

Current Anti-Amyloid- β Therapy for Alzheimer's Disease Treatment: From Clinical Research to Nanomedicine

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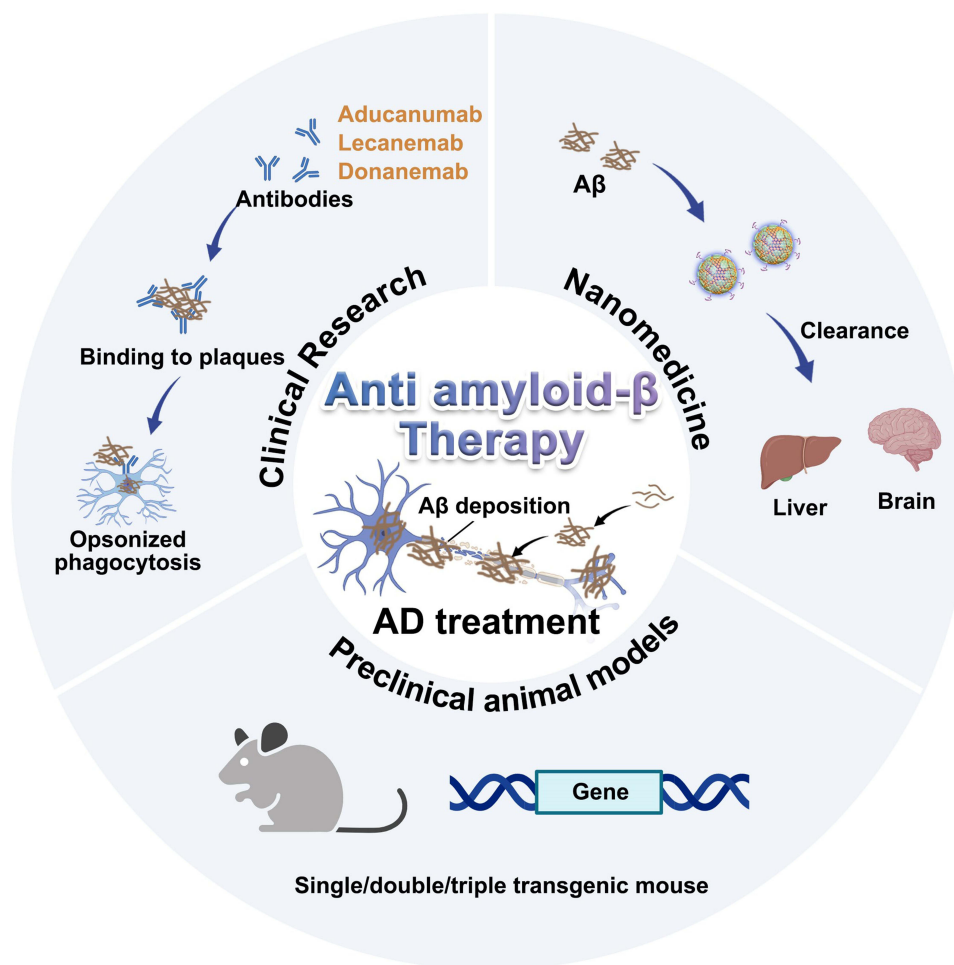
Abstract: Recent successive approval of anti-amyloid- β (A β) monoclonal antibodies as disease-modifying therapies against Alzheimer's disease (AD) has raised great confidence in the development of anti-AD therapies; however, the current therapies still face the dilemma of significant adverse reactions and limited effects. In this review, we summarized the therapeutic characteristics of the approved anti-A β immunotherapies and dialectically analyzed the gains and losses from clinical trials. The review further proposed the reasonable selection of animal models in preclinical studies from the perspective of different animal models of A β deposition and deals in-depth with the recent advances of exploring preclinical nanomedical application in A β targeted therapy, aiming to provide a reliable systematic summary for the development of novel anti-A β therapies. Collectively, this review comprehensively dissects the pioneering work of A β -targeted therapies and proposed perspective insight into AD-modified therapies.

Keywords: Alzheimer's disease, amyloid- β , anti-amyloid- β monoclonal antibody, disease-modifying therapeutics, nanomedicine

Introduction

Alzheimer's disease (AD) is the most common type of dementia causing tens of millions of elderlies suffering from cognitive decline and behavioral impairment.¹ Although it has been identified for more than 100 years, the cognitive deterioration of AD has remained frustratingly to develop disease-modifying therapeutics despite massive efforts. Most of the current research has been philosophically guided by the connection of the hallmark histopathology of AD, including the senile plaques (SP) formed by deposited amyloid- β (A β) and neurofibrillary tangles (NFTs) formed by hyperphosphorylated microtubule-associated protein tau. Before the first approval of A β antibody (aducanumab) in 2021 as the first disease-modifying therapy,² the Food and Drug Administration (FDA) had only approved a handful of drugs to relieve the condition, including acetylcholine esterase (AChE) inhibitors and N-methyl-D-aspartic acid (NMDA) receptor modulators. This class of symptomatic remission drugs has been reported to be gradually discontinued by patients due to serious adverse reactions and weak disease remission,³ thus it is not hard to understand the encouraging success of the groundbreaking approval of aducanumab for confronting AD. Based on clarifying the shared role of A β deposition in AD as the clearest instance of cross-talk between neurodegenerative and cerebrovascular processes, lecanumab, the next generation of A β antibodies, was approved two years later with overwhelming support. These achievements have, to some extent, cleared the cloud in the development of AD therapies and represent the effectiveness of therapeutic strategies with A β as the mainstream intervention target.⁴ In this context, there is great potential to extend the entire life cycle of A β production and clearance metabolism and to develop advanced anti-AD therapies. Therefore,

Graphical Abstract



there is extremely necessary to systematically review the regulation of A β metabolism and clarify the key to the development of therapeutic strategies targeting A β .

