1. Introduction

Near-infrared (NIR) dyes with a high quantum yield, good biocompatibility, and a fast clearance have gained extensive attention in materials chemistry, optical imaging, and relevant biomedical applications.\(^1\-^3\) Because of its easy operation, low cost, and real-time monitoring, near-infrared fluorescence (NIRF) imaging is widely used for detection in various pathophysiological procedures and pre-clinical applications.\(^4\,^5\)

However, there are few functional NIR dyes that are currently feasible for the detection of specific diseases due to three main weaknesses: (1) poor properties such as suboptimum optical absorption, insufficient stability, poor hydrophilicity, and a low quantum yield are still fatal obstacles for imaging biological tissues \textit{in vitro} and \textit{in vivo};\(^6\-^8\) (2) most NIR dyes possess extremely low disease targeting and binding capacities, which greatly preclude their use in bioapplications;\(^9\) (3) there is no rational or ideal approach for developing an NIR dye with complementary imaging capabilities for positron emission tomography (PET)/photoacoustic tomography (PAT) or PET/PAT/NIR fluorophores on one platform.

Cerebral amyloid angiopathy (CAA) occurs in most patients with Alzheimer’s disease (AD), which is characterized by amyloid beta (A\(\beta\)) deposition in the walls of leptomeningeal arteries, cortical arteries and veins in the brain.\(^10\) Considering that A\(\beta\) deposits can occur simultaneously or separately in both blood vessel walls and brain parenchyma, it is critical to develop a precise strategy to map the abnormality for further pinpointing CAA \textit{in vivo}.\(^11\) However, the development of effective amyloid-specific optical agents still remains a great challenge. Bacskai \textit{et al.} reported that vascular amyloid deposits in mice were labeled by thioflavin-T, which, however, has drawbacks of nonspecific binding and a short absorption peak at 500 nm.\(^12\) In addition, there are fewer binding sites associated with the A\(\beta\) plaque of the vessels than those reported for the AD brain.\(^13\) Therefore, this demands a diagnostic technique suitable for
vascular imaging with a high sensitive agent that is specific to plaque.

PET, a powerful tool for the early detection of AD, can detect Aβ deposits with high sensitivity using amyloid specific tracers, enabling in vivo pathological and molecular diagnosis. However, limited research has been done on the development of imaging agents for the effective diagnosis of CAA. For instance, [11C]Pittsburgh compound B and [18F]Florbetapir ([18F]AV-45) have shown promise for non-invasive amyloid imaging in AD patients. Yet, these two compounds bind to Aβ aggregates in both blood vessels and parenchymal brain tissue. In order to target Aβ located in vasculature, polyethylene glycol modified bivalent and trivalent ligands or a 64Ga/99mTc conjugated complex was synthesized either with a larger molecular weight or negative electric charges to limit their permeation of blood vessels. On the downside, these compounds demanded an extra efficient ligand and were only examined in vitro but not in any living subjects. Moreover, PET is impeded by a low spatial resolution and requires an onsite cyclotron along with a radioactive source. In contrast to nuclear imaging, optical imaging offers the benefits of high sensitivity, rapid frame rates, and detection safety.

PAT, a new and emerging technique, has demonstrated prosperous imaging capabilities for samples ranging from organelles to organs, and especially for blood vessels since this method is intrinsically sensitive to hemoglobin. By converting a pulsed laser into ultrasound, it has opened up a new frontier for bioimaging of the brain with improved spatial resolution and increased penetration depth. With endogenous contrast, Yao et al. applied a functional photoacoustic (PA) microscopy system to image the cerebral metabolism and vascular morphology of a mouse brain. However, the lack of amyloid-specific contrast has confined its use in further disease-oriented applications. Therefore, the combination of using an intrinsic specific probe and noninvasive imaging method is urgent in visualizing the morphology and pathology of cerebral vascular plaque.

