Picomolar-sensitive β-amyloid fibril fluorophores by tailoring the hydrophobicity of biannulated π-elongated dioxaborine-dyes

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ABSTRACT

The pathological origin of Alzheimer’s disease (AD) is still shrouded in mystery, despite intensive worldwide research efforts. The selective visualization of β-amyloid (Aβ), the most abundant proteinaceous deposit in AD, is pivotal to reveal AD pathology. To date, several small-molecule fluorophores for Aβ species have been developed, with increasing binding affinities. In the current work, two organic small-molecule dioxaborine-derived fluorophores were rationally designed through tailoring the hydrophobicity with the aim to enhance the binding affinity for Aβ fibrils—while concurrently preventing poor aqueous solubility—via biannulate donor motifs in D→A dyes. An unprecedented sub-nanomolar affinity was found (Kd = 0.62 ± 0.33 nM) and applied to super-sensitive and red-emissive fluorescent staining of amyloid plaques in cortical brain tissue ex vivo. These fluorophores expand the dioxaborine-curcumin-based family of Aβ-sensitive fluorophores with a promising new imaging agent.

1. Introduction

Ablative deposits of amyloid beta (Aβ) peptides are a major pathological hallmark in Alzheimer’s disease (AD) [1,2]. Aβ pathology first starts to occur in the asymptomatic stage of AD onset, and the extracellular Aβ accumulation in matured Aβ aggregates, known as amyloid plaques (or senile plaques), is primarily observed in the frontal and temporal lobes. This pathology gradually progresses to affect several regions of the neocortex, hippocampus, as well as cerebrovasculature [3–5]. Since AD is accompanied by a long prodromal phase of approximately 20 years; aberrant Aβ deposits can be found at least a decade before pathological symptoms, including memory loss, confusion, and cognitive decline of dementia [6,7]. Early diagnosis of AD is essential for establishing a prevention strategy [8,9]. At the same time, currently uncertainty remains with regards to treatment options and aggressive efforts are directed at finding a cure for AD [10]. Given this wide time gap between the discovery of deposited Aβ and the appearance of cognitive dysfunction, this provides a potential opportunity to diagnose and initiate treatment of AD. However, it requires an accurate and reliable visualization method. Additionally, detecting senile plaques in postmortem brain sections is an indicator of AD progression. Accurately and sensitively detecting this insoluble analyte is indispensable in histological and pathological studies of AD, thereby improving the quantification and detection of the neuropathological lesions [11]. Among the current tools to reveal AD pathology, fluorescence spectroscopy is an incomparably convenient and economical experimental technique, usually in a non-invasive manner [12]. Hence, fluorescence spectroscopy has been established as a cornerstone in basic and pre-clinical research as a fundamental tool at (sub-)cellular and small-animal scales. Above all, a high degree of sensitivity is required to enable an

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excellent performance of these fluorescent dyes toward potential targets in biological environments [13]. This sensitivity can be achieved through various means, for instance, functional selectivity where the fluorescent behavior of the fluorophore depends on environmental factors, conformational changes or in situ liberation of a dye, or superior binding affinity [14]. The latter has the advantage that tight and selective binding of a fluorophore would potentially enable future developments into other imaging modalities that could find clinical applications, e.g. Positron Emission Tomography (PET) probes as 11C or 18F-labeled agents [15,16].