The cerebral A β is the hydrolysate of amyloid precursor protein (APP) via proteolysis of β -/ γ - secretase and is cleaved into 39–43 length peptides according to the different cutting sites (eg, A β_{1-40} or A β_{1-42}).⁵ The extracellular A β monomers gradually aggregate to form soluble oligomers, fibrils and fibers until insoluble plaques and eventually lead to neuronal degeneration.⁶ The subsequent catabolism of A β was mediated by central or peripheral pathway to obtain the homeostasis of A β , including the glial immunophagocytosis in the central or direct draining by the circulatory system to the periphery.^{7–9} Abundant clues have suggested that the imbalance between the production and removal of anabolism and catabolism of A β is the main culprit of amyloid plaques.¹⁰ Therefore, therapies that block the production of A β or mobilize its clearance are the eternal theme of A β -targeting strategies, such as the current A β monoclonal antibodies which are specific IgG1s binding to A β with affinity and complete the degradation of A β by promoting cell recognition and phagocytosis. In addition, researchers have developed a series of small-molecule drugs, nucleic acid drugs, proteins/peptides, and others to regulate the level of enzyme activity of A β secretion or to mobilize the clearance and degradation of A β at the molecular biological level (Figure 1).¹¹

However, such treatment strategies are not without pitfalls. For example, amyloid-related imaging abnormalities (ARIA), the most controversial adverse effect of A β antibodies in clinical anti-A β immunotherapy, warn of the malignant events caused by excessive central burden in the process of A β clearance.¹² The development of nanotechnology and

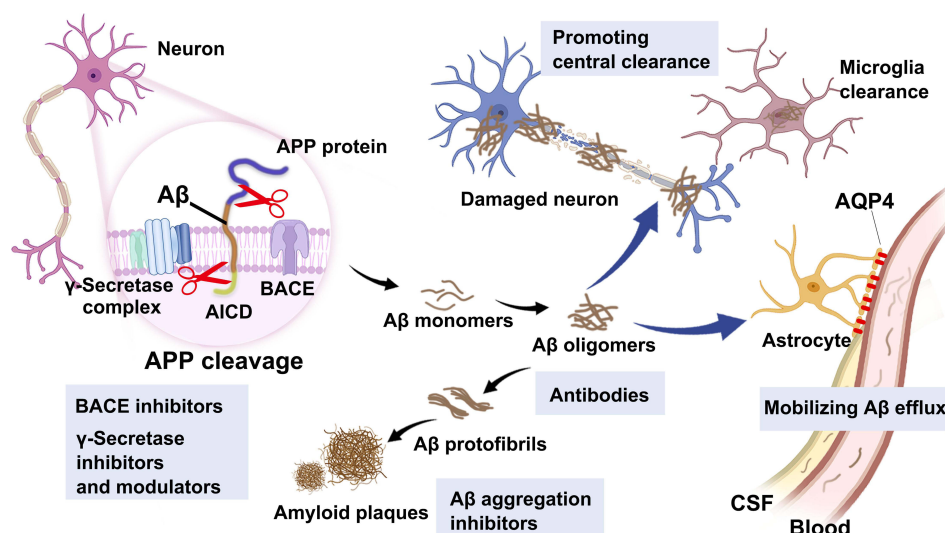


Figure 1 Illustration of the process of A β production, aggregation, and clearance in central and peripheral. APP is cleaved by β - and γ -secretase on neuron cells to release A β monomers, following self-aggregated into oligomers and fibrils to form plaques. Cerebral A β can be degraded by microglia phagocytosis or transported out of the brain for degradation.

targeted drug delivery systems in recent years has effectively facilitated the development of novel AD therapeutic strategies. In the past decade or so, nanomedicine for AD treatment has shown a continuous development process along with the progress of disease cognition. Previous nanomedicine focused on high-potency drug encapsulation to reduce peripheral adverse reactions of clinical medication or promoting central delivery by utilizing the advantageous size or functionalized surface to achieve the blood-brain barrier crossing, thus promoting the effective enrichment of potential anti-AD active drugs in the lesion.^{13,14} In recent years, as the elaboration of more detailed crosstalk relationship of AD pathogenic targets, the application of nanocarriers to efficiently support multiple therapeutic components has become a mainstream, especially the diverse drug carrying potential of the nanocarriers exhibits great advantages in the combinational delivery of biological macromolecules and active compounds. Meanwhile, with the successive approval of A β monoclonal antibodies, more studies have focused on the limitations of current antibody therapies, and a variety of synthetic nanoparticles have recently been demonstrated to act as highly efficient A β “chaperones” to regulate A β aggregation or facilitate clearance, attempting to provide alternatives to A β antibodies.^{15,16} Furthermore, the specific transport and enrichment rules of nanomedicine in vivo provide carrier tools for the systemic clearance of A β to realize the metabolism of A β outside the brain to achieve a safer and more efficient A β clearance pattern. Therefore, the rational application of nanomedicine strategies to reasonably design and construct AD-adapted A β targeted regulation of active components or delivery platforms to achieve safe and efficient A β clearance is the key challenge for the clinical transformation of A β nanomedicine. Finally, the success rate of drug clinical transformation is closely related to the selection of preclinical animal models. Therefore, it is essential to understand the complexity, advantages, and limitations of various animal models, as well as the irrational use, in order to enhance preclinical research on translational therapy for AD. In this review, we start from the gain and loss of clinical A β antibodies and elaborate on the key of targeted A β therapy. Following the introduction of applicable animal models in preclinical anti-A β studies, we have explored the endogenous clearance pathway of A β as well as the corresponding current AD modification strategies, aiming to provide a reliable approach for the development of A β -targeted therapies.

Literature Search Strategy

An electronic search was conducted using the PubMed, Clinical Trail, and Scopus databases, supplemented with a manual search using various combinations of Medical Subject Headings (MeSH) terms, including Alzheimer’s disease, amyloid- β , antibody, nanomedicine, and transgene mice model. The inclusion criteria for the search were as follows: articles published between 2010 and 2023, English as the language of publication, studies concerning Alzheimer’s

disease, clinical studies, and nanomedicine. In order to include relevant studies on classic AD transgenic animal models, the search scope for this part was expanded to include articles published from January 1990 to September 2023. The exclusion criteria for the search included opinion papers, case reports, duplicate articles, and studies on biosensors or in vivo diagnostic reagents. For included articles, it should help answer at least one research question. Thus, the inclusion criteria developed relate to what the document covers:

1. Analysis of the application of A β clinical therapy and the causes of adverse reactions;
2. Construction method, characteristics and applicable scope of AD animal model;
3. The application advantages of nanomedicine for AD compared with insufficient clinical treatment;
4. Interaction between A β and other AD pathogenic targets, and the consideration of multi-drug combination.