In our study, we designed a crotonium dye for amyloid (CDA) with limited but sufficient permeability to specifically bind with plaque in vessels (Scheme 1). This probe exhibits very strong absorption in the NIR range and produces a much stronger photoacoustic signal than other prominent materials of the same mass concentration. Moreover, the unique material is able to strongly bind the hydrophobic amino acid of the plaque with a strong binding energy, which can significantly improve the detection sensitivity without antibody labeling. After irradiation with an NIR laser, the CDA-3 solution displayed a surging temperature increase from 26 to 63 °C. The hyperthermia that has a great effect on amyloid formation can act as a promising way to dissolve Aβ deposits.

Multimodality imaging that provides both anatomical and functional information has brought a new avenue in precision medical research. To the best of our knowledge, this is the first report on the application of unique crotonium dye labeling with an efficient radiotracer for in vivo versatile PAT/PET/NIR fluorescence imaging. As a noninvasive method, PAT provides endogenous absorption contrast from hemoglobin and exogenous molecular contrast from the amyloid specific NIR dye, permitting the real-time visualization and monitoring of cerebral vasculature plaque. Herein, we also report for the first time the evaluation of a specific PAT probe without antibody-labeling for Aβ plaque targeting imaging in a transgenic (Tg) mouse model.

In addition, our proposed [18F]CDA-3 is among the very few PET probes that can specifically bind plaque in CAA and is available for in vivo whole-body imaging. As expected, significant advantages such as excellent sensitivity, a high spatial resolution and superior quantification are apparent in our NIR fluorescent PAT integrated with PET strategy. Our study may provide a new tool to detect brain plaque at an early stage and unveil the exact mechanisms lying behind neurovascular dysfunctions, for example, the narrowing and rupture of vessels, and related dementia.

2. Results and discussion

Inspired by the “aromatic ring-conjugated bridge-aromatic ring” planar backbone structure, we fabricated a series of biocompatible dyes with the crotonium or squarylium motif (Fig. 1a and b). The detailed synthesis processes are schematically illustrated in Schemes S1 & S2.† All of the intermediates, the final CDA and the squarylium dye for the amyloid (SDA) were characterized using 1H nuclear magnetic resonance (NMR), 13C NMR and high-resolution mass spectrometry (HRMS) (Fig. S1–S9†). The CDA compounds were further dissolved in an NaHCO3/KHCO3 solution for subsequent applications.

CDA with an electron donor–π–acceptor–π–donor structure exhibits excellent NIR absorption properties (604 nm for CDA-1, 779 nm for CDA-2, and 798 nm for CDA-3). The crotonium ring of CDA is a stable annulus mesoionic compound with an electron acceptor heart and an electron donor wing, leading to the strong NIR absorption (Fig. 1c–e). The two thiophene structures are also donor groups, and contribute to the high affinity of Aβ through binding to the hydrophobic channel. The stronger electron withdrawing ability of the croconic acid ring suggests that CDA may exhibit a more bathochromic shift and a higher sensitivity than its squaraine analogues. In Fig. S10† the peak absorption wavelengths of SDA-1 and SDA-2 were 491 nm and 658 nm, which show a 113 nm and 121 nm blue-shift for CDA-1 and CDA-2, respectively.

Scheme 1 Schematic illustration of intrinsic targeting [18F]CDA-3 for the ultrasensitive PAT/PET/NIR imaging of brain plaque.
and binding energy were simulated using a molecular docking technique, a powerful tool for studying protein–ligand interactions. As shown in Fig. 1f, the optimized geometry of the CDA compounds was a trapezoid plane with two hydrophilic terminal groups stretching out. The docking result indicated that only one site was located on the symmetrical surfaces of the fibril. CDA-3 is lying above Val18, and parallel leaning to Lys16 with hydrogen bonds formed between the carboxylate ion of CDA-3 and the carboxyl of lysine (Fig. 1g).