Unlike enzymes or receptors with specific binding sites or reactivity, pernicious, various amyloid proteins involved in amyloidosis such as Aβ, α-synuclein, insulin, and hyperphosphorylated tau protein (p-Tau), are atypical proteins that exhibit less predictable binding modes with small-molecular ligands [17,18]. More concretely, previous studies suggested that the Aβ polypeptide is regularly ordered in a structural characteristic ‘cross-β-sheet’ scaffold, in which adjacent chain segments are tightly folded in an anti-parallel way within the fiber lattice [19]. These β-strand segments’ peptide chains run perpendicular to the long fibril, and van der Waals forces and electrostatic forces preserve its conformational stabilization in an aqueous system [20]. This fiber structure possesses approximately 25% of uninterrupted hydrophobic residues, and the compact coil structure displays metastability [6]. These characteristics could contribute to a global conformational rearrangement and intermolecular hydrophobic interaction, such as β-sheet secondary structure caused by fibrilization. Especially, Aβ-42 assumes a shape of a double-horseshoe-like cross-β-sheet as a separate entity with maximally buried hydrophobic side chains, in which residues 1–14 are organized and adopt a β-strand conformation [17]. Hence Aβ-42 aggregates much faster and dominates in the core of senile plaques [21,22]. Based on the intrinsic hydrophobicity of Aβ, the diverse small-molecule fluorescent dyes developed so far have been postulated to bind via hydrophobic π–π stacking interactions to these cross-β-sheets surfaces or intercalate into hydrophobic grooves and channels of Aβ aggregates [23,24]. Hydrophobic interactions arise from behavior that maximizes hydrogen bonding between water molecules and minimizes the area of contact between water and non-polar molecules [25,26]. Given the significant hydrophobicity of amyloid proteins, strong hydrophobicity of ligand molecules could effectively enhance binding affinity to cross-β-sheets or hydrophobic grooves through a hydrophobic effect in aqueous environments [27].

To date, several fluorescent dyes sensitive toward various states of Aβ aggregation have been reported to display a high binding affinity. Structural modifications, mainly π-elongation, have commonly been introduced as a critical strategy for small-molecular fluorophores to make bathochromic fluorescence emission and increase hydrophobicity in an aqueous medium [28–36]. However, the increased hydrophobicity associated with an extended π-conjugation also inevitably leads to poor aqueous solubility and increases the ligands’ promiscuity with regard to other hydrophobic proteins and cellular components. Thus, the major challenge in the design of new red-emissive fluorophores with a stronger affinity for Aβ fibrils is not only to maintain adequate aqueous solubility but also specificity and sensitivity.

A wide range of dissociation constants (K_d) have been reported so far for dyes binding Aβ fibrils, ranging from the nanomolar to micromolar affinities [23,24]. Among them, dioxaborine-based curcumin structures have been incredibly successful, most notably the CRANAD series by Ratnavat group, and have been reported to show good binding affinities due to curcumin’s intrinsic affinity of several Aβ species [37–40]. In 2009, the first reported dye of this series, CRANAD-2, demonstrated an strong affinity (K_d = 36.7 nM) for Aβ-40 fibrils and adequate photophysical properties (Table S1) [41]. The versatility of this fluorophore class has been demonstrated during the last decade, with recent work describing CRANAD-58, able to detect both soluble and insoluble Aβ-42 species in vitro and even in vivo [42]. In parallel, Liu et al. investigated a separate class of dyes, known as the DANIR series, with one of these demonstrating a K_d value of 1.9 nM, with remarkable fluorescent sensitivity [43]. To the best of our knowledge, DADNIR-2 from Cui’s group showed the highest binding affinity (K_d = 1.04 ± 0.03) reported for an Aβ-42 aggregate fluorescent probe thus far. However, its flat elongated structure and relatively high molecular weight may result in solubility issues [44].

In the current study, components of both the dioxaborine-based half-curcumin structures and the high-affinity DANIR dyes were rationally combined. These dyes were synthesized via a condensation reaction of a beta diketone with two N,N-dimethyl biannulated electron-donating groups: BZ-OB (benzofuran) and NAP-OB (napthalene) (Fig. 1). Due to the relatively small number of carbon atoms and introduced oxygen atom into the aromatic ring of benzofuran, which induces a different hydrophobicity for the whole molecule structure, compared with naphthalene. Importantly, this particular diketone moiety was chosen as it previously demonstrated an excellent aqueous solubility for the corresponding dioxaborine-based half-curcumin structures via the suppression of intermolecular π–π stacking interaction by the isobutyl group’s innate steric bulk [45].

