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# **Review Paper**

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# Systematic review of curcumin-based optical imaging for amyloid- $\beta$ detection in Alzheimer's disease models

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#### Keywords

Alzheimer's Disease; Amyloid-Beta; Optical Imaging; Curcumin; Fluorescent Probes; Preclinical

### Abstract

**Background:** Alzheimer's disease (AD) is a neurodegenerative disorder characterized by amyloid-beta (A $\beta$ ) plaque accumulation and cognitive decline. Early and precise A $\beta$  detection is vital for effective therapeutic intervention. Curcumin-based fluorescent probes offer high specificity, non-invasive imaging compatibility, and deep tissue penetration, making them promising tools for optical A $\beta$  imaging. This systematic review evaluates preclinical studies on curcumin-based fluorescent probes to assess their photophysical properties, imaging capabilities, and potential applications in detecting A $\beta$  plaques in mouse models of AD.

**Methods:** A comprehensive literature search was performed in PubMed and ScienceDirect (2000-2024). Eligible studies were original English-language articles

using curcumin-based probes for optical imaging of  $A\beta$  in Alzheimer's mouse models. Data extraction focused on imaging parameters such as binding affinity [dissociation constant (Kd)], emission wavelength, quantum yield, fluorescence enhancement, and delivery methods.

**Results:** Thirteen preclinical studies met the inclusion criteria and were analyzed. CRANAD-102 probe showed the highest binding affinity (Kd = 7.5 nM) while CRANAD-3 achieved the most significant fluorescence intensity (39.5-fold). Emission wavelengths averaged 690 nm, with longer wavelengths facilitating deeper tissue imaging. Quantum yields ranged from 0.011 to 0.40, with the highest yield (20.31) observed in CH<sub>2</sub>Cl<sub>2</sub> and effective doses averaging 2.0 mg/kg.

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Innovative delivery methods, such as aerosolized formulations and micelle-based probes, expanded diagnostic applications, including non-invasive retinal imaging.

**Conclusion:** Curcumin-based fluorescent probes exhibit high specificity for  $A\beta$  aggregates, effective deep tissue imaging, and non-invasive delivery potential, making them promising tools for preclinical Alzheimer's diagnostics. However, their clinical translation requires further validation in standardized preclinical and translational studies.

#### Introduction

Alzheimer's disease (AD) is a progressive neurodegenerative disorder marked by cognitive decline, memory loss, and behavioral dysfunction, predominantly affecting the elderly. As the global population ages, AD poses a growing burden due to increasing rates of disability and mortality.<sup>1-3</sup> Despite advances in symptomatic treatment, no curative therapy exists, making early and accurate diagnosis essential for timely intervention and slowing disease progression.

One of the earliest pathological hallmarks of AD is the extracellular accumulation of amyloid-beta (A $\beta$ ) plaques, formed through aberrant cleavage of amyloid precursor protein (APP) by  $\beta$ - and  $\gamma$ -secretases. Notably, this accumulation begins up to two decades before clinical symptoms emerge,<sup>4</sup> highlighting the critical need for sensitive, non-invasive, and scalable diagnostic tools capable of detecting A $\beta$  pathology in the preclinical stages.

Conventional imaging modalities such as positron emission tomography (PET), singlephoton emission computed tomography (SPECT), and magnetic resonance imaging (MRI) have been employed to visualize Aβ burden. However, these approaches are limited by high cost, radiation exposure, limited accessibility, and technical complexity.1 Optical imaging, particularly using near-infrared (NIR) fluorescence probes, offers several advantages including non-invasiveness, high sensitivity, real-time imaging capability, and cost-effectiveness. Ideal NIR probes for AB detection must exhibit high specificity, strong yield, quantum appropriate lipophilicity, long-wavelength emission (> 650 nm), and minimal off-target interactions.<sup>5,6</sup>

Curcumin, a natural polyphenol from Curcuma longa, has emerged as a promising scaffold for  $A\beta$ -targeting fluorescent probes due to its ability to cross the blood-brain barrier (BBB), bind

 $\beta$ -sheet-rich A $\beta$  aggregates, and emit in the visible to NIR range. Numerous curcumin derivatives have been developed to overcome limitations such as poor bioavailability and rapid metabolism, with several demonstrating success in detecting A $\beta$  plaques in the brain and retina of transgenic (Tg) AD mouse models.

Despite promising preclinical data, the clinical translation of curcumin-based probes remains Challenges include species-specific limited. pharmacokinetics, variability in fluorescence performance across formulations, and lack of regulatory standardization. Nonetheless, these probes offer a strong foundation for the design of cost-effective, non-radioactive, and diagnostic tools for AD. To address this translational gap, the present systematic review synthesizes preclinical evidence on curcuminbased fluorescent probes for AB imaging. Specifically, we aim to characterize their photophysical properties, imaging performance, and delivery strategies in Tg AD mouse models.