Immunotherapies with Anti-A β Monoclonal Antibodies in AD Patients

The development of amyloid pathology in AD involves a series of steps, starting with the aggregation of A β monomers, followed by the formation of dimer/trimers, oligomers, protofibrils, fibrils, and ultimately the deposition of insoluble amyloid plaques.¹⁷ A β monoclonal antibodies (mAbs) can target different stages of this process and facilitate the clearance of A β .¹⁸ Recently, the FDA has approved two mAbs, aducanumab and lecanemab. Another promising mAb, donanemab, has submitted a marketing application to the FDA and received priority review by the Center for Drug Evaluation (CDE) in China in November 2023. The following sections provide a summary and discussion of the therapeutic mechanisms and clinical advancements of these antibodies, along with other mAbs that show promise for future approval.

Aducanumab

Aducanumab, the mAb designed to target residues 3–7 in the A β N-terminus, obtained accelerated approval from the FDA in June 2021, marking a significant milestone in AD treatment.¹⁹ Aducanumab exhibits low monovalent affinity but strong avidity for epitope-rich aggregates, allowing it to selectively bind to A β oligomeric or fibrillar aggregates.²⁰ This binding activates microglia, leading to the clearance of A β from the brain.

The Phase I clinical trial (NCT01397539) revealed that patients generally tolerated doses up to 30 mg/kg well, with no occurrence of severe adverse events. A subsequent phase Ib clinical trial (PRIME, NCT01677572) further demonstrated that a dose range of 3 to 10 mg/kg of aducanumab reduced amyloid plaque and was safe enough to be used in the Phase III trials for efficacy.²¹ Based on the results of the PRIME trial, two clinical trials, EMERGE (NCT02484547) and ENGAGE (NCT02477800) trials, were initiated in 2015, both of which were phase III studies conducted over 18 months in individuals with prodromal and mild AD.²² Unfortunately, these two trials and an additional Phase II trial (EVOLVE, NCT03639987) that was designed to assess safety were prematurely ended following an interim futility analysis of phase III trial.²³ In October 2019, Biogen announced that the interim futility analysis was incorrect. An analysis of a larger dataset showed that the high-dose aducanumab in the EMERGE trial met its primary endpoint, though the low-dose aducanumab did not exhibit significant benefits compared to the placebo group. Besides, while the ENGAGE trial did not meet its primary endpoint, it was observed that individuals who received 10 or more 10 mg/kg doses in the ENGAGE trial showed a slower decline similar to the participants in the EMERGE trial. The disparity in outcomes between the EMERGE and ENGAGE trials can be attributed to differences in the duration of exposure to high-dose aducanumab.

According to safety data from the EMERGE and ENGAGE trials, about a third of aducanumab-treated participants developed amyloid-related imaging abnormalities (ARIA), of which a quarter were symptomatic and 3% were severe.²³ ARIA-E (vasogenic edema) and ARIA-H (microhemorrhages and hemosiderosis) were commonly observed in patients receiving a dosage of 10 mg/kg, with a 19.1% and 35% incidence rate, respectively. Besides, APOE ϵ 4 carriers had a higher incidence compared to noncarriers.²² Notably, ARIA is a common phenomenon associated with anti-A β immunotherapies.^{24,25} This may be due to the antibody promoting the disaggregation of amyloid plaques into soluble forms when it binds with them, triggering glial cell-mediated neuroinflammation. At the same time, some soluble A β may enter blood vessels and deposit, leading to vascular pathology (Figure 2).

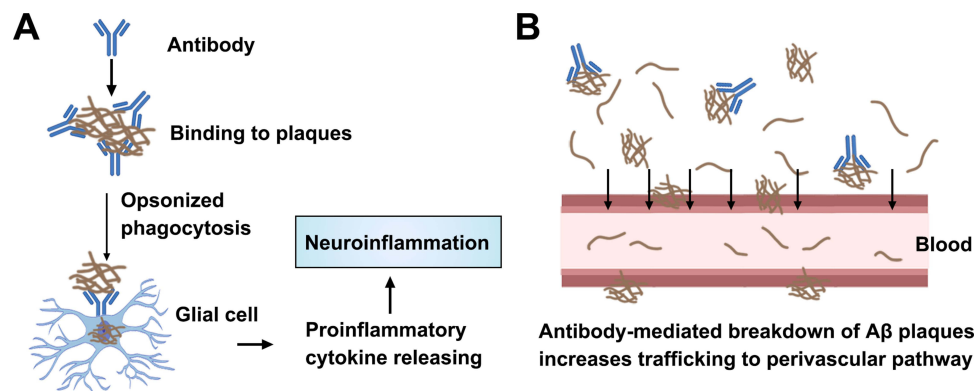


Figure 2 The process of antibody-mediated A β clearance and production of vascular amyloid pathology. **(A)** Antibodies recognize and bind to amyloid plaques, prompting their transformation into a soluble form, which then promotes phagocytosis of microglia, during which microglia are activated and neuroinflammation occurs. **(B)** Soluble A β flows to the blood vessels and deposits, resulting in vascular pathology.

Despite controversies, aducanumab received accelerated approval from the FDA based on the surrogate endpoint of A β reduction, representing a significant milestone in the field of AD therapeutics. It signifies the culmination of almost 25 years of research into the amyloid hypothesis and ushers in a new era for AD treatment, bringing hope to patients and researchers. However, the Centers for Medicare & Medicaid Services (CMS) currently limits coverage for aducanumab outside of approved clinical trials, reducing its accessibility in clinical settings. Moreover, it should be noted that further post-approval studies are still needed to confirm its clinical benefits.