2. Materials and methods

2.1. Quantum yield and determination of fluorescence emission and wavelength maxima

To avoid the inner filter effect during measurement of fluorescence emission, the spectra were recorded with the absorbance lower than 0.1 at wavelengths longer or equal to the excitation wavelength with HPLC grade solvents including 1,2-dichlorobenzene, 1-butanol, 1-propanol, 2-methoxy ethanol, acetone, acetonitrile, chloroform, diethyl ether, ethyl acetate, ethylene glycol dimethyl ether, ethanol, methanol, tetrahydrofuran, or toluene). The fluorescence quantum yields of BZ-OB and NAP-OB were measured versus 4-(dicyanomethylene)-2-methyl-6-(4-dimethylaminostyryl)-4H-pyran (DCM) in acetonitrile. (Φ_F1 = 0.43) [46], using a previously reported procedure [47].

2.2. Spectroscopy in a solvent with increasing viscosity

The emission spectra of BZ-OB, NAP-OB, and Thioflavin T (ThT) [48] (10 μM) in the various v/v % of ethylene glycol and glycerol (30, 50, 60, 70, and 80% of glycerol in ethylene glycol) at 25 °C were recorded. BZ-OB and NAP-OB stock solutions were prepared in dimethyl sulfoxide (DMSO), respectively, and all solutions contain a final concentration of 1% DMSO.

2.3. Solubility of BZ-OB and NAP-OB in aqueous solution

The absorbance of BZ-OB and NAP-OB was recorded at 495 and 523 nm, respectively, in phosphate buffered saline (PBS) (10 mM, pH = 7.4) containing 1% DMSO at different concentrations in the 0 – 50 μM range.

2.4. Spectroscopy in the presence of Aβ-42 and potential interferents (metal ions, amino acids, thiols, and albumin)

Stock solutions of BZ-OB and NAP-OB were prepared in DMSO and added to solutions of the analytes in PBS (Aβ-42 or bovine serum albumin (BSA)) or deionized water (DIW) (metal ions, amino acids, or thiols) to finally obtain a solution containing 1% DMSO. All emission signals were obtained after 5 min incubating at 37 °C with 300 rpm orbital shaking.

2.5. pH dependent absorbance of BZ-OB and NAP-OB

The absorbance and fluorescence spectra of BZ-OB (10 μM) and NAP-OB (10 μM) were recorded in DIW with the pH adjusted by hydrochloric acid or sodium hydroxide to the pH 2, 4, 6, 8, 10, or 12.
2.6. Photostability

A solution of BZ-OB or NAP-OB with an absorbance of 1.0 at the maximum absorbance wavelength was prepared in 1% DMSO/PBS. A 3100 K halogen lamp (Olympus LG-PS2; 12 V, 100 W) was used for irradiation, and the absorbance was recorded at 10 min intervals for 60 min.

2.7. Theoretical calculations

Density functional theory (DFT) calculations were performed using the Gaussian 16 software package [49]. The structures of NAP-OB and BZ-OB were optimized using the ωB97XD functional at the N07D level of theory, with the default IEPPCM solvation model of acetonitrile [50,51]. Molecular orbital were visualized using Gaussianview 6.1 [52]. To determine the amount of intramolecular charge transfer (ICT) character upon vertical excitation to the first excited state, a calculation based on the simplified formula can be used where the fluorescence intensity is directly proportional to the concentration of the Aβ fibril relative to the protein fibril. Assuming different binding stoichiometries resulted in binding affinities within the same order of magnitude but increased uncertainty estimation ranges.

