This is because the lifetime and preexponential factors are proportional to the emission intensity of each component and ultimately to the concentration of the free and A β -bound species. The total photoluminescence intensity (for a two lifetimes decay) can be determined by integrating the decay law (eqn (1)) as:

$$I = I_1 + I_2 = \int_0^{+\infty} (A_1 \exp(-t/\tau_1) + A_2 \exp(-t/\tau_2)) dt$$

= $A_1 \tau_1 + A_2 \tau_2$ (2)

where I_1 and I_2 are the intensities of component 1 and 2, A_1 and A_2 are the amplitudes of component 1 and 2, and τ_1 and τ_2 are the lifetimes of component 1 and 2, respectively.

To study the interaction of $[Ru(bpy)_2(dpqp)]^{2+}$ and A β , we performed a titration experiment where the concentration of $[Ru(bpy)_2(dpqp)]^{2+}$ is held constant and the integrated photoluminescence is plotted as a function of the concentration of A β fibrils. Interestingly, the emergence of the 1082 ns component can be tracked clearly (Fig. 2a red curve) and shows a behavior similar to previous amyloid binding dyes that have been fitted to the one-site model below:

$$I_{\text{bound}} = \frac{1}{2} \delta_{Ru-A\beta} \left[K_{\text{d}} + [Ru]_{0} + \frac{[A\beta]_{0}}{n} - \sqrt{\left(K_{\text{d}} + [Ru]_{0} + \frac{[A\beta]_{0}}{n} \right)^{2} - 4[Ru]_{0} \frac{[A\beta]_{0}}{n}} \right]$$
(3)

where, $\delta_{\text{Ru}-\text{A}\beta}$ is the proportionality constant that relates the photoluminescence signal and the concentration of [Ru(bpy)₂(-dpqp)]²⁺, K_d is the dissociation constant, [Ru]₀ and [Aβ]₀ are the total concentrations of [Ru(bpy)₂(dpqp)]²⁺ and Aβ fibrils respectively, and n is the number of Aβ monomers that come together to form the binding site. A one-site model implies that as the 1082 ns lifetime component increases in intensity, the 446 ns

component would decrease proportionally. However, when we examine the intensity of the 446 ns component, the lifetime seems to decrease initially before reaching to a steady-state and not decaying to zero. While the 1082 ns component intensity seems to be leveling off near 80 μ M A β fibrils (shown in Fig. 2a, red curve), the 446 ns component still shows more than 60% of its original intensity (shown in Fig. 2a, blue curve). This behavior is completely unexpected and only observable due to the ability of time-resolved photoluminescence spectroscopy to provide information about the different species present. This new information led us to conclude that a different binding model is necessary to explain the data in Fig. 2a.

The behavior of the data in Fig. 2a can be explained by considering an additional binding site (site 2) that would allow the binding of $[Ru(bpy)_2(dpqp)]^{2+}$ to A β without influencing its lifetime. This is in contrast with what happens when the probe binds to site 1, which promotes an increase in lifetime to *ca.* 1082 ns. Due to the relative rigidity of the A β fibril's backbone (in comparison with well-folded proteins), we will assume these binding sites are independent of each other. For a two-site non-cooperative binding model, we will use the expression from Wang & Jiang, ⁴⁷ as implemented by Brautigam and co-workers: ⁴⁸

$$[Ru]_{b} = [Ru]_{b1} + [Ru]_{b2} = \frac{\frac{[A\beta]_{0}}{n}[Ru]}{\frac{R_{d1}}{K_{d1}} + [Ru]} + \frac{\frac{[A\beta]_{0}}{m}[Ru]}{\frac{R_{d2}}{K_{d2}} + [Ru]}$$
(4)

where $[Ru]_b$ is the total concentration of $[Ru(bpy)_2(dpqp)]^{2^+}$ bound to $A\beta$, $[Ru]_{b1}$ and $[Ru]_{b2}$ represent the concentration of $[Ru(bpy)_2(dpqp)]^{2^+}$ bound to sites 1 and 2 respectively, n and m represent the number of monomers making up binding sites 1 and 2 respectively, K_{d1} and K_{d2} are the dissociation constants for sites 1 and 2 respectively, [Ru] is the concentration of free $[Ru(bpy)_2(dpqp)]^{2^+}$, and $[A\beta]_0$ is the total $A\beta$ concentration. By substituting the mass conservation equation in (4) we get:

$$[Ru]^3 + a[Ru]^2 + b[Ru] + c = 0$$
 (5)