To our knowledge, this is the first systematic review to integrate photophysical characteristics, in vivo imaging outcomes, and delivery strategies of curcumin-based probes in Tg AD mouse models. Unlike previous reviews, our work provides a comparative synthesis focused on probe structure-function relationships and their in vivo applicability, offering practical insights for the rational design of next-generation diagnostic agents. This review highlights how curcumin derivatives may inform future probe selection and contribute to the refinement of in vivo imaging technologies for early AD diagnosis.

## **Materials and Methods**

This systematic review followed the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) 2020 guidelines to ensure methodological transparency and reporting completeness.

Search strategy: A systematic literature search was conducted using two major biomedical databases – PubMed and ScienceDirect – to identify relevant preclinical studies published between January 2000 and March 2024. In addition to database queries, Google Scholar citation tracking tools were used to identify further eligible studies through backward and forward reference searching. This approach was employed to enhance the breadth of the review by capturing relevant publications that may have been indexed

in other platforms.

The search strategy incorporated a combination of Medical Subject Headings (MeSH) and free-text keywords to ensure comprehensive coverage. The primary search terms included: "Alzheimer Disease" (MeSH) OR "Alzheimer's", "Amyloid beta-Peptides" (MeSH) OR "amyloid- $\beta$ " OR "A $\beta$ ", "Curcumin" (MeSH), and "Optical Imaging" (MeSH) OR "fluorescence imaging". Boolean operators (AND, OR) were applied systematically to combine these terms and optimize retrieval sensitivity and specificity.

The search was restricted to original, peerreviewed articles published in English during the specified period. Only full-text articles were included, while reviews, editorials, conference abstracts, and studies without accessible full texts were excluded.

The complete search strategy is provided in table 1, and the overall article selection process is depicted in figure 1.

*Selection criteria:* Articles were screened for relevance based on their titles and abstracts. The extracted data from the title and abstract had to meet the following criteria:

- 1) The study utilized optical imaging to evaluate and identify bilateral AD,
- 2) Curcumin was employed as a fluorescent probe for detecting  $A\beta$ ,
- 3) The research was conducted using an animal model (mice).

# Inclusion and exclusion criteria

Studies were included if they:

- 1) Utilized mouse models of AD,
- 2) Employed fluorescent imaging techniques for the detection of  $A\beta$  plaques.

Studies were excluded if they:

- 1) Were reviews or non-original research,
- 2) Did not provide full-text access.

Data extraction and analysis: Full-text articles were thoroughly reviewed, and studies that did not address the research question or provided insufficient data were excluded. No studies were excluded due to marginal eligibility; all exclusions were based on clearly defined criteria during full-text screening.

Ultimately, 13 studies were selected for analysis based on their focus on optical imaging in animal models of AD and their exploration of curcumin's effects on  $A\beta$ .

Outcomes and variables of interest: The primary outcomes extracted from each study included: (1) binding affinity to A $\beta$  aggregates [expressed as dissociation constant (Kd)], (2) quantum yield of fluorescence, (3) emission wavelength ( $\lambda$ max), (4) fluorescence signal enhancement, and (5) effective dose in mg/kg.

Additional variables extracted included: (1) physicochemical properties such as partition coefficient (logP), (2) delivery methods [e.g., intravenous (IV) injection, aerosol, micelle-based formulation], (3) cytotoxicity profiles, and (4) imaging targets such as brain cortex or retina. In cases where specific values were not reported, no assumptions were made, and such fields were documented as "not available (NA)".

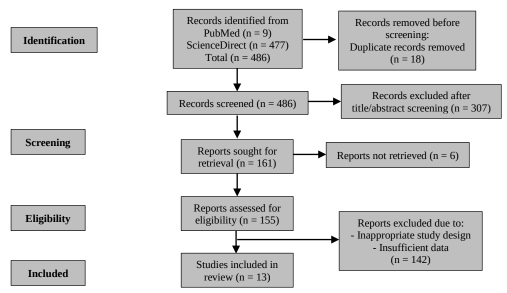
*Effect measures:* For each included study, the primary effect measures extracted were:

- 1) Kd (in nM) to reflect probe binding affinity to  $A\beta$ ,
- 2) Fluorescence signal enhancement ratios [e.g., fold-change vs. control or wild-type (WT)],
- 3) Emission wavelength ( $\lambda$ max, in nm) as a measure of optical penetration potential, and
- 4) Quantum yield as an indicator of fluorescence efficiency.

These metrics were used to qualitatively and quantitatively compare the performance of different curcumin-based probes across studies.

Table 1	1.	Search	Strategy
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Search Strategy					
Databases	PubMed, ScienceDirect				
Time frame	January 2000-March 2024				
Keywords and MeSH	("Alzheimer Disease" [MeSH Terms] OR "Alzheimer" [All Fields] OR "Alzheimer*" [All				
terms used	Fields]) AND ("Amyloid beta-Peptides" [MeSH Terms] OR "amyloid-beta" [All Fields] OR				
	"amyloid-β"[All Fields] OR "Aβ"[All Fields]) AND ("Curcumin"[MeSH Terms] OR				
	"curcumin"[All Fields]) AND ("Optical Imaging"[MeSH Terms] OR "optical imaging"[All				
	Fields] OR "fluorescence imaging"[All Fields])				
Search methodology	Boolean operators (AND, OR) were systematically applied to combine the search terms and				
	ensure both sensitivity and specificity in retrieval. Filters were applied to include only peer-				
	reviewed original articles published in English between 2000 and 2024. Additionally,				
	citation tracking (both backward and forward) was used to identify further relevant studies				
through reference lists of included articles.					