2.8. Docking studies

The ground-state B3LYP-optimized structures of BZ-OB and NAP-OB were used as the ligands for docking studies. The cryogenic electron microscopy (cryo-EM) structure of Aβ1-42 (PDB ID: 5OQV) [56] was prepared by removing either the Glu18 or Val18 facing conformation of Phe20, and one entire protofibril was encompassed within the search area (38 × 50 × 32 Å) using AutoDock Vina [57] as the docking software, with an exhaustiveness of 1,000. The identified docking sites were then reexamined using a smaller search area of (14 × 14 × 26 Å) for the tunnel and (14 × 20 × 26 Å) for the binding sites adjacent to Phe20. The input for the docking calculations was prepared using AutoDockTools 4.2, [58] and figures were generated using the Python Molecule Viewer 1.5.6 software package [59].

9. Aβ1-42 fibril formation

0.5 mL of 2 mM sodium hydride aqueous solution was added into the purchased Aβ1-42 kit and treated with sonication at 0 °C for 30 min. 0.6 mL of 20 mM PBS was added to the sonicated solution, then incubated with an orbital shaker (300 rpm) at 37 °C for 72 h. The solution was diluted to experimental concentration with 10 mM PBS (pH = 7.4) before each experiment.

2.10. Fitting of Aβ1-42 saturation titration analysis

Aβ1-42, BZ-OB, and NAP-OB are virtually non-fluorescent, and a simplified formula can be used where the fluorescence intensity is directly proportional to the concentration of the Aβ1-42–probe (BZ-OB or NAP-OB) complex only [60]:

\[
I = \frac{1}{2} F \left( \left( [P] + N[Aβ] + K_d \right) - \sqrt{\left( [P] + N[Aβ] + K_d \right)^2 - 4N[Aβ][P]} \right)
\]

With F: a fluorescence proportionality factor, [P]: the initial concentration of BZ-OB or NAP-OB, [Aβ]: the initial concentration of Aβ1-42, N: the number of equivalent binding sites on the Aβ fibril relative to Aβ monomer, and K_d: the dissociation constant. The results in Fig. 2E and F were obtained following a multi-parameter optimization of the experimental results following the fluorescence intensity at 642 or 621 nm, respectively, using a concentration of 1 or 0.5 μM Aβ1-42 and total concentrations of BZ-OB or NAP-OB in PBS containing 1% DMSO. The slit width was set at 2.5/2.5. The best fittings were obtained for N = 0.1, i.e., one binding site per 10 Aβ1-42, consistent with the hypothesized high affinity cross-fibril binding site on the protein fibril. Assuming different binding stoichiometries resulted in binding affinities within the same order of magnitude but increased uncertainty estimation ranges.

2.11. Preparation of AD mouse brain tissues

APP/PS1/TauP301L-BiFC triple transgenic mice presenting both amyloid and tau pathology were produced by the crossbreeding of APP/PS1 (APPsw/PSEN1E9 mice) with TauP301L-BiFC mice [61]. APP/PS1/TauP301L-BiFC triple transgenic mice were used in this study.
and the littermate APP/PS1, and wild-type mice were used as controls. For tissue preparation, mice were perfused with 0.9% saline and fixed with PBS containing 4% paraformaldehyde. Frozen brain samples embedded in O.C.T. were prepared at 30 μm thickness with a cryostat (Leica).

2.12. Fluorescent probe stain and image acquisition

To evaluate amyloid staining properties of BZ-OB and NAP-OB, 10 month-old of APP/PS1 and wild type brain tissue sections were stained with 10 μM of each fluorescent probe for 30 min and mounted onto glass glides. Fluorescence images were acquired using an Axio Scan Z1 (Zeiss) (BZ-OB: λ<sub>ex</sub> = 540–546 nm and λ<sub>em</sub> = 560–640 nm). To evaluate the amyloid staining property of NAP-OB in the presence of tau aggregation, brain tissue sections of 20 month-old of APP/PS1/TauP301L–BiFC brain tissue sections were stained with Sudan Black B solution (70% ethanol containing 0.05% Sudan Black B) for 10 min to reduce auto-fluorescence of aged brain tissue. Then the tissues were washed three times with PBS containing 0.1% Triton X-100 and washed with distilled water. For counterstaining nuclei, brain tissues were stained with Hoechst (0.5 μg/mL dissolved in distilled water) for 30 min. BiFC fluorescence (λ<sub>ex</sub> = 460–490 nm and λ<sub>em</sub> = 500–640 nm) images of whole brain tissues were acquired using Operetta (PerkinElmer) and Axio Scan Z1 (Zeiss). Then, the tissues were stained with NAP-OB (300 nM) for 30 min for staining amyloid plaques. Fluorescence images were acquired using an Axio Scan Z1 (Zeiss) (λ<sub>ex</sub> = 540–546 nm and λ<sub>em</sub> = 578–640 nm).