Lecanemab

Lecanemab, a humanized IgG1 antibody derived from the mouse antibody mAb158, received accelerated approval by the FDA in January 2023 and full approval in July 2023 for the treatment of mild cognitive impairment (MCI) or mild dementia associated with AD.²⁶ Lecanemab possesses an exceptional affinity for selectively binding to soluble A β aggregates (including oligomers and protofibrils), approximately 1000 times greater than A β monomers and 10–15 times than insoluble fibrils.²⁷

The Phase I clinical trial (NCT01230853) of lecanemab conducted on 80 patients with mild-to-moderate AD, evaluated its safety, tolerability, and pharmacokinetics. Lecanemab demonstrated good tolerability at all tested doses, up to 10 mg/kg administered every two weeks for four months. The incidence of amyloid-related imaging abnormalities (ARIA-E, ARIA-H) was similar to placebo. Besides, pharmacokinetic analysis showed dose-proportional response when lecanemab dose was ≥ 10 mg/kg, with a serum elimination half-life of 7 days.²⁸ Thereafter, a Phase IIb trial (NCT01767311) enrolled 856 subjects to determine the dosage and efficacy of the treatment. At the 12-month mark, the trial did not reach its primary endpoint or futility criteria. Hence, the trial continued until the 18-month mark. Results showed that patients receiving lecanemab at a dosage of 10 mg/kg twice a month experienced a statistically significant reduction in cognitive decline at 6, 12, and 18 months. This dosage led to a 93% reduction in brain amyloid plaques, a 47% slower decline in cognitive function as measured by ADAS-Cog, and a 30% slower decline in cognitive function as measured by ADCOMS.²⁹

The phase III CLARITY AD trial (NCT03887455) enrolled 1795 participants with early AD, who were randomly assigned to receive either placebo or intravenous lecanemab at a dosage of 10 mg/kg every two weeks.³⁰ The randomization process took into account clinical subgroups (MCI or mild AD) and concomitant use of approved AD symptomatic medication. Assessing clinical decline using CDR-SB as the primary endpoint, the lecanemab group showed a substantial reduction in clinical decline. Furthermore, lecanemab exhibited a highly statistically significant decrease in clinical decline on the global cognitive and functional scale at 18 months, with reduced amyloid plaque burden, a 24% deceleration in disease progression, and a 37% deceleration in decline in activities of daily living. Aside from ARIA-H and ARIA-E, infusion-related reactions are the most commonly reported adverse events, of which the

majority reactions tended to be mild to moderate, with most occurring (75%) after the first dose. These results provide valuable insights into the efficacy and safety profile of lecanemab as a potential treatment for early AD.

Collectively, lecanemab stands out as a highly promising therapeutic option for AD due to its ability to reduce brain A β levels, mitigate cognitive decline, and exhibit a lower incidence of ARIA-E, thus ensuring improved safety. However, despite its moderate therapeutic effect and enhanced safety profile, lecanemab failed to demonstrate clinically significant benefits in patients with clinically manifest or prodromal dementia. Further research and investigation are essential to explore the efficacy and safety of lecanemab thoroughly.

Donanemab

Donanemab is a humanized IgG1 antibody designed to specifically target a pyroglutamated form of A β (N3pG-A β) that is aggregated in amyloid plaques.³¹ This truncated form of N-terminal A β is highly toxic and hydrophobic, leading to faster aggregation with β -sheet stabilization and resistance to degradation. The specificity of donanemab allows it to effectively bind these stable plaques, overcoming the issue of soluble A β forms that can impede recognition of deposited forms.³² Notably, donanemab has received priority review by CDE in November 2023 and is expected to become the third mAb approved by the FDA in the near future.

The Phase Ia clinical trial (NCT01837641) involved six different doses ranging from 0.3 to 10 mg/kg administered intravenously, and one group receiving donanemab through subcutaneous injection.³³ The results showed that monthly intravenous administration of donanemab at a dose of 10 mg/kg reduced A β plaques by approximately 40–50%. The Phase Ib clinical trial (NCT02624778) enrolled 61 participants and used intravenous single or multiple dosing of donanemab at doses ranging from 10 to 40 mg/kg.³⁴ All dosing cohorts showed a reduction in brain A β at 72 weeks. In a phase II trial (TRAILBLAZER-ALZ, NCT03367403) involving 272 patients with early symptomatic AD and confirmed tau and amyloid deposition, participants were randomly assigned to receive either donanemab or placebo intravenously every 4 weeks for up to 72 weeks. The primary outcome, measured by the change in the Integrated Alzheimer's Disease Rating Scale (iADRS) score at 76 weeks, suggested that donanemab possessed a positive effect on cognitive and functional impairment. Donanemab also showed greater reductions in amyloid plaque levels and global tau load compared to placebo at 76 weeks. Notably, nearly complete clearance of A β plaques was observed within 6 months.³⁵ In a Phase III trial, TRAILBLAZER-ALZ 2 (NCT04437511), 1736 patients were randomly assigned to either the donanemab treatment group (860 patients) or the placebo control group (876 patients). Over a 72-week period, patients received donanemab or placebo treatment every four weeks. At the 76-week mark, donanemab treatment showed a 35.1% reduction in disease progression risk compared to placebo in early AD patients with low/medium tau pathology, as measured by iADRS scores. Additionally, among early AD patients with low/medium tau pathology, 47% remained stable after one year of donanemab treatment, while only 29% of those receiving placebo treatment remained stable.

Donanemab was generally well tolerated but was reported to be associated with the occurrence of ARIA, with approximately 40% of participants in donanemab groups experiencing ARIA, and about 26.1% of them showing symptomatic ARIA. In contrast, symptomatic ARIA-E was observed in only 0.8% of participants receiving placebo.³⁶ These data led to the FDA granting donanemab a breakthrough therapy designation and form the basis of donanemab's application for traditional approval shortly thereafter.