**Figure 1.** Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) flowchart summarizing the search strategy and study selection process

**Data preparation:** No data conversions or transformations were required, as all relevant outcomes were directly reported in the included studies.

Synthesis of results: A narrative synthesis approach was employed to compare and interpret findings across studies. No statistical pooling or meta-analysis was performed. No subgroup analysis or formal assessment of heterogeneity was conducted, as the review was not designed for quantitative synthesis. Sensitivity analyses were not applicable given the descriptive nature of the synthesis.

Quality assessment: To ensure methodological rigor and transparency, the risk of bias for each included preclinical study was systematically assessed using SYRCLE's Risk of Bias Tool, a domain-based instrument specifically developed for animal research. The assessment covered five key domains: (1) random sequence generation (selection bias), (2) allocation concealment (selection bias), (3) blinding of investigators and outcome assessors (performance and detection bias), (4) completeness of outcome data (attrition bias), and (5) selective outcome reporting (reporting bias).

Two independent reviewers conducted the assessments, and any disagreements were resolved through discussion and consensus. Risk of bias judgments were applied across all included studies and summarized narratively in the manuscript. Studies with a high overall risk of bias were excluded

to ensure the internal validity of the findings.

No formal assessment of certainty or confidence in the body of evidence (e.g., using GRADE) was performed, given the descriptive nature of the synthesis.

#### Results

A total of 13 studies employing curcumin-based fluorescent probes for optical imaging of  $A\beta$  plaques in AD were reviewed. These studies evaluated photophysical properties, in vivo fluorescence measurements, and  $A\beta$  specificity in Tg mouse models. Key data are summarized in table 2.

Probe 9 (Fang et al.²) showed a 1.5-fold higher fluorescence intensity in APP/presenilin 1 (PS1) mice compared to WT [emission wavelengths: monomers = 690 nm, oligomers = 688 nm, aggregates = 697 nm; binding constants: monomers =  $11.16 \pm 0.79 \text{ nM}$ , oligomers =  $36.59 \pm 2.69 \text{ nM}$ , aggregates =  $14.57 \pm 1.27 \text{ nM}$ ; quantum yield = 20.31 (in CH<sub>2</sub>Cl<sub>2</sub>)]. Fluorescence signal increased with mouse age.

CAQ (Wu et al.¹) in 5×FAD mice showed 1.57-fold higher fluorescence versus WT at 30 minutes post-injection (binding constant = 78.89 nM, logP = 3.08, quantum yield = 0.011).

Analog 8b (Park et al.<sup>5</sup>) exhibited 2.26-fold stronger fluorescence in  $5\times FAD$  mice versus WT at 10 minutes (emission  $\lambda = 667$  nm, > 20-fold enhancement post-A $\beta$  binding). Quantum yield was not reported.

**Table 2.** Summary of photophysical properties and fluorescence imaging characteristics of curcumin-based probes in the reviewed studies (Part I)

Study	Dose (mg/kg)	Quantum yield	LogP	Binding constant (Kd, nM)	Mouse model	Imaging technique
Wu et al. <sup>1</sup>	0.15	0.01	3.08	78.89	5×FAD Tg (10 months, female) WT (10 months, female)	- In vivo/ex vivo fluorescent imaging - FIHC
Fang et al. <sup>2</sup>	1.00	20.31 in CH <sub>2</sub> Cl <sub>2</sub>	2.14	Aβ oligomers: $36.59 \pm 2.69$ Aβ aggregates: $14.57 \pm 1.27$ Aβ monomers: $11.16 \pm 0.79$	APP/PS1 Tg (14 months, male) WT (14 months, male)	<ul> <li>In vitro fluorescent staining</li> <li>In vivo NIRF imaging</li> <li>Upright fluorescence microscopic imaging</li> </ul>
Sidiqi et al. <sup>3</sup>	0.75	N/A	N/A	N/A	APP/PS1 Tg (6-18 months) WT (6-18 months)	-IHC -Whole-mount immunofluorescence - In vivo retinal fluorescence imaging with curcumin
Park et al. <sup>5</sup>	0.40	N/A	N/A	$91.20 \pm 3.28$	5×FAD Tg (15 months, female) WT (15 months, female)	<ul><li>In vivo NIRF imaging</li><li>Ex vivo and histological imaging</li></ul>
Ni et al. <sup>6</sup>	2.00	N/A	N/A	N/A	arcAβ Tg (18-24 months, both sexes) WT (18-24 months, both sexes)	- MSOT - Hybrid vMSOT-fluorescence imaging - MRI IHC
Si et al. <sup>7</sup>	3.00	0.37	N/A	1360.00	APP/PS1 Tg (17 months)	- In vivo NIRF imaging
Ran et al.8	5.00	0.40	3.00	38.00	2576 Tg (19 months) WT (19 months)	- Fluorescence intensity-based NIR imaging
Chibhabha et al. <sup>9</sup>	N/A	N/A	N/A	N/A	APPswe/PS1ΔE9 Tg C57BL	- Ex-vivo double immunofluorescence staining - In vivo imaging
Li et al. <sup>10</sup>	2.00	0.01	N/A	Soluble A $\beta$ : $7.50 \pm 10.00$	APP/PS1 Tg (five months, female)	- In vivo NIRF imaging
Zhang et al. <sup>11</sup>	0.50	N/A	2.50	Insoluble A $\beta$ : 505.90 ± 275.90 A $\beta$ monomers: 24.00 ± 5.70 Dimers: 23.00 ± 1.60 Oligomers: 16.00 ± 6.70 Aggregates: 27.00 ± 15.80	WT (five months, female) APP/PS1 Tg (14 months, female) WT (14 months, female)	- Two-photon imaging - In vivo NIRF imaging
Zhang et al. <sup>12</sup>	2.00	N/A	1.94	Aβ40: 105.80 Aβ42: 45.80	APP/PS1 Tg (4 months, female) WT (4 months, female)	- In vivo NIR imaging
Koronyo-	7.50	N/A	N/A	N/A	APPswe/PS1ΔE9 Tg	- IHC
Hamaoui et al. <sup>13</sup>					(18 months, female) WT (18 months, female)	- In vivo fluorescence imaging
McClure et al. <sup>14</sup>	5.00	N/A	N/A	N/A	5×FAD Tg (8 months) C57BL/6 WT (8 months)	- Ex vivo fluorescence brain imaging - IHC