2.13. Image J analysis

The number and sizes of fluorescence-detected amyloid plaques were quantified by using the “analyze particle” tool of Image J (NIH) software. Fluorescence intensity profiles of amyloid plaques stained with each probe were generated by using the “Interactive surface plots” tool of Image J (NIH) software. Signal to Noise ratio (S/N) was calculated from cortical area of three represent brain tissue images stained with each probe (Thioflavin S (ThS), BZ-OB, and NAP-OB). Error ranges represent SD (n = 3), slit 5/5.

3. Results and discussion

BZ-OB was synthesized via α-alkylation of deacetalized 2-(2,2-dithioethoxy)benzaldehyde with acetic acid followed by dehydration.
followed by an aldol condensation reaction of the obtained 6-(dimino)-2-naphthaldehyde (aldehyde synthesis reaction from aryl 6-halide forms of naphthalene, of the resulting ether and NAP-OB was synthesized through a Bouveault aldehyde synthesis reaction from aryl 6-halide forms of naphthalene, followed by an aldol condensation reaction of the obtained 6-(dimethylamino)benzofuran-2-carbaldehyde (BZ-CHO) or 6-(dimethylamino)-2-naphthaldehyde (NAP-CHO) with 6-methyl heptane-2,4-dione as seen in Fig. 1 and Scheme S1. Fluorophores BZ-OB and NAP-OB, as is the case for the majority of currently described fluorescent dyes for Ajβ, are D-π-A dyes and their significant Stokes’ shifts, combined with a relatively broad featureless absorbance and emission spectra suggests they likely exhibit ICT behavior upon excitation. Therefore, these dyes’ photophysical properties, particularly their ICT behavior, were investigated using several organic solvents (Table S2). The maximum wavelengths of absorption and emission of both fluorophores were measured concerning the solvent polarity, using the polarizability (induced dipole) and the dipolarity (permanent dipole) descriptions of solvent polarity, as reported in the Catalaní data set [62]. The maximum absorption wavelength correlated well with the solvent polarizability, derived from the solvent-associated electronic stabilization of the Franck-Condon state. In contrast, the maximum fluorescence emission wavelength corresponded well with the solvent dipolarity, indicating that solvent cage rearrangement, following the geometrical relaxation of the excited states of BZ-OB and NAP-OB (Figs. S15 and S16), plays a major role. Collectively, these solvatochromic effects obviously support an excited state with a considerably higher dipole moment than the ground state, which is consistent with displaying ICT behavior in the excited state [63].

Additionally, the quantum yield of fluorescence is the highest in the most apolar solvents (~100%) and gradually decreases with the solvent polarity, with only 2.1 and 1.2% observed in methanol for BZ-OB and NAP-OB, respectively, which is a common observation in ICT-based fluorescent dyes [64,65]. Given the poor fluorescence quantum yield in aqueous media, the fluorophores could exhibit potentially considerable fluorescent turn-on ratios in the presence of the hydrophobic Ajβ aggregates’ interior. Importantly, to properly perform their function as an Ajβ sensing probe and allow for potential in vivo experiments, these dyes need to maintain a reasonable solubility in aqueous media. The strategy of increasing hydrophobicity through π-elongation is a double-edged sword that can increase the binding affinity for hydrophobic analytes but simultaneously results in decreasing aqueous solubility, inducing aggregation-caused quenching (ACQ). As a maneuver to overcome an ACQ issue, an isobutyl moiety was introduced in both BZ-OB and NAP-OB, which prevented intermolecular aggregation through interference with crystal packing. The high aqueous solubility that followed from the introduction of this moiety was validated using concentration-dependent absorbance spectra. A linear relationship between concentration and absorbance was found at concentrations up to 50 µM for both BZ-OB and NAP-OB (Fig. S17), confirming the absence of aggregation in this concentration range, despite the relatively long π-conjugation of fluorophores BZ-OB and NAP-OB.