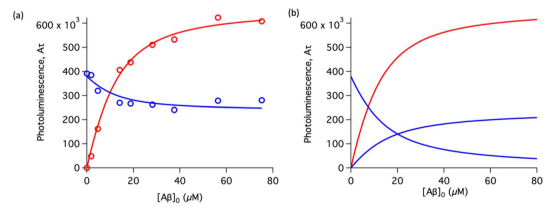


Fig. 2 Binding equilibrium experiments. (a) Titration curves showing the different components of the decay of $[Ru(bpy)_2(dpqp)]^{2+}$ with different concentrations of Aβ fibrils. The red circles represent the intensity of the component with *ca.* 1082 ns lifetime. The blue circles represent the intensity of the component with *ca.* 446 ns lifetime. Red and blue solid lines represent non-linear least-square fits to eqn (11) and (12) respectively, fitted simultaneously using the same floating parameters. (b) Simulated emission of $[Ru(bpy)_2(dpqp)]^{2+}$ bound to site 1 (increasing red curve), site 2 (increasing blue curve), and free in solution (decreasing blue curve) using eqn (1) and the data in Table S1a.† The sum of both blue curves in (b) reproduces the blue curve in (a).

where,

$$a = K_{d1} + K_{d2} + \frac{[A\beta]_0}{n} + \frac{[A\beta]_0}{m} - [Ru]_0$$
 (6)

$$b = K_{d1}K_{d2} + K_{d2}\frac{[A\beta]_0}{n} + K_{d1}\frac{[A\beta]_0}{m} - (K_{d1} + K_{d2})[Ru]_0$$
 (7)

$$c = -K_{d1}K_{d2}[Ru]_0 \tag{8}$$

where [Ru]₀ is the total concentration of [Ru(bpy)₂(dpqp)]²⁺. This equation yields only one physically meaningful root:⁴⁷

$$[Ru] = -\frac{a}{3} + \frac{2}{3} \left(\sqrt{a^2 - 3b} \right) \cos\left(\frac{\theta}{3}\right) \tag{9}$$

$$\theta = \arccos\left(\frac{-2a^3 + 9ab - 27c}{2\sqrt{(a^2 - 3b)^3}}\right)$$
 (10)

Combining (9) with mass action equations, it is possible to derive a mathematical model that fits the data in Fig. 2. The saturation curve for the 1082 ns component can be fitted to (see ESI† appendix 1 for details):

$$A_{1}\tau_{1} = \frac{[Ru]_{b1}}{\delta_{1}} = \frac{1}{\delta_{1}} \left(\frac{[Ru]}{K_{d1}}\right) \left(\frac{[Ru]_{0} - \frac{[A\beta]_{0}}{n} - [Ru]}{\left(\frac{[Ru]^{2}}{(K_{d1})(K_{d2})} - 1\right)}\right) \left(1 + \frac{[Ru]}{K_{d2}}\right)$$

where δ_1 is a proportionality constant that relates the intensity of the 1082 ns component with its concentration and [Ru] is given by (9). Similarly, the saturation curve for the 446 ns component (composed of free and bound ligand) can be fitted to:

$$A_2\tau_2 = \frac{[\mathbf{R}\mathbf{u}]_{b2} + [\mathbf{R}\mathbf{u}]}{\delta_2}$$

$$= \frac{1}{\delta_{2}} \left(\left(\frac{[Ru]}{K_{d2}} \right) \left(\frac{[Ru]_{0} - \frac{[A\beta]_{0}}{m} - [Ru]}{\left(\frac{[Ru]^{2}}{(K_{d1})(K_{d2})} - 1 \right)} \right) \left(1 + \frac{[Ru]}{K_{d1}} \right) + [Ru] \right)$$
(12)

where δ_2 is a proportionally constant that relates the intensity of the 446 ns component with its concentration and [Ru] is given by (9). Here, since the lifetime of $[Ru(bpy)_2(dpqp)]^{2+}$ does not change between free in solution *versus* bound to site 2, we make the reasonable assumption that the quantum yield of free and bound $[Ru(bpy)_2(dpqp)]^{2+}$ is similar and a single δ_2 value is adequate to describe the trend. While (11) and (12) could be fit independently to $A_1\tau_1$ and $A_2\tau_2$ respectively, the real challenge is to fit the two sets of data with two equations simultaneously, while minimizing the parameters K_{d1} , K_{d2} , δ_1 and δ_2 . Parameters n and n have been selected to be 5 for simplicity and implies a single ruthenium complex is bound to 5 A β monomers (see MD simulation below for rationale). Fig. 2a, S5a and b \dagger shows