Table 2. Summary of photophysical properties and fluorescence imaging characteristics of curcumin-based probes in the reviewed studies (Part II)

Wu et al.   Small animal live imager   Text   Te	Study	nmary of photophysical properties and fluorescence imaging of photophysical instrument	Emission wavelength	Probe	Brief result
Fang et al. 2					
Strong binding to Aβ aggregates					
Fang et al.					
et al. <sup>2</sup> - Olympus VS200 microscope - Aggregates: 697 nm	Fang	- Leica TCS SP8 CLSM	Oligomers: 688 nm	Probe 9	
Sidiqi		- Olympus VS200 microscope			
et al. 3 - Fluorescence scanning laser ophthalmoscopy (fSLO) Park et al. 5 - Maestro 2.0 in vivo imaging system et al. 5 - CCD camera  Ni et al. 6 - Confocal microscope - A hybrid vMSOT-fluorescence imaging system Si et al. 7 - IVIS Lumina system  Ran et al. 8 - Fluorescence reflectance (also known as epifluorescence) and tomography (FMT)  Chibhabh a et al. 9 - Carl Zeiss LSM 780 - Micron IV retinal imaging microscope Li et al. 10 - IVIS spectrum animal imaging system  Zhang - Spectrum animal imaging system  Zhang - Spectrum animal imaging system  Amax: 635 nm  Amax: 635 nm  Amax: 635 nm  Amax: 635 nm  Dye 2 - CRANAD-2  Slower fluorescence 2 hours post-injection, high specificity for Aβ deposits, low cytotoxicity  Slower fluorescence decay in Tg2576 mice vs. WT, higher signal intensities at multiple time points  Labeled retinal Aβ plaques, robust deposition in retina and hippocampus, low toxicity  The control of the properties of the control of the plane of the control of the plane of the control of the plane of the control		- IVIS Lumina XR III animal imaging system	Aβ Monomers: 690 nm		affinity, low cytotoxicity
Park et al. 5 - Maestro 2.0 in vivo imaging system et al. 5 - CCD camera	Sidiqi	- Confound fluorescent imaging	N/A	Curcumin	Higher cortical and retinal Aβ signal intensity
et al. 5  Order of the tal. 6  Ni et al. 6  Ni et al. 6  Si et al. 7  Fluorescence reflectance (also known as epifluorescence) and tomography (FMT)  a et al. 9  A et al. 9  A et al. 9  A cranada curcumin polymeric micelles  Li et al. 10  A loss pectrum animal imaging system  A round 730 nm  Et al. 11  Apply 1  Apa affinity, NIR emission  Dose-dependent signal intensity increase, significant AUC in cortical regions of arcAβ mice  Peak fluorescence 2 hours post-injection, high specificity for Aβ deposits, low eytotoxicity  Slower fluorescence decay in Tg2576 mice vs. WT, higher signal intensities at multiple time points  Labeled retinal Aβ plaques, robust deposition in retina and hippocampus, low toxicity in retina and hippocampus, low toxicity or soluble Aβ species  Zhang  - IVIS spectrum animal imaging system  Around 730 nm  CRANAD-3  CRANAD-3  CRANAD-58  1.3-fold higher NIRF signal intensity in Tg mice vs.  Aβ species and age groups  CRANAD-58  Labeled retinal Aβ plaques, robust deposition in retina and hippocampus, low toxicity mice vs. WT at 30 minutes, selective for soluble Aβ species  CRANAD-102  Around 730 nm  CRANAD-58  1.3-fold higher NIRF signal intensity in Tg mice vs.	et al. <sup>3</sup>	- Fluorescence scanning laser ophthalmoscopy (fSLO)			in Tg mice, correlating with cortical Aβ loads
Ni et al. 6  - Confocal microscope - A hybrid vMSOT-fluorescence imaging system  Si et al. 7  - IVIS Lumina system  Amax: 635 nm  Dye 2  Peak fluorescence 2 hours post-injection, high specificity for Aβ deposits, low cytotoxicity Fan et al. 8  - Fluorescence reflectance (also known as epifluorescence) and tomography (FMT)  a et al. 9  - Carl Zeiss LSM 780  Li et al. 10  - IVIS spectrum animal imaging system  Around 730 nm  CRANAD-3  Aβ affinity, NIR emission  Dose-dependent signal intensity increase, significant AUC in cortical regions of arcAβ mice  Peak fluorescence 2 hours post-injection, high specificity for Aβ deposits, low cytotoxicity Slower fluorescence decay in Tg2576 mice vs. WT, higher signal intensities at multiple time points  Labeled retinal Aβ plaques, robust deposition in retina and hippocampus, low toxicity in retina and hippocampus, low toxicity  I 1.22-fold higher signal intensity in APP/PS1 mice vs. WT at 30 minutes, selective for soluble Aβ species  Zhang  - IVIS spectrum animal imaging system  Around 730 nm  CRANAD-3  Apax ~ 750 nm  CRANAD-58  Labeled retinal Aβ plaques, robust deposition in retina and hippocampus, low toxicity  and plaques, robust deposition in retina and hippocampus, low toxicity  and plaques, robust deposition in retina and hippocampus, low toxicity  and plaques, robust deposition in retina and hippocampus, low toxicity  and plaques, robust deposition in retina and hippocampus, low toxicity  and plaques, robust deposition in retina and hippocampus, low toxicity  and plaques, robust deposition in retina and hippocampus, low toxicity  and plaques, robust deposition in retina and hippocampus, low toxicity  and plaques, robust deposition in retina and hippocampus, low toxicity  and plaques, robust deposition in retina and hippocampus, low toxicity  and plaques, robust deposition in retina and hippocampus, low toxicity  and plaques, robust deposition in retina and hippocampus, low toxicity  and plaques, robust deposition in retina and hippocampus, low toxicity  and plaques,	Park	<ul> <li>Maestro 2.0 in vivo imaging system</li> </ul>	λmax: 667 nm	8b curcumin analog	2.26-fold signal intensity enhancement in