Most of the existing probes share the same site with 6-iodo-2-(4-dimethylamino)phenylimidazo[1,2-a]pyridine (IMPy) and nestle into the hydrophobic Val18_Phe20 channel. The calculated binding energy for CDA-3 was –9.3 kcal mol$^{-1}$, which is much more negative compared to –4.9 kcal mol$^{-1}$ for IMPy to Aβ$_{1-42}$, and suggests that the CDA compound fits rigorously into the binding pocket. This Lys16_Val18_Phe20 hydrophilic–hydrophobic dual-channel runs longitudinally to the fibril axis. The channel length and potential binding sites would increase with fibril elongation as more monomers are added. Therefore, it is forcefully confirmed that this binding site is mostly responsible for the binding of CDA-3 to the Aβ$_{1-40}$ fibrils.

To examine the feasibility of CDA-3 for detecting CAA diseases, we performed \textit{in vitro} fluorescence staining and \textit{in vivo} noninvasive NIR and PA imaging. The excitation wavelength and emission wavelength of CDA-3 was 760 nm and 811 nm, respectively. A very weak interaction was observed in the presence of the Aβ$_{1-40}$ monomers and HSA as shown in Fig. S14.$^{†}$ In contrast, the fluorescence intensity increase for CDA-3 upon binding to the Aβ$_{1-40}$ aggregates suggested an obvious interaction between CDA-3 and the Aβ$_{1-40}$ aggregates (Fig. 1h).

Here, we also evaluated this croconium dye as a novel PA contrast agent for imaging Aβ plaque in the brain. The PA properties of CDA-3 were compared with some of the most commonly used probes such as indocyanine green (ICG), Prussian blue (PB), and gold nanorods (GNRs). The absorption peaks of all of these dyes fall in the NIR window (Fig. 2a). After irradiation with an 808 nm laser (1.0 W cm$^{-2}$, 10 minutes), the temperature of the CDA-3 solution increased quickly from 26 to 63 °C and the solution exhibited no measurable change in ultraviolet (UV) absorbance after laser irradiation. Under the same conditions, the GNRs and PB produced a much smaller temperature increase (from 27 to 40 °C) (Fig. 2b and c). Most of the cyanine dyes (e.g. ICG) have obvious drawbacks such as photobleaching and the loss of heating effect as time increases (Fig. 2c).$^{45,52}$ In contrast, the dramatic absorption spectral shifts of ICG were not found for CDA at the same time. Our result illustrated that CDA-3 was designed as an ideal organic dye with high performing NIR photothermal properties and photostability.$^{28}$

As an NIR organic dye with an excellent photothermal effect, CDA can efficiently generate PA signals under 800 nm pulsed laser irradiation (Fig. 2d). Fig. 2f displayed the PA signal dependence of CDA as a function of concentration. In addition, we compared the PA properties of CDA with three prominent representative probes, ICG, PB and GNRs. At the same concentration, CDA provided the highest PA signal intensity.
that was approximately 1.4, 1.5, and 4.5 times higher than that of ICG, PB and GNR, respectively (Fig. 2e). After exposure to a 1.0 W cm⁻² laser for 10 min, the PA intensity declines for PB and GNRs were 18.0% and 7.3% respectively. Furthermore, the PA amplitude of ICG dramatically decreased by ~80%, illustrating its susceptibility to laser-induced bleaching (Fig. 2g).

Among these four agents, there was a very slight signal decline for CDA (only ~8%), revealing its superior potential for long-term imaging. To study the biocompatibility of CDA-3, an MTT test was performed using human neuronal cells (SH-SY5Y). Fig. 2h showed that CDA-3 exhibited negligible cytotoxicity even at a relatively high concentration of 0.5 mg mL⁻¹. The acute lethal dose (LD₅₀) of CDA-3 via intravenous administration was calculated to be 324.3 mg kg⁻¹ for Kunming mice (n = 8), suggesting an extremely low acute toxicity. The SH-SY5Y cells were then incubated with CDA-3 and fluorescence images were acquired using confocal microscopy (FV1200, Olympus, Japan). Fig. S15† illustrated that CDA-3 can be effectively taken up by cells, indicating the high potential application of this probe for biological imaging.