To support the existence of an ICT process in these dyes, the energy levels of their molecular orbitals in acetonitrile were calculated using DFT calculations (Fig. S18). The electron density distributions in the frontier orbitals unveiled an electron redistribution from the dimethylaniline benzofuran or dimethylamino naphthalene moieties in the highest occupied orbital (HOMO) to the dioxygen core in the lowest unoccupied orbital (LUMO). The involvement of an ICT process was further confirmed using TDDFT calculations, focusing on the degree of ICT based on changes in electron density. Both fluorophores clearly exhibited a charge transfer upon excitation to the first excited state, which was dominated by the HOMO to LUMO transition, with a change in polarity upon excitation of 17.028 Debye and 14.598 Debye for NAP-OB and BZ-OB, respectively, resulting from a larger partial charge redistribution and larger distance between the donor and acceptor barycenters for NAP-OB (0.778e- , 4.556 Å) as compared to BZ-OB (0.737e- , 4.125 Å). These results correlate well with the observed stronger sensitivity of the Stokes’ shift and maximum emission window to the polarity for NAP-OB (Fig. S16).

The absorbance and fluorescence emission spectra were determined in the presence of Ajβ fibrils. As can be seen from Fig. 2A, a maximum absorbance wavelength was observed at 523 nm and 495 nm for BZ-OB and NAP-OB, respectively, with a Stokes’ shift of approximately 120 nm for both dyes, as the maximum emission bands were observed at 642 nm and 621 nm for BZ-OB and NAP-OB, respectively (βex = 523 nm for BZ-OB and 495 nm for NAP-OB, Figs. 2A and S19). This fluorescence response emerges by two potential pathways: conformational restriction or polarity changes. As the fluorescence intensity hardly changed in function of increased viscosity in binary glycerol/ethylene glycol mixtures (Table S3 and Fig. S20), we assume that the fluorescence intensity mainly depends on microenvironmental polarity changes upon hydrophobic interactions with the β-sheet surface of Ajβ fibril aggregate. Using the absorbance and fluorescence wavelengths of these dyes as a guide, the polarity of the environment sensed by these fluorophores matched best with diethy ether for BZ-OB and tetrahydrofuran for NAP-OB, in accordance with previous reports on the polarity of the Ajβ fibril aggregates’ interior [43,60,66].

Next, a time-dependent coincidence with Ajβ pre-aggregates was carried out for BZ-OB and NAP-OB, as well as with the common Ajβ aggregate fluorescent dye ThT, as a reference (Fig. 2B). As can be seen, the fluorescence intensity increased as the aggregate was formed, and BZ-OB and NAP-OB mimicked the fluorescent behavior of ThT, confirming their selectivity for the fibril forms of Ajβ. The increased fluorescence of BZ-OB and NAP-OB was determined to be linearly dependent on the concentration of the Ajβ fibril, with a maximum of 20.5-fold and 69.4-fold fluorescence enhancement, respectively (inset of Fig. 2C and D). The fluorescent response of BZ-OB and NAP-OB to potential interferants, including metal ions, amino acids, and thiols, was negligible (Fig. 2C and D). The absorbance and fluorescence emission spectra were measured in the pH 2 to 12 range, and there was almost no change observed in the pH 4 to 10 range, while significant changes were observed at pH 2 or 12 (Fig. S23). Thus importantly, both fluorophores do not exhibit a pH dependence of the fluorescence under physiological pH conditions. BZ-OB and NAP-OB showed only a relatively weak fluorescence enhancement in the presence of BSA (Fig. S24), an essential consideration in polarity-dependent fluorescent probes for biomolecules, as albumin exhibits several hydrophobic sites and is a major carrier protein for hydrophobic drugs. In addition, a fluorescence titration of BZ-OB and NAP-OB with Ajβ fibrils revealed a binding affinity of Kd = 21.56 ± 9.45 nM for BZ-OB and 0.62 ± 0.33 nM for NAP-OB (Fig. 2E and F). This value of NAP-OB represents a significantly higher affinity for Ajβ fibrils compared to the reported literature data for other dioxaborine-based curcumin structures (Table S1). To our best knowledge, this is the highest binding affinity as yet reported for all Ajβ fibril small-molecular fluorophores.