Ni et al. 6  - Confocal microscope - A hybrid vMSOT-fluorescence imaging system  Si et al. 7  - IVIS Lumina system  Amax: 635 nm  Dye 2  Peak fluorescence 2 hours post-injection, high specificity for Aβ deposits, low cytotoxicity for Aβ deposits, low cytotoxicity specificity for Aβ deposits, low	et al. <sup>5</sup>	- CCD camera			5×FAD mice vs. WT at 10 minutes, high
- A hybrid vMSOT-fluorescence imaging system  Si et al. <sup>7</sup> - IVIS Lumina system  \[ \lambda \text{Amax: 635 nm} \\ \text{Peak fluorescence 2 hours post-injection, high specificity for A\beta deposits, low cytotoxicity specificity for A\beta deposition in retinal A\beta places, robust deposition in retina and hippocampus, low toxicity in retina and hippocampus, low toxicity specificity for soluble A\beta species  \[ \text{Li et al.}^{10} \] - IVIS spectrum animal imaging system  \[ \text{Around 730 nm} \] - IVIS spectrum animal imaging sys					
Si et al. 7 - IVIS Lumina system	Ni et al. <sup>6</sup>		N/A	CRANAD-2	
Si et al. 7 - IVIS Lumina system		<ul> <li>A hybrid vMSOT-fluorescence imaging system</li> </ul>			
Ran et al. <sup>8</sup> - Fluorescence reflectance (also known as epifluorescence) and tomography (FMT)  Chibhabh a et al. <sup>9</sup> - Carl Zeiss LSM 780 - Micron IV retinal imaging microscope  Li et al. <sup>10</sup> - IVIS spectrum animal imaging system  Zhang - Spectrum animal imaging system  Chibhabh - CLSM - Carl Zeiss LSM 780 - IVIS spectrum animal imaging system  Around 730 nm  Et al. <sup>11</sup> - Spectrum animal imaging system  Around 730 nm  CRANAD-102  Slower fluorescence decay in Tg2576 mice vs. WT, higher signal intensities at multiple time points  Labeled retinal Aβ plaques, robust deposition in retina and hippocampus, low toxicity  in retina and hippocampus,					
Ran et al. <sup>8</sup> - Fluorescence reflectance (also known as epifluorescence) and tomography (FMT)  Chibhabh a et al. <sup>9</sup> - Carl Zeiss LSM 780 - Micron IV retinal imaging microscope  Li et al. <sup>10</sup> - IVIS spectrum animal imaging system  Zhang - IVIS spectrum animal imaging system  The species and age groups  Zhang - Spectrum animal imaging system  Slower fluorescence decay in Tg2576 mice vs. WT, higher signal intensities at multiple time points  Labeled retinal Aβ plaques, robust deposition in retina and hippocampus, low toxicity  In retina and hippocampus, low toxicity  Slower fluorescence decay in Tg2576 mice vs. WT, higher signal intensities at multiple time points  Labeled retinal Aβ plaques, robust deposition in retina and hippocampus, low toxicity  In retina and hippocampus, low toxicity mice vs. WT at 30 minutes, selective for soluble Aβ species  Zhang - IVIS spectrum animal imaging system  Around 730 nm  CRANAD-3  CRANAD-3  2.29-fold higher signal intensity in APP/PS1 mice vs. WT, effective across  Aβ species and age groups  Amax ~ 750 nm  CRANAD-58  1.3-fold higher signal intensity in Tg mice vs.	Si et al.	- IVIS Lumina system	λmax: 635 nm	Dye 2	
epifluorescence) and tomography (FMT)  Chibhabh a et al. 9  Carl Zeiss LSM 780  - Carl Zeiss LSM 780  - Micron IV retinal imaging microscope  Li et al. 10					
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Li et al. 10  - IVIS spectrum animal imaging system  N/A  CRANAD-102  1.22-fold higher signal intensity in APP/PS1 mice vs. WT at 30 minutes, selective for soluble Aβ species  Zhang  - IVIS spectrum animal imaging system  et al. 11  Around 730 nm  CRANAD-3  CRANAD-3  2.29-fold higher NIRF signal intensity in APP/PS1 mice vs. WT, effective across Aβ species and age groups  Zhang  - Spectrum animal imaging system  λmax ~ 750 nm  CRANAD-58  1.3-fold higher signal intensity in Tg mice vs.	a et al.				in retina and nippocampus, low toxicity
mice vs. WT at 30 minutes, selective for soluble $Aβ$ species  Zhang - IVIS spectrum animal imaging system et al. 11  Around 730 nm  CRANAD-3  2.29-fold higher NIRF signal intensity in APP/PS1 mice vs. WT, effective across $Aβ$ species and age groups  Zhang - Spectrum animal imaging system $λmax \sim 750$ nm  CRANAD-58  1.3-fold higher signal intensity in Tg mice vs.	Li at al 10		NI/A		1.22 fold higher signal intensity in ADD/DC1
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Zhang - Spectrum animal imaging system $\lambda \text{max} \sim 750 \text{ nm}$ CRANAD-58 1.3-fold higher signal intensity in Tg mice vs.	ct ai.				
	7hang	- Spectrum animal imaging system	λmax ~ 750 nm	CRANAD-58	
or at a month, low our target interactions	et al <sup>12</sup>		Miles 750 IIII	CRITIVID 30	
Koronyo Micron II retinal imaging microscope N/A Curcumin Retinal Aβ signal intensity correlates with		12111	N/A	Curcumin	
Hamaoui - Carl Zeiss Axio Imager Z1 fluorescence microscope disease progression in Tg mice.	•		- 1/	001 0011111	
et al. 13 - Leica TCS SP5 double-spectral confocal microscope					F
McClure - Fluorescence tomographic imaging system N/A FMeC1 FMeC1 binds amyloid plaques expressed in			N/A	FMeC1	FMeC1 binds amyloid plaques expressed in
et al. <sup>14</sup> - Dual channel fluorescence microscopy the hippocampal areas and cortex.					