Next, we explored whether CDA-3 specifically labeled Aβ plaque in cerebral blood vessels. The formaldehyde-fixed brain of WT (wild type) and Tg mice was first stained with anti-Aβ antibody (6E10). As shown in Fig. S16,† the brain sections of Tg mice were easily subjected to labeling by 6E10. Staining Aβ deposits within cerebral blood vessels and parenchyma was observed on brain sections from the Tg mice and AD patient for CDA-3, which was in good agreement with thioflavin-S (Th-S) staining on the same section (Fig. 3a and d). However, there was no apparent labeling in the brain section of WT mice (Fig. S17†). To further examine the influence of CDA-3-positive Aβ plaque on blood vessels, staining was performed using a vascular smooth muscle cell (VSMC) marker (phalloydin-Alexa 594) on the paraffin-embedded brain sections. As displayed in Fig. 3b and c, the Aβ deposits in the blood vessels of Tg mice were clearly labeled with the VSMC marker. VSMCs were distributed discontinuously in these cerebral vessels, which is a typical hallmark of CAA.

Brain grey and white matter are high scattering and absorbing media and pose a great challenge for deep brain imaging. As an NIR fluorescent dye, the penetration depth of CDA-3 was examined with various brain tissue overlays (0, 1, 2, 3 mm) (Fig. 4a–c). As the thickness of the brain tissue increased, the fluorescence intensity decreased significantly but still could be detected at the depth of 3 mm. The quantification of the fluorescence intensity is demonstrated in Fig. 4d. This result suggested that CDA-3 could be used as an excellent probe for brain imaging. To detect the biodistribution of CDA-3 dye in vivo, NIRF imaging was performed after an intravenous (i.v.) injection of 0.1 mg mL⁻¹ of CDA-3 was administered to transgenic mice with five familial Alzheimer’s disease (5 × FAD) and littermate WT mice. As can be seen from Fig. 4e and f, there is an obvious difference in brain accumulation between the 5 × FAD and WT groups. At 4 h postinjection, the 5 × FAD group showed a maximal fluorescence in the brain (Fig. 4g). However, the biodistribution of CDA-3 showed an ignorable brain retention in the WT group, demonstrating that the free dye was quickly eliminated from the body (Fig. 4e).
concentration–time curve of CDA-3 in serum shown in Fig. S18† implies its rapid metabolism in living mice. These results suggested that the high fluorescence signal in the brains of the 5×FAD mice was attributable to the specific binding capacity between the CDA-3 dye and the β-amyloid plaque.

However, in strongly scattering brain tissue, incident light is mainly attenuated by the cranium, which greatly impedes the application of NIRF imaging for locating the β-amyloid plaque. Unlike fluorescence imaging, PA imaging can ultrasonically break the strong optical scattering, and simultaneously retain a high penetration depth and spatial resolution.36 Owing to the excellent PA performance of CDA-3 in vitro, we then studied the in vivo PA imaging in a CAA model. The PA images of the mice brains are shown in Fig. 5 at different time points after the [18F]CDA-3 injection. From the PA imaging results, it is observed that the functionalized dye achieved better accumulation in the CAA model than in the control group at 1 hour post-injection. After 4 hours from administration, the PA signal of the organic dye displayed a significant improvement in brain blood vessels for the Tg group, suggesting high Aβ binding (Fig. 5a). In contrast, the PA signals from the WT group remained almost the same as before (Fig. 5c).

To further highlight the brain region, differential PA imaging was performed by subtracting the raw PA images before the agent injection from the PA images at different time points. As shown in Fig. 5b, an outstanding enhanced PA image contrast of the sagittal sinus was observed in the Tg group by comparing the PA signals before and 4 h after injection. This may be due to the high affinity to Aβ plaque and the gradual accumulation of the dye in brain blood vessels. However, there is no obvious change in the PA signal at different time points for the WT group (Fig. 5d).

Quantitative data was also obtained from the same region of interest (ROI) of brain blood vessels. Specifically, the ratio of the PA signals between the Tg and WT mice [PA(Tg)/PA(WT)] reached 2.1 at 4 h, indicative of specific binding by Aβ plaque (Fig. 5e).