Other Anti-A β mAb Candidates Under Clinical Trials

Remternetug, a humanized mAb developed by Eli Lilly, specifically targets N3pG-A β and represents a next-generation targeted therapy for AD. In July 2022, a global Phase III clinical trial (NCT05463731) was initiated to evaluate the safety, tolerability, and efficacy of Remternetug in early-stage AD patients. In 2023, Eli Lilly commenced an international multicenter Phase III trial (CTR20230785) employing a double-blind, randomized, placebo-controlled, parallel design to assess its safety and efficacy through subcutaneous injections in individuals with early AD symptoms. Additionally, the positive results of a Phase I clinical trial (NCT04451408) have been announced and indicated the efficacy of Remternetug in amyloid protein clearance.

Trontinemab is a humanized mAb as an improved version of the gantenerumab developed by Genentech. Trontinemab combines the gantenerumab with a protein domain that can bind to transferrin receptors, aiming to enhance the

antibody's ability to cross the BBB in to the brain.³⁷ Trontinemab possesses 50-fold higher BBB penetration and A β plaque binding efficiency compared to unmodified gantenerumab. Currently, Trontinemab is undergoing Phase II clinical trials (NCT04639050) for the treatment of mild-to-moderate AD. Preliminary results indicate that trontinemab can facilitate rapid amyloid plaque clearance at substantially lower doses than typical amyloid-targeting antibodies. After 28 weeks of treatment with the maximum dose of 1.8 mg/kg, 75% of patients exhibited a reduction of amyloid levels to below normal detectable limits.

ABBV-916 is a mAb developed by AbbVie, which recognizes the N3pG-A β aggregated in amyloid plaques. The preclinical data for this antibody has not been publicly disclosed yet. The clinical development of ABBV-916 began in August 2022, and the trial was initially listed as a phase I/II trial with an enrollment of 288 participants. It was later changed to a phase II trial (NCT05291234) with a target enrollment of 195 participants. This trial is randomized, placebo-controlled, double-blind, and conducted across multiple centers, which is being conducted in 88 centers across North America, Europe, and Japan, with an expected completion date of December 2024.

ACU193, a humanized mAb developed by Acumen, is the first A β oligomers-selective immunotherapeutic undergoing human clinical trials for the treatment of early-stage AD, aiming to investigate the A β oligomer hypothesis in AD.³⁸ In clinical settings, the binding of anti-A β mAbs to amyloid plaques has been associated with a higher incidence of ARIA.³⁹ ACU193 has a higher specificity for A β oligomers and shows minimal or no binding to amyloid plaques. Therefore, ACU193 is predicted to carry a lower risk of ARIA compared to other A β antibodies. In October 2022, ACU193 was granted Fast Track designation by the FDA for the treatment of MCI or early-stage AD. A randomized, placebo-controlled, double-blind Phase I clinical trial (NCT04931459) for ACU193 is complete.⁴⁰ Data showed significant reductions in amyloid plaque levels as detectable by brain amyloid PET scans in patients receiving higher doses of ACU193 after 6–12 weeks. Besides, ACU193 demonstrated favorable tolerability and safety profiles, with 10.4% of patients experiencing ARIA-E, of which 2.1% were symptomatic. A Phase II clinical trial (NCT05291234) investigating the safety, efficacy, pharmacokinetics, and pharmacodynamics of ABBV-916 for the treatment of early-stage AD is currently recruiting participants.

In summary, A β monoclonal antibodies show promise in targeting A β pathology in AD (Table 1), despite challenges including disease stage limitations, target specificity, blood–brain barrier penetration,⁴¹ adverse effects,⁴² and the need for combination therapies.⁴³ Ongoing research and advancements offer potential solutions to these challenges. The current immunotherapy approach for lowering amyloid levels in AD is inadequate in providing a comprehensive treatment. Consequently, there is a persistent quest for novel therapeutic options to address the complex nature of AD pathology.

Table 1 The Therapeutic Mechanisms and Clinical Effect of Anti-A β mAbs

mAbs	Manufacturer	Types of A β Binding	Indication	Clinical Status
Aducanumab	Eisai, Biogen	Oligomeric A β or fibrillar aggregates	Mild AD	FDA accelerated approval
Lecanemab	Eisai, Biogen	Soluble A β protofibrils	MCI or mild AD	FDA full approval
Donanemab	Eli Lilly	N3pG-A β A β plaques	Early symptomatic AD	Submit the application for market approval to the FDA; Receive priority review by CDE
Remternetug	Eli Lilly	N3pG-A β A β plaques	Early-stage AD	Phase III
Trontinemab	Genentech	A β oligomers, fibers, and plaques	Mild to moderate AD	Phase II
ABBV-916	AbbVie	N3pG-A β A β plaques	Early-stage AD	Phase II
ACU193	Acumen	A β oligomers	MCI or Early-stage AD	Phase I

Transgenic Mouse Models in Preclinical Studies

Despite concerted efforts from both academia and industry, a potent curative intervention for AD remains elusive.⁴⁴ The high failure rate observed in clinical drug development is significantly related to the choice of animal models used in preclinical studies that demonstrate only limited aspects of AD pathology.^{45,46} Hence, understanding the intricacies, advantages, and limitations of each model while evaluating AD therapies across multiple models represents a crucial step in enhancing the translation of treatments from preclinical studies to patients. For that, we succinctly delineate several AD mouse models based on A β pathology, including mutation sites, pathological characteristics, advantages, disadvantages and applications, to provide reference and guiding significance for fundamental AD investigations (Table 2).

Single Transgenic APP Mouse Model

The utilization of the APP single transgenic mouse emerges as a prominent animal model in AD research. This paradigm involves the integration of a mutated variant of the human APP gene within the murine genome, thus orchestrating an emulation of the aberrant A β deposition within the cerebral milieu.⁴⁴ These mutations can be those associated with familial AD, such as the Swedish (K670N/M671L) mutation, which results in abnormal production and accumulation of A β .⁵⁸ APP single transgenic murine models predominantly manifest an augmented accrual of A β , a cardinal pathological hallmark intricately linked to AD. In the cerebral milieu of these murine counterparts, A β progressively accumulates, orchestrating the emergence of amyloid plaques that potentially precipitate neuronal dysfunction and nerve impairment.^{59,60} These encompass deficits in learning and memory faculties, impediments in spatial orientation, and alterations in socio-behavioral tendencies. Concomitantly, within the milieu of APP single transgenic murine paradigms, the incipient onset of cellular detriment mechanisms becomes discernible, notably encompassing neuroinflammation and oxidative stress.⁴⁷ These deleterious processes engender an intricate interplay that might further erode neuronal viability and functional efficacy.⁶¹ Analogous to the symptomatic manifestation of AD within human patients, these quintessential features proffer these murine models as a formidable armamentarium for delving into the realms of cognitive impairment etiology within the AD spectrum.