Data are presented as mean  $\pm$  standard deviation (SD)

Kd: Dissociation constant; N/A: Not available; NIR: Near-infrared; Tg: Transgenic; WT: Wild-type; FIHC: Fluorescence immunohistochemistry; MSOT: Multispectral optoacoustic tomography; IHC: Immunohistochemistry; MRI: Magnetic resonance imaging; NIRF: Near-infrared fluorescence; TEM: Transmission electron microscopy; CLSM: Confocal laser scanning microscope; LSM: Laser scanning microscope; FMT: Fluorescence molecular tomography; CCD: Charge-coupled device; FMeC1: F-methyl-curcumin-1; APP/PS1: Amyloid precursor protein/presenilin 1; AUC: Area under the curve; Aβ: Amyloid-beta; vMSOT: Volumetric multispectral optoacoustic tomography; DMSO: Dimethyl sulfoxide

CRANAD-2 showed dose-dependent fluorescence for  $A\beta_{1-42}$  fibrils ( $r^2 = 0.991$ ). For  $A\beta_{40}$  aggregates, fluorescence increased 70-fold (logP = 3.0).

CRANAD-102 had binding constants of  $7.5 \pm 10.0 \, \text{nM}$  (soluble A $\beta$ ) and  $505.9 \pm 275.9 \, \text{nM}$  (insoluble). In APP/PS1 mice, fluorescence was 1.22-fold higher than WT at 30 minutes.

1,2-distearoyl-sn-glycero-3-phosphoethanolamine-N-[methoxy(polyethylene glycol)-2000] (DSPE-PEG2000) (DSPE-PEG2000) micelles labeled retinal plaques; hemolytic activity was reported as 1.79%.