Additionally, the limit of detection (LOD) was found to be 13.071 and 10.813 nM for BZ-OB and NAP-OB, respectively (Fig. S25). This observation is somewhat surprising given the massive difference in Kd as well as a 3.4-fold higher fluorescence turn-on ratio for NAP-OB. However, a LOD of approximately 10 nM could be rationalized due to the relatively high instrument detection limit (IDL) of the fluorescence spectrometer caused by the turbidity induced by suspended Ajβ aggregates, thus increasing the standard deviation of the blank.

A near-atomic-resolution cryo-EM structure of Ajβ fibril was used as a protein scaffold for docking studies (Fig. 2A). As suggested before for other Ajβ fluorophores, the hydrophobic tunnel architecture along the Ajβ fibril axis, consisting of the residues of Phe19, Asn27, Gly29, and Ile31, could be the major binding site of BZ-OB and NAP-OB as cross-β-sheet binding ligands (Fig. 3B, D, and E for BZ-OB and 3C, F, and G for NAP-OB). However, the results of these theoretical studies need to be interpreted with some level of caution, as they do not provide a direct rationale for the observed binding affinity difference of BZ-OB and NAP-OB.

Next, we investigated whether probes BZ-OB and NAP-OB could
potentially label amyloid plaques in biological tissue samples. Whole-brain tissue sections were prepared from 10 month-old APP/PS1 mice and littermate controls. Each brain tissue was incubated with 300 nM of BZ-OB and NAP-OB for 30 minutes. As anticipated, BZ-OB and NAP-OB-stained dot-shaped amyloid plaques in APP/PS1 brain tissue slices (Fig. 4A, S26–27). In contrast, no specific staining occurred in wild-type brain slices, in which Aβ is absent. High magnification images also showed a prominent staining ability for NAP-OB in detecting various sizes of amyloid plaques which co-localized well with ThS into compact and diffuse plaques (Fig. S28). An interactive surface plot analysis showed that NAP-OB has an outstanding S/N, which is comparable to that of ThS (NAP-OB: S/N = 3.5 ± 0.4, and ThS: S/N = 3.9 ± 0.6) (Fig. 4B). In comparison, BZ-OB was found not to stain small-sized plaques well, and its S/N was also lower than NAP-OB (BZ-OB: S/N = 1.8 ± 0.2). Quantification of the number and sizes of fluorescence-detected spots indicated that the amyloid labeling capacity of probe NAP-OB was comparable to that of ThS by presenting 68 ± 5 spots, with sizes below 100 μm² and 96 ± 11 spots, with sizes above 100 μm² (Fig. 4C). In comparison, BZ-OB detected a smaller number of amyloid plaques by showing only 12 ± 6 spots below 100 μm² and 43 ± 18 spots above 100 μm². Violin plots of spot areas indicated the distinguished staining pattern of NAP-OB comparable to that of ThS (Fig. 4D). These observations support NAP-OB as a reliable, red-colored amyloid tracer.