Although the APP single transgenic AD mouse model is important for the study of AD, it also has some shortcomings. AD, in its entirety, engenders a labyrinthine neurodegenerative pathos intrinsically underpinned by a myriad of pathological signatures, spanning neurofibrillary tangles, inflammatory cascades, and neuronal attrition. Thus, mimicking A β accumulation alone may not fully reflect multiple aspects of AD.⁶² In addition, APP single transgenic mouse models may be deficient in neurofibrillary tangles. Neurofibrillary tangles are strongly associated with abnormal Tau aggregation, which may not be fully simulated in a single transgenic model. However, Mesquita et al discovered that the single transgenic APP mouse (J20) has more pronounced lymphatic defects when investigating the connection between meningeal lymphatic vessels and cognitive dysfunction, making them a more suitable choice as a preclinical research model.⁶³ Consequently, a judicious amalgamation of diverse models and methodological approaches is imperative to yield a panorama both comprehensive and veracious in its essence.

Double Transgenic APP/PS1 Mouse Model

APP/PS1 double transgenic mice, an animal model widely used to study AD, are unique in that they harbor mutations in both the human APP and Presenilin 1 (PS1) genes, thus mimicking multiple AD-related pathological features.⁶⁴ Specifically, the mutations within the APP genes give rise to an abnormal A β production trajectory, while those in the PS1 genes, associated with familial AD, engender dysfunction within the γ -secretase complex, thereby engendering perturbations in A β metabolism and aggregation kinetics.⁵⁷ Notably, APP/PS1 double transgenic mice typically manifest heightened cognitive and behavioral aberrations compared to their single transgenic counterparts, displaying potential anomalies in domains such as learning, memory, spatial cognition, and socio-behavioral tendencies. These deviations align more closely with the clinical symptomatic profile exhibited by human AD patients. APP/PS1 double transgenic mice are commonly used to evaluate various drug therapies. Because these mice mimic multiple AD pathological features, researchers can test drug effects on various aspects of A β accumulation, cognitive function, and more. Oakley et al invented 5 $\times$ FAD mice with five FAD mutations in APP/PS1 double transgenic mice (APP_{K670N/M671L/V717I/I716V} Tg

Table 2 Comparison of Current APP Mouse Models of AD

Category	Model	Mutation(s)	Genetic background	Plaques	Pathological Features	Disadvantages	Applications
Single transgenic APP-Tg	PDAPP	APP ^{V717F}	C57B6×DBA2	8 M: Cx, HIP ^{47,48}	Weaker cognitive and behavioral deficits; A β Deposition; Neuronal degeneration and death in some models	Single pathological features; Random integration of transgenes; Overexpression of APP fragments; No complete negative control mice were available; Mixed genetic backgrounds; No neurofibrillary tangle formation.	Pathological mechanism research; Evaluation and screening of drugs targeting A β ; The biomarkers for early diagnosis and monitoring of AD; The effects of A β plaques on neuronal and cellular functions were investigated.
	Tg2576	APP ^{K670N, M671L}	B6; SJL mixed background	7–8 M: focal plaques in Cx ⁴⁹ 11–13 M: Cx, HIP, CB ⁵⁰			
	APP23	APP ^{K670N, M671L}	C57BL/6	6 M: rare plaques in Cx and SUB ⁵¹ 12 M: Cx, HIP, THAL, AMYG ⁵² Majority fibrillar			
	J20	APP ^{K670N, M671L, V717F}	C57BL/6	5–7 M: diffuse plaques in HIP 10 M: plaques HIP, Cx ⁵³			
Double transgenic APP/PS1-Tg	APP/PS1	APP ^{K670N, M671L} PSEN1 ^{L166P}	C57BL/6J	1.5 M: Cx 3 M: Cx, HIP 3–5 M: THAL, Striatum, Brain stem Majority fibrillar ⁵⁴	Obvious cognitive and behavioral deficits; Larger A β plaques formation; Neuronal degeneration and death in some models	Single pathological features in some cases; Random integration of transgenes; Overexpression of APP fragments; No complete negative control mice were available; Mixed genetic backgrounds; neurofibrillary tangle formation.	Pathological mechanism research; Evaluation and screening of drugs targeting A β ; The biomarkers for early diagnosis and monitoring of AD; The effects of A β plaques on neuronal and cellular functions were investigated; Early diagnostic markers of AD.
	APP/PS1	APP ^{K670N, M671L} PSEN1 ^{M146L}	C57BL/6J	6 M: Cx, HIP, all fibrillar 12 M: THAL, Striatum ⁵⁵ brain stem, all diffuse			
	5×FAD	APP ^{K670N, M671L} APP ^{V717I} APP ^{I716V} PSEN1 ^{M146L} PSEN1 ^{L286V}	(C57BL/6×SJL) F1 and C57BL/6J	2 M: SUB, Cx With age: HIP, OB, THAL, brain stem, none in CB ⁵⁶			
Triple transgenic	3×Tg-AD	APP ^{K670N, M671L} PSEN1 ^{M146V} MAPT ^{P301L}	C57BL/6J	6–12 M: Cx, HIP Diffuse and compact ⁵⁷	More pronounced cognitive deficits; neurofibrillary tangle formation; Significant Neuronal loss	Random integration of transgenes; Overexpression of APP fragments; No complete negative control mice were available; Mixed genetic backgrounds; Cognitive impairment before A β accumulation.	Pathological mechanism research; Evaluation and screening of drugs targeting A β and Tau; Identification of CSF biomarkers; Analysis of cell death.