Dye 2 reached peak fluorescence at 2 hours post-injection (quantum yield = 0.37, cell viability > 85%).

CRANAD-3 showed 12.3-39.5-fold fluorescence increases across A $\beta$  species (2.29-fold higher in APP/PS1 vs. WT, emission  $\approx$  730 nm).

CRANAD-58 (Zhang et al.<sup>12</sup>): Binding constants:  $A\beta_{40} = 105.8 \text{ nM}, A\beta_{42} = 45.8 \text{ nM}, \log P = 1.94.$ 

F-methyl-curcumin-1 (FMeC1) (McClure et al.<sup>14</sup>): Aerosol delivery targeted hippocampal/cortical plaques; fluorescence data were not quantified.

Other studies (Sidiqi et al.³, Koronyo-Hamaoui et al.¹³) reported retinal and cortical fluorescence signals in Tg models, with retinal plaques detectable at  $\geq$  2.5 months.

# **Summary metrics**

- Binding constants (Kd): Range = 7.5-1360 nM, mean ≈ 40.8 nM
- Quantum yields: Range = 0.011-20.31, mean ≈ 4.22. This observed variability is influenced by experimental factors such as solvent polarity, aggregation state of the probes, and the specific photophysical environment, which are not always standardized across studies.
- Emission wavelengths: Range = 635-760 nm, mean ≈ 690 nm
- Doses: Range = 0.15-7.5 mg/kg, mean  $\approx 2.0$  mg/kg

Noticeable variability may relate to solvent systems, probe aggregation, and delivery methods.

Risk of bias assessment: All studies were assessed using SYRCLE's Risk of Bias Tool. Domains related to random sequence generation, allocation concealment, random housing, and blinding were rated as "unclear" due to insufficient reporting. Remaining domains (baseline comparability, outcome completeness, selective reporting, and other biases) uniformly showed "low risk". No study was rated "high risk" in any domain.

#### Discussion

Developing non-invasive imaging tools for detecting  $A\beta$  deposits with high resolution is essential for understanding AD mechanisms and advancing  $A\beta$ -targeted therapies. Fluorescence imaging has been the primary application for most  $A\beta$  probes, including NIAD-4, AOI-987, boron-dipyrromethene (BODIPY), THK-265, Donor-acceptor near-infrared fluorophores (DANIR), luminescent conjugated oligothiophenes (LCOs), (E)-4-(4-(dibutylamino)styryl)-1-(2-hydroxyethyl quinolin-1-ium chloride (DBA-SLOH), and curcumin-derived CRANAD series.<sup>2</sup>

Curcumin, a natural compound capable of crossing the BBB, has shown promise in staining senile plaques and modulating amyloid pathology in Tg AD models. However, its use in NIR imaging has been limited by short emission wavelengths, low water solubility, poor BBB permeability, and rapid metabolic degradation. Strategies such as borate complexation have successfully induced red shifts in emission spectra, thereby improving imaging performance.<sup>5</sup>

This review highlights significant variability complementarity among the curcumin-based fluorescent probes. CRANAD-102 exhibited the highest binding affinity for soluble A $\beta$  species (Kd = 7.5 ± 10.0 nM), suggesting potential for detecting early pathological changes, whereas CAQ, despite lower affinity (Kd = 78.89 nM), offered advantages in safety and in vivo stability due to its low cytotoxicity and favorable logP (3.08).<sup>1,10</sup> In terms of fluorescence efficiency, probe 9 demonstrated the highest quantum yield (20.31 in CH<sub>2</sub>Cl<sub>2</sub>), supporting its use in high-signal imaging applications.<sup>2</sup> However, CRANAD-3 provided the broadest fluorescence enhancement (12.3-39.5-fold) across A $\beta$  species and ages, and its NIR emission (~ 730 nm) made it suitable for deeper tissue imaging.11 While analog 8b reached > 20-fold enhancement upon Aβ binding and performed well in rapid-onset models like 5×FAD mice,5 CRANAD-2 stood out for its excellent BBB permeability (logP = 3.0) and dose-dependent signal increases.6,8

terms of delivery and practicality, DSPE-PEG2000 micelle formulations improved solubility and biocompatibility, making them viable for retinal imaging,9 while aerosolized FMeC1 enabled non-invasive midbrain imaging – an advantage over IV delivery routes.14 These findings illustrate specific how probe modifications red-shifting, polymeric (e.g.,

encapsulation, or solubility enhancement) can optimize imaging for different diagnostic contexts, from early detection to longitudinal monitoring.

Innovative delivery systems such as aerosolized curcumin derivatives and micellebased formulations have expanded the diagnostic potential of these probes. FMeC1 delivered via aerosol enabled midbrain imaging without IV injection, while DSPE-PEG2000 micelles correlated well with retinal A $\beta$  load, enabling non-invasive early diagnostics.