Table 1 Averaged values of parameters for the binding of $[Ru(bpy)_2(dpqp)]^{2+}$ to A β fibrils described by a non-cooperative two-sites model

Parameter	Values ^a
K_{d1}	$3\pm1~\mu\mathrm{M}$
$K_{ m d2}$	$2.2\pm0.5~\mu\mathrm{M}$
δ_1	$(2.6\pm0.4) imes10^{-6}~{ m M}$
δ_2	$(10.9 \pm 0.5) imes 10^{-6} \mathrm{\ M}$

 a The two data sets in Fig. 2a, S5a and b were simultaneously fit to eqn (11) and (12) (global fit) to obtain the dissociation constants and proportionality constants.

curve fits of the data, and Table S1† displays the obtained parameters. Table 1 shows the averages and standard deviations of these three sets of experiments. Using the data in Table S1a,† we simulated $[Ru(bpy)_2(dpqp)]^{2+}$ binding to site 1, site 2, and free in solution at different concentrations of A β (Fig. 2b). The two blue curves, one increasing and one decreasing, represent the binding of $[Ru(bpy)_2(dpqp)]^{2+}$ to site 2 and the disappearing of free $[Ru(bpy)_2(dpqp)]^{2+}$ as the A β concentration increases, respectively. The sum of the two blue curves in Fig. 2b, yield the blue curve in Fig. 2a.

The ability to extract the different species contributing to an emission signal has led us to propose the existence of two different binding sites on AB: site 1 that affects the lifetime of $[Ru(bpy)_2(dpqp)]^{2+}$ and site 2, which shows a smaller dissociation constant and does not present a change in lifetime upon binding to Aβ. To investigate the nature of these binding sites, we combined molecular docking with all-atom MD simulations. We observed in the energetically stable poses (Fig. S1 and S2†) provided by the docking procedure that the metal complex was bound to the Val18-Phe20 region (Fig. 3) and on the surface near Asn27 on the Aβ fibril (Fig. 4). In the Val18-Phe20 site, the bpy and dpqp ligands of the metal complex associate with the fibrils through hydrophobic interactions. In detail, both bpy and dpqp ligands interact with the side chains of multiple Phe20 via CH- π and π - π interactions, and the aromatic ring of dpqp ligand interacts with the side chain of Val18 via CH- π interaction. This binding site largely remained unchanged throughout the 200 ns MD simulations in explicit aqueous solution (Fig. S3†). In the equilibrated structure, there was only a slight shift of the metal complex towards Phe20 residues (Fig. 3a). In particular, both CH- π and π - π interactions between the dpgp ring and Phe20 became stronger, while CH- π interaction with the side chains of Val18 got weaker (Fig. 3b). At this site, the dpqp ligand is deeply buried in the hydrophobic cleft of the fibril, whereas both bpy fragments are located on the opposite side towards the solvent (Fig. 3c). This binding site was similar to the one proposed for the binding of [Ru(bpy)2(dppz)]2+11 and $[Re(CO)_3(dppz)(Py)]^{+16}$ to the A β fibril. In this site, the [Ru(bpy)₂(dpqp)]²⁺ complex was found to interact with at least four Aβ monomers. The same site was provided by simulations performed using the AMBER 99-ILDN/TIP4P-EW method (Fig. S6a†). For the plots in Fig. 2, we used one more Aβ monomer to establish the binding sites (n = m = 5) in order to