Abbreviations: HIP, hippocampus; Cx, cortex; CB, cerebellum; WM, white matter; THAL, thalamus.

and PSEN1^{M146L/L286V}).⁴⁴ The “5×FAD” mouse model was designed to introduce multiple genetic mutations into a single animal model to more fully mimic multiple pathological features of AD, especially Aβ plaque formation. Locci et al observed that the percentage of area occupied by plaques was significantly higher in 5xFAD and APP/PS1 mice compared to APP^{NLGF} mice. Despite having the lowest Aβ plaque load, APP^{NLGF} mice exhibit robust memory deficits, anxiety and depression-like behaviors, and impaired social behaviors, and are an effective preclinical model to study AD neuropathology, memory deficits, and affective behavioral changes.⁶⁵ Therefore, it is essential to better understand the advantages and disadvantages of each model and to select the appropriate model for preclinical studies.

Triple Transgenic AD Mouse Model

This model configuration encompasses a triad of mutant genes, namely APP, PS1, and tau, thereby facilitating the manifestation of NFTs induced by Aβ deposition and perturbed tau phosphorylation, rendering it more consonant with the intricate spectrum of AD pathology.⁶⁶ The 3×Tg-AD murine model has been meticulously devised to emulate a constellation of pathological features characteristic of AD, encompassing the genesis of Aβ plaques, the atypical aggregation of tau proteins, and the ensuing neurofibrillary tangle formations.^{57,67} This elaborate architecture permits a more comprehensive emulation of the multifaceted pathological panorama intrinsic to AD, thus furnishing an invaluable substrate for the explication of pathogenic mechanisms and the formulation of therapeutic modalities within the AD domain.

Some models may not fully mimic the complexity of the pathological development of AD in humans; Aβ deposition is not the only pathological feature of AD, and abnormal aggregation of tau protein is also closely related to the pathogenesis of AD.^{68,69} Therefore, rational selection of models, combining the advantages of multiple animal models, will contribute to a more comprehensive understanding of the pathogenesis of AD and the search for more effective treatment strategies. Taken together, when choosing the appropriate model, we should combine the research purpose and problem, and make full use of the advantages of different models to better promote the basic research of AD and provide a scientific basis and guidance for the treatment strategy of AD.

Exploration in Aβ-Targeted Preclinical Nanomedicine

The Intervention of Aβ Production

Though various interpretations including elevated tau phosphorylation, neuroinflammation, and so on have been proposed, the Aβ plaque is still a conspicuous pathological character and compelling cause for AD evolved.⁷⁰ Especially, the imbalance between accumulation and removal of Aβ has been considered to contribute to accumulation a lot.⁷¹ Hence, intervention of Aβ production from the source would seem to be a promising way to decrease intracerebral Aβ. There are two main enzymes, beta-site amyloid precursor protein cleaving enzyme 1 (BACE1) following γ-secretase, cleave the transmembrane APP and achieve amyloidogenic cleavage (Figure 3A). Of them, the BACE1 plays a rate-limiting role in reducing Aβ production.⁷² The high enzymatic activity of BACE1 corresponding to the high levels of Aβ reconfirmed its vital position in the AD process. Given that numerous inhibitors have been extensively developed to lower the Aβ concentrations in patients. Due to the similar structure of BACE1 with other aspartyl proteases, and extensive location in the brain, high selectivity as well as high BBB penetration efficiency to implement minimal off-target side effects are crucial for inhibitor design.

Strategies to Inhibit BACE1

Over the past few decades, many pharmaceutical companies have invested time and effort in developing the BACE1 inhibitor pipeline, and 13 small-molecule inhibitors have been introduced into the clinic. Unfortunately, early clinical trial results did not support the BACE1 inhibitor, leading to its cessation of production. BACE1 inhibitors can be divided into two categories named peptidomimetic and non-peptidomimetic based on structure. The former class is characterized by an amide bond or its isosteres. Among them, OM99-2 is the earliest BACE1 inhibitor designed by mimicking the APP cleavage site sequence, firstly demonstrating a potent inhibitory activity of BACE1 and providing much insight into the development of subsequent inhibitors.⁷⁵ However, due to the peptide-based analogs, the poor brain permeability, short half-life leading to rapid clearance, and low bioabsorption limit the in vivo therapeutic effect.⁷⁶ Nonpeptidomimetic