Several studies also emphasized age-dependent imaging: retinal Aβ plaques were detectable as early as 2.5 months in Tg AD mice, aligning with early pathological events involving soluble Aβ species.3 Importantly, variability in Tg mouse models may influence the apparent efficacy of these probes. For instance, 5×FAD mice develop dense and aggressive A $\beta$  pathology by 2-3 months of age, providing a more robust imaging target, whereas APP/PS1 mice exhibit slower and more regionally restricted plaque development beginning around 5-6 months. These distinctions affect probe binding dynamics, fluorescence intensity, and temporal imaging windows, and should be considered when interpreting crossstudy differences in diagnostic performance.

In relation to established clinical approaches, curcumin-based fluorescent probes offer several advantages over Food and Drug Administration (FDA)-approved Aβ PET tracers, such as florbetapir and flutemetamol. These include lower cost, absence of radioactivity, potential for realtime and longitudinal imaging, and even retinalbased detection. However, they are limited by shorter tissue penetration depth, formulationdependent variability, rapid metabolism, and the lack of regulatory standardization. PET tracers benefit from validated pharmacokinetics, reproducibility, and broad clinical acceptance. Therefore, while curcumin probes show promise for early-stage or point-of-care diagnostics, their clinical utility will depend on further optimization, including enhancements in bioavailability, safety, and standardized imaging protocols.

Limitations: While curcumin-based fluorescent probes show considerable promise, a number of methodological and technical limitations inherent in the existing studies must be acknowledged. The included studies exhibited substantial methodological heterogeneity, including variations in probe structure, administration routes, imaging equipment, and

animal models with differing AB pathology profiles. For instance, 5×FAD mice develop early and aggressive plaque accumulation, whereas APP/PS1 models display more gradual and regionally restricted pathology, complicating cross-study comparisons. Sample sizes were generally small, and few studies conducted direct, head-to-head comparisons under standardized conditions, limiting the ability to benchmark probe performance systematically. Moreover, long-term biodistribution, and accumulation were seldom evaluated, leaving gaps in the safety profile of these probes. Curcumin-based agents also face intrinsic technical challenges, such as photobleaching, poor aqueous solubility, rapid metabolic degradation, and formulation-dependent variability, all of which may compromise imaging consistency and translational reliability. These limitations highlight need for further standardization and optimization to ensure the safe and effective clinical application of curcumin-based optical imaging strategies.

While risk of bias was assessed at the individual study level using the SYRCLE's Risk of Bias Tool, no formal assessment of reporting bias was performed at the synthesis level due to the descriptive, non-quantitative nature of this review. Likewise, the certainty of the evidence was not evaluated using formal frameworks such as GRADE. Finally, no review protocol was registered prior to the conduct of this systematic review, and therefore no protocol amendments were applicable.

Future directions and clinical translation: Although current evidence from preclinical studies is encouraging, the clinical applicability of curcumin-based fluorescent probes remains challenges such as rapid constrained by metabolism, limited solubility, and formulation variability. This review helps bridge the translational gap by identifying probes with favorable photophysical profiles and highlighting delivery strategies – such as polyethylene glycol (PEG) ylated micelles and liposomal systems – that enhance bioavailability and imaging performance. Importantly, our findings inform rational probe selection based on quantitative metrics (e.g., binding affinity, emission wavelengths, quantum yields) and model-dependent outcomes, which are critical for advancing in vivo imaging technologies.

In support of clinical relevance, multiple translational studies have demonstrated the

feasibility of curcumin-based probes in human contexts. Liu et al. reported that a bivalent curcumin-cholesterol ligand (BMAOI-14) crossed the BBB and specifically stained Aß plagues in both Tg mouse models and human AD brain tissue. 15 More recently, An et al. developed a quinolinecurcumin-based probe [quinoline-derived halfcurcumin-dioxaborine (Q-OB)] that sensitively detected A<sub>β</sub> oligomers in the cerebrospinal fluid (CSF) of patients with AD, distinguishing disease stages through fluorescence signal variation.<sup>16</sup> Complementing these findings, den Haan et al. showed that curcumin and its clinically bioavailable derivatives selectively bound fibrillar Aβ in post-mortem AD brains, with minimal crossneurodegenerative reactivity other pathologies.<sup>17</sup> Collectively, these studies reinforce the translational promise of curcumin derivatives for in vivo diagnostics. Future directions include expanding cross-species validation, regulatory harmonization, and integration with artificial intelligence (AI)-assisted image analysis to optimize signal quantification and clinical applicability. These efforts will be critical for advancing curcumin-based probes toward safe

and effective use in early AD detection.

#### Conclusion

Curcumin-based fluorescent probes show promise for preclinical applications in AD research, particularly due to their specificity for AB aggregates, potential for deep tissue imaging, and compatibility with non-invasive delivery methods. However, these findings are derived from heterogeneous preclinical studies with notable limitations, including variability in experimental protocols, probe formulations, and animal models, as well as limited assessment of long-term safety and clinical applicability. As such, while the reviewed evidence highlights their potential utility in early detection and disease monitoring, further standardized investigations – including validation in large animal or human models - are needed before these probes can be considered for clinical translation.

#### **Conflict of Interests**

The authors declare no conflict of interest in this study.

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None.

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