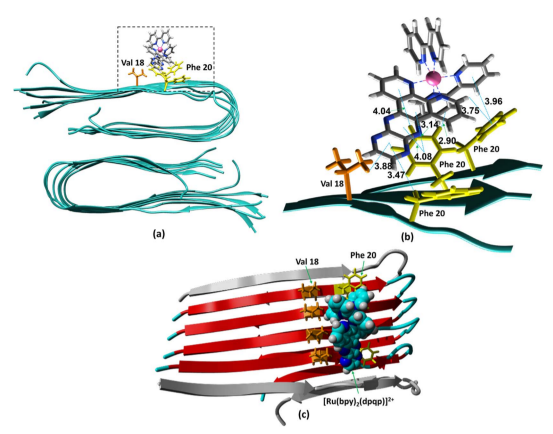


Fig. 3 Binding of $[Ru(bpy)_2(dpqp)]^{2+}$ to $A\beta$ site 1 by MD simulations. (a) The most representative structure of $[Ru(bpy)_2(dpqp)]^{2+}$ bound to a 2-fold $A\beta_{9-40}$ fibril after MD simulations, and (b) zoomed view showing binding interactions. (c) The projection of the $[Ru(bpy)_2(dpqp)]^{2+}$ complex on the $A\beta$ axis (the red colored peptides represent the projection).

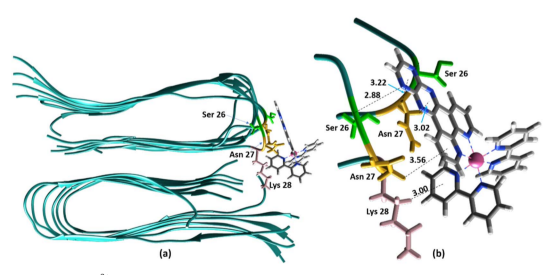


Fig. 4 Binding of $[Ru(bpy)_2(dpqp)]^{2+}$ to the site 2 of A β by MD simulations; (a) the most representative structure of $[Ru(bpy)_2(dpqp)]^{2+}$ bound to a 2-fold A β_{9-40} fibril after MD simulations and (b) zoomed view showing binding interactions at site 2.

provide a minimum separation between consecutive metal complexes for fittings to eqn (11) and (12). Given the strong change in polarity around the dpqp ligand, and knowing that the $[Ru(bpy)_2(dpqp)]^{2+}$ excited state is localized in the dpqp ligand, binding to this site is consistent with the change in the

lifetime of $[Ru(bpy)_2(dpqp)]^{2^+}$. It is noticeable that the dissociation constant calculated for $[Ru(bpy)_2(dpqp)]^{2^+}$ (Table 1, K_{d1}) is larger than that calculated for $[Ru(bpy)_2(dppz)]^{2^++1}$ and $[Re(CO)_3(dppz)(Py)]^{+16}$. This is consistent with the larger polarity of the dpqp ligand in comparison with dppz, which leads to

a weaker binding to the Val18-Phe20 hydrophobic pocket (and therefore larger dissociation constants).

Alternatively, for the other site (site 2), the backbone of Ser26 shows H bonds with the N atom on the dpqp ring. Here, the ruthenium complex is surrounded by hydrophilic amino acids such as Ser, Asn and Lys, which is likely the reason why its lifetime is barely affected. Previously, we proposed this hydrophilic binding site for [Ru(bpy)₂(dppz)]²⁺, but it was discarded at the time because the location of amino acids 1-8 made this site less accessible to [Ru(bpy)₂(dppz)]²⁺.11 It must be noted that amino acids 1-8 assume a random coil conformation and are not present in the model used.^{28,49,50} Simulations on this truncated peptide $(A\beta_{9-40})$ using two different force fields and water models provide similar structures for this binding site (Fig. S6b†). The binding free energy (BFE) computed utilizing λ dynamics of this site is -18.4 kcal mol^{-1} , which is 11.0 kcal mol⁻¹ more favorable than the one computed for site 1 (Table S2†), and can compensate for the steric constraints of amino acids 1-8. The BFE calculated through the MM/PBSA method for the AMBER 03/TIP3P and AMBER 99-ILDN/TIP4P-EW methods is also favorable by -16.7 and -11.8 kcal mol⁻¹, respectively, in comparison to the site 1. This difference in BFE is caused by stronger hydrogen bonding at this site in comparison to the interactions at site 1. Interestingly, the more negative BFE value observed for site 2 agrees qualitatively with the smaller dissociation constant obtained experimentally for this site. However, quantitatively, the calculated BFE values indicate that binding to site 2 is more than twice as favorable as site 1, while the difference between the calculated K_{d1} and K_{d2} are small. The larger than expected K_{d2} can be explained due to the presence of amino acids 1-8 in the real peptide that is expected to partially block the access to Ser26, Asn27, and Lys28 reducing the affinity of site 2 to $[Ru(bpy)_2(dpqp)]^{2+}$.