Although PA imaging is a promising technology, whole-body mapping is still a huge challenge in large animals and humans. As the PET technique is widely used for whole-body imaging in clinics, we explored the [18F] based PET method as a complementary imaging technique. After being analyzed using high performance liquid chromatography (HPLC), we injected a 1.85 million becquerel (MBq)/62.5 μg dose of purified [18F]CDA-3 via the tail vein. As can be seen from Fig. 6a, there was a clear distribution of [18F]CDA-3 in the brains of the Tg mice
que between blood vessels and parenchyma. In contrast, owing to the lack of Aβ plaque, there was rarely specific binding in WT mice brains.

To further illustrate this point, study-based analyses for the WT and Tg mice were also carried out using [18F] fluorodeoxyglucose ([18F]FDG) and [18F]AV-45. As expected, the [18F]FDG PET images showed that there was a higher glucose metabolic rate in WT mice than in Tg mice, implying hypometabolism in the brain of AD model. [18F]AV-45, an FDA approved PET probe for AD, also exhibited excellent uptake in the Tg mice. However, the nonspecific diffusion of the [18F]AV-45 probe in the whole brain hinders its further application for differentiating Aβ plaque between blood vessels and parenchyma. In contrast, owing to the amyloid specific tracer, [18F]CDA-3 PET showed approximately a 2-fold higher uptake (0.93% injection dose per gram (% ID per g)) in brain blood vessels of the Tg mice than in the WT group (0.53% ID per g) (Fig. 6b). Since PET has no tissue penetration limitation, 0.9 mm-thick consecutive coronal slices through the whole mouse were obtained 1 h after the [18F]CDA-3 injection was administered (Fig. 6c). However, notable defluorination was found for [18F]CDA-3, tomographic imaging of Tg mice revealed obvious distribution of [18F]CDA-3 in brain in three dimensions, as opposed to low brain accumulation in WT group. Taken together, these results suggest that [18F]CDA-3 is a promising PET probe for the monitoring and early diagnosis of CAA in vivo.

All animal studies were conducted in compliance with the animal care and use guidance established by Xiamen University animal care committee.

3. Conclusions

In summary, this is the first study to evaluate the potential of croconium dye as a building block for PA/PET/NIR fluorescence tri-modality imaging for the diagnosis of brain plaque. Depending on the molecular structure, [18F]CDA has higher selective affinity for cerebrovascular Aβ plaque than neuritic plaque, which does not require an extra antibody or ligand. With its electron donor–π–acceptor–π–donor architecture, CDA exhibits excellent NIR absorption and PA response. Under 760 nm excitation, CDA-3 showed enhanced brain accumulation in the Tg group where the fluorescence intensity in the brain was significantly improved by more than 4 fold compared with the WT group.

Featuring good photostability and photo-thermal properties, CDA-3 generates a higher PA signal than most of the currently reported PA contrasting nanoagents, such as ICG, PB and GNRs, on a per mass basis. These virtues may permit sensitive PA imaging in living mice at a low systemic injection mass and provide a potential means for the photothermal inhibition of amyloid formation. Moreover, CDA-3 exhibited obvious PA improvement in the cerebral vessel other than parenchyma, which makes it possible for the non-invasive amyloid imaging of CAA. Labeled with a radiotracer, [18F]CDA-3 is feasible for CAA-selective PET imaging in vivo. Compared with conventional Aβ imaging tracers, [18F]CDA-3 shows great potential for selectively binding CAA deposits in the brain.

The innovation of this study is to explore the potentials of croconium dye as a PET/PA/NIR fluorescent tri-modality imaging ligand for selective affinity towards brain plaque. With enhanced NIR absorption, effective photothermal property, sensitive PA response, and whole-body PET imaging, CDA-3 plays an important role in monitoring cerebrovascular amyloid and will provide powerful support to develop effective prevention and treatment strategies for CAA and other cerebral blood vessel disorders. By replacing croconic acid, 18F, and thiophene with other functional groups, this unique building block will shed light on investigating the underlying mechanisms of brain disorders.

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References