Next, we applied NAP-OB for staining amyloid plaques in the brain tissues of 20 month-old APP/PS1/TauP301L-BiFC mice, which present both Aβ plaques and fluorescently labeled tau aggregates [67]. Before the NAP-OB stain, Sudan Black B staining was performed to reduce autofluorescence of the aged mouse brain, and Tau-BiFC fluorescence was scanned by using a Zeiss Axio Scan (λex = 460–490 nm and λem = 500–550 nm). Tau-BiFC fluorescence indicates tau aggregates ranging from oligomers to insoluble aggregates (Fig. 5A). Then, the tissues were stained with NAP-OB (300 nM) and scanned by using a Zeiss Axio Scan.
\( \lambda_{\text{ex}} = 540-546 \text{ nm and } \lambda_{\text{em}} = 578-640 \text{ nm} \). Spot analysis indicated a total of 544 amyloid plaques stained with NAP-OB in half brain tissue of 20-month-old APP/PS1/Tau-BiFC mouse. Out of 544 plaques, 239 were tau-BiFC positive, indicating that tau proteins were co-aggregated with amyloid plaques in the brain. The median areas of tau negative and positive plaques were 51 \( \mu \text{m}^2 \) and 136 \( \mu \text{m}^2 \), respectively (Fig. 5B). This observation indicates that tau proteins are co-aggregated with large amyloid plaques, of which sizes are over 100 \( \mu \text{m}^2 \) rather than small ones. High magnification images of the subiculum show large amyloid plaques co-stained with tau-BiFC fluorescence (yellow arrows). In contrast, tau-BiFC fluorescence was not detected in most small amyloid plaques, below 100 \( \mu \text{m}^2 \) (Fig. 5C). These observations support that NAP-OB, unlike many other lipophilic pi-extended fluorescent probes, could reliably visualize senile plaques in the presence of other lipophilic protein aggregates, such as pTau tangles. In addition, its red-shifted fluorescence properties make these probes complementary with the green-fluorescent tau-BiFC system, while the emission windows of ThT and ThS overlap with BiFC and cannot be used. Thus, NAP-OB could find applications in differentiating aggregates, to study the co-occurrence of both protein aggregates in more detail, particularly so in the Tau-BiFC system.
4. Conclusions

In conclusion, two novel dioxaborine-based fluorescence dyes were designed and synthesized. In particular, the strategy of applying bian-nulated $\pi$-elongation to electron donor groups has successfully offered significantly lowered $K_d$ values, i.e., higher binding affinity, for $\text{A}\beta_{1-42}$ fibrils. To the best of our knowledge, fluorophore NAP-OB exhibits the most potent affinity for $\text{A}\beta_{1-42}$ aggregates reported to date. The precise detection of $\text{A}\beta$ species and knowledge about their emergence, accumulation, and evolution is indispensable for a detailed understanding of these neurodegenerative diseases, and the development of diagnostics and clinical intervention strategies, especially for early-stage disease. The structural modification presented here to enhance the sensitivity for $\text{A}\beta$ may provide a rational path to the development of future small-molecule fluorophores. In terms of application, due to the efficient $^{18}\text{F}/^{19}\text{F}$ isotopic exchange at the central difluoro-boron complex structure, NAP-OB could find further applications as a dual fluorescence and PET imaging agent.

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CRediT authorship contribution statement

Jusung An: Project administration, Conceptualization, Data curation, Investigation, Software, Formal analysis, Writing – original draft. Peter Verwilst: Investigation, Data curation, Resources, Writing – review & editing, Funding acquisition. Hira Aziz: Data curation, Software, Formal analysis, Writing – original draft. Jinwoo Shin: Data curation, Software, Formal analysis. Sungsu Lim: Data curation, Formal analysis. Ilwha Kim: Methodology, Investigation. Yun Kyung Kim: Project administration, Conceptualization, Supervision, Funding acquisition, Writing – review & editing. Jong Seung Kim: Project administration, Conceptualization, Supervision, Funding acquisition, Writing – review & editing.

Appendix A. Supplementary data

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