Time-resolved photoluminescence could also be used as a tool to detect Aβ aggregation in real-time. For this, we incubated Aβ at 37 °C with orbital shaking at 600 rpm and analyzed aliquots at different time points using time-resolved spectroscopy. The photoluminescence decay curves for [Ru(bpy)₂(dpqp)]²⁺ were obtained at different incubation times. The lifetime decay curves were fitted to eqn (1) (biexponential decays were only used when monoexponential decays did not satisfactorily fit the data), allowing for the photoluminescent intensity of $[Ru(bpy)_2(dpqp)]^{2+}$ bound to site 1 to be selectively determined (eqn (2)). The real-time assay is shown in Fig. 5 and displays a typical sigmoidal curve for Aβ aggregation. The intensity is low during the lag phase (0-100 minutes) where fibril growth does not take place, and AFM shows only small spherical structures, which correspond to Aβ monomers and oligomers. In this lag phase, $[Ru(bpy)_2(dpqp)]^{2+}$ is in the presence of soluble AB (which could contain monomers and oligomers), 15 but there was no observed difference in lifetime. This is followed by a phase where fibrils are formed and elongated (propagation) which occurs from 100-320 minutes. AFM images obtained from samples at 120 minutes incubation, shortly after the lag phase, showed some short A\beta fibrils. A saturation phase is reached after 240 minutes where most A β has been converted to fibrils. AFM of the 240 minutes incubation sample displays

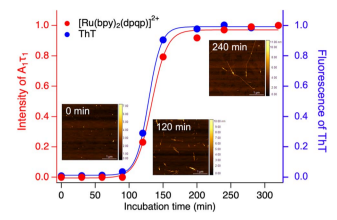


Fig. 5 Monitoring the formation of $A\beta$ fibrils using photoluminescence lifetime (red circle) and ThT fluorescence (blue circle). Inserted images are the AFM images of samples incubated for 0, 120 and 240 minutes.

abundant long A β fibrils. Parallel experiments done by measuring the fluorescence of ThT in the same samples show good agreement between the ThT fluorescence and the time-resolved deconvolution of the $[Ru(bpy)_2(dpqp)]^{2+}$. The extracted halftime of aggregation from the sigmoidal curves in Fig. 5 were 128 min and 133 min for ThT fluorescence measurement and $[Ru(bpy)_2(dpqp)]^{2+}$ lifetime measurement, respectively. This corroborates that both probes respond similarly to the aggregation of A β .

Throughout this paper, we have shown that time-resolved spectroscopy can be used as a tool to extract information on molecular binding not accesible through conventional steadystate spectroscopy. To consolidate these observations, we explored two common polypyridine metal complexes without a conventional Aβ binding element, for example, dppz or dpqp. We used commercially available [Ru(bpy)₃]²⁺ and synthesized [Ir(ppy)2(bpy)]+, and obtained the time-resolved decays as a function of the concentration of AB (ESI Fig. S7a† and 6a respectively). [Ru(bpy)₃]²⁺ showed a 380 ns lifetime in airequilibrated aqueous solution. Similar to [Ru(bpy)₂(dpqp)]²⁺, the lifetime decay of [Ru(bpy)₃]²⁺ became biexponential with a second component of 774 ns when AB nanofibrils were added (Fig. S7a†). The photoluminescence intensities of the different components (obtained from eqn (2)) as a function of AB concentration were plotted in Fig. S7b.† The intensities of the 380 ns component did not reach zero, similar to the short lifetime component of [Ru(bpy)₂(dpqp)]²⁺. Therefore, we fitted the data with the two-binding site model based on eqn (11) and (12). The resulting dissociation constants for 774 ns and 380 ns were 21 μM and 4.6 μM, respectively (Table S3a†). In the case of $[Ru(bpy)_2(dpqp)]^{2+}$, the dissociation constant of the longer lifetime component was significantly smaller than $[Ru(bpy)_3]^{2+}$. This supported that the extended π system is important for the hydrophobic interaction between Val18-Phe20 of Aβ and metal complexes. For [Ir(ppy)₂(bpy)]⁺, the decay of the complex without A β was monoexponential (ca. 41 ns), while with A β , the decay shows three components with lifetimes of ca. 41 ns, 252 ns, and 572 ns (Fig. 6a). The latter two are due to the complex