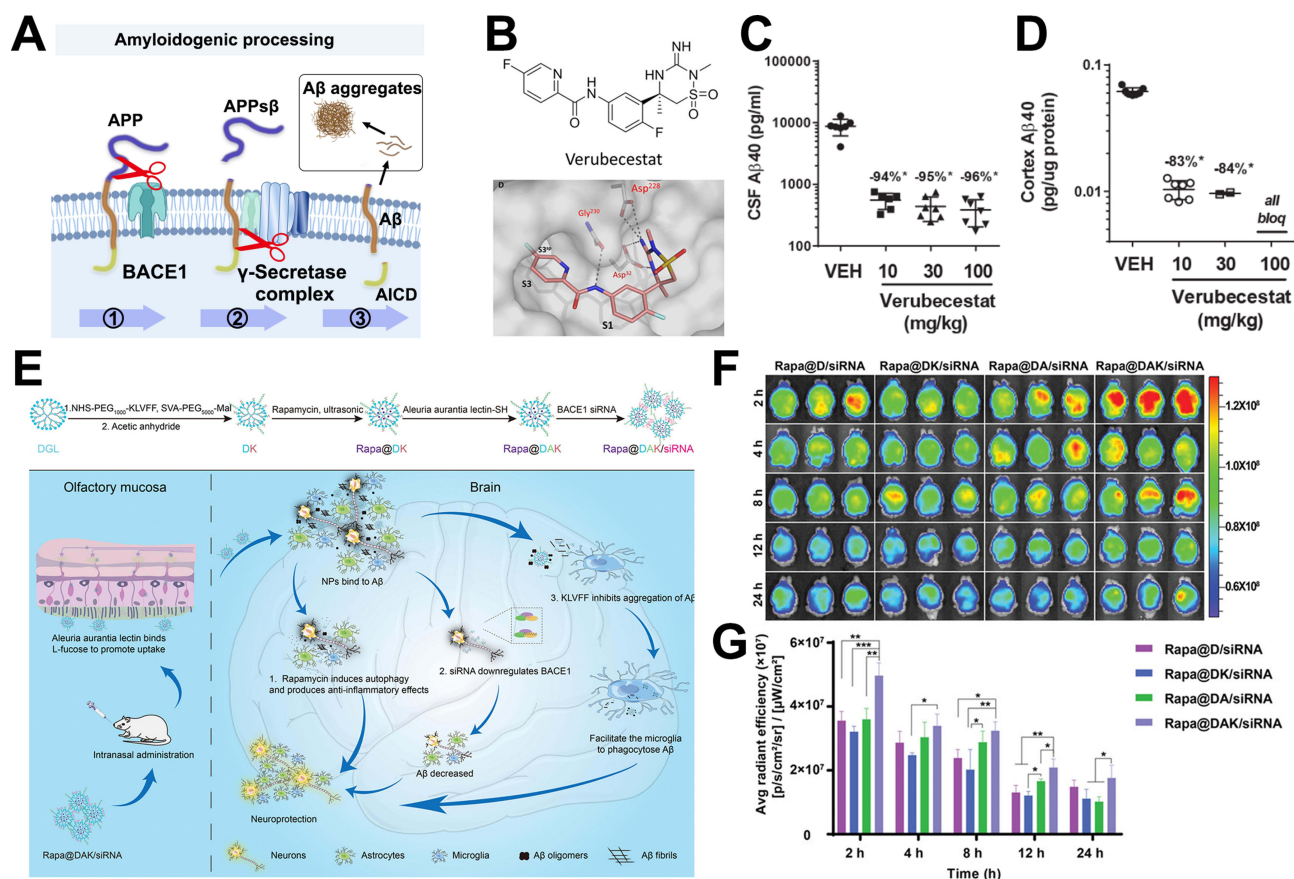


Figure 3 Inhibition of BACE1 is an effective AD treatment strategy to reduce A β production considering the rate-limiting role of BACE1. **(A)** A β is released from the APP after being cleaved by BACE1 and γ -secretases, subsequently self-aggregating into oligomers and fibrils to form plaques. **(B)** The chemical formula and structure-based drug design of verubecestat. **(C)** CSF levels of A β ₁₋₄₀ were reduced. **(D)** Cortex levels of A β ₁₋₄₀ were reduced. Reproduced with permission from Kennedy ME, Stamford AV, Chen X, et al. The BACE1 inhibitor verubecestat (MK-8931) reduces CNS β -amyloid in animal models and in Alzheimer's disease patients. *Sci Transl Med.* 2016;8:363ra150. ⁷³ Copyright 2016, The American Association for the Advancement of Science. **(E)** Nanoparticles (Rapa@DAK/siRNA) for nasal administration and achieving BACE1 silence. **(F)** Ex vivo imaging of the brain tissue of AD model mice after intranasal administration with different preparations at series time points. **(G)** The semiquantitative analysis data of fluorescence intensity from **(F)**. *p < 0.05 and **p < 0.01 and ***p < 0.001. Reproduced with permission from Yang X, Yang W, Xia X, et al. Intranasal Delivery of BACE1 siRNA and Rapamycin by Dual Targets Modified Nanoparticles for Alzheimer's Disease Therapy. *Small.* 2022;18:e2203182. ⁷⁴ © 2022 Wiley-VCH GmbH.

inhibitors, showing metabolic stability and sufficient lipophilicity to cross the both barriers of BBB and endosomal membranes, are usually found via high-throughput screening (HTS) and followed by lead optimization. For example, on the basis of isothioureia, a weak BACE1 ligand, the amidine moiety and diaryl amide substituent were modified to provide the high affinity for the continuous hydrophobic subsite of BACE1, as well as a high permeability for oral absorption, and MK-8931 (verubecestat) guanidine containing scaffold was designed (Figure 3B). In addition to its satisfying drug properties, it could significantly reduce A β ₁₋₄₀ content in cerebrospinal fluid and cortex by oral administration at the dose of 10 or 30 mg/kg (Figure 3C and D), thereby rescuing the memory loss of AD. Favorable data facilitated verubecestat to reach Phase III, but it was all overthrown by the unchanged cognitive ability along with adverse events.⁷³ Other several of these BACE1 inhibitors have entered into clinical trials. The atabecestat and LY2886721 successfully alleviated the cognitive decline in the phase II/III trial but was discontinued owing to hepatic-related adverse events.⁷⁷ A phase II trial of lanabecestat aiming at prodromal AD and mild AD participants was suspended because of futility.⁷⁸ Clinical trials of BACE1 inhibitors can be described as a graveyard of new drug discovery for AD. Pharmaceutical companies seem to have given up on BACE1 inhibitors altogether as none are currently listed in any company's experimental or clinical development pipeline.

In general, the safety of BACE1 inhibition therapy is a topic of concern. The over-inhibition of BACE1 may result in shortened lifespan, epilepsy and nerve desheathing according to transgene mice which are born to lack BACE1.⁷⁹ However, recent research shows that this kind of BACE1-knockout therapy in mature mice or brain-target-knockout did

not exhibit above adverse reaction, though the impairment of the formation of myelin was observed in the hippocampus, which may result in the Axon disorders in hippocampal mossy fiber pathway. Although the levels of BACE1 inhibition and A β reduction required for disease modification are unknown, they could be deduced from data collected in the clinical trials in progress. Data from clinical trials and preclinical works indicated that roughly 50% of BACE1 inhibition might be an appropriate therapeutic level.^{79–81} In sum, all the failure of latest drug candidates targeting BACE1 is mainly on account of its persistent side effects and the inefficacy in humans which might be in connection with genetic background discrepancy between species.⁸² Attention may be supposed to focus on appropriate dosage determination, avoiding negative results and pertinent animal models in further development.

Compared with small-molecule inhibitors, the small interfering RNA (siRNA) developed these years can directly block the disease-causing gene expression