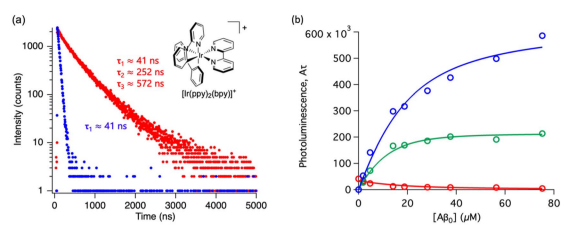


Fig. 6 Photoluminescence lifetime of $[lr(ppy)_2(bpy)]^+$ with Aβ fibrils. (a) Time decay curves of $[lr(ppy)_2(bpy)]^+$ in aqueous solution (blue) and in the presence of Aβ fibrils (red). (b) Titration curve showing the different components of the decay curves of $[lr(ppy)_2(bpy)]^+$ with different concentrations of Aβ fibrils. The red circles represent the intensity of the component with *ca.* 41 ns lifetime. The blue circles represent the intensity of the component with *ca.* 252 ns. Curves are fits to eqn (S27)–(S29).†

bound to A β . The curves in Fig. 6b were fitted to a two-binding site model based on eqn S27 to S29 resulting in dissociation constants of ca. 9 and 1.8 μ M (Table S3b†). These experiments resulted in two unprecedented observations: (1) ruthenium and iridium complexes without an amyloid binding element can bind to A β , and (2) they bind to two different sites on the fibrils.

Conclusion

In summary, our studies have shown that time-resolved photoluminescence spectroscopy can be used to deconvolute binding sites of $[Ru(bpy)_2(dpqp)]^{2+}$ to A β fibrils. The dissociation constants for each binding site can be mathematically determined and reveal specific information about [Ru(bpv)₂(dpqp)]²⁺ bound to Aβ. By considering a two sites non-cooperative binding model, we propose the existence of two different binding sites on Aβ: one that affects the lifetime of [Ru(bpy)₂(dpqp)]²⁺ (driven by hydrophobic interactions) and one in which the lifetime is unchanged (driven by polar interactions). MD simulations lead us to propose that these two sites are the hydrophobic cleft between Phe20 and Val18 and the fibril side nearby Asn27. We also showed that we can use [Ru(bpy)₂(dpqp)]²⁺ and time-resolved spectroscopy to monitor the aggregation of $A\beta$ in real-time, which was confirmed by ThT assays and AFM. Finally, we used time-resolved spectroscopy to demonstrate that the common polypyridine complexes, [Ru(bpy)₃]²⁺ and [Ir(ppy)₂(bpy)]⁺, can bind to Aβ. This study demonstrates that timeresolved photoluminescence spectroscopy can be a powerful tool not only for excited-state dynamics but also for sensing and uncovering binding sites, which could be hidden in steady-state photoluminescence studies. The method developed in this paper can be extrapolated to study a variety of biomolecules, potentially revealing new binding sites. In addition, these principles can be used to enhance the application of FLIM to the study of amyloid aggregates. Understanding amyloid binding sites is central to the development of A\beta-binding drugs that inhibit or quench the toxicity of Aβ to neural cells.

Data availability

We have included most of the data used for this manuscript in the paper's body or the ESI.† Specific numeric values or files are available upon request from the authors.

Author contributions

Conceptualization, B. J., U. U., R. P., and A. A. M.; methodology, B. J., U. U., R. P., and A. A. M.; investigation, B. J., U. U., A. A., V. M. J. A., L. S., Z. M. S. H., and K. M. W.; writing – original draft, B. J., U. U., V. M. J. A., L. S., R. P., and A. A. M.; writing – review & editing, B. J., U. U., A. A., V. M. J. A., L. S., Z. M. S. H., and K. M. W., R. P., and A. A. M.; funding acquisition, R. P. and A. A. M.; resources, R. P., and A. A. M.; supervision, R. P. and A. A. M.

Conflicts of interest

There are no conflicts to declare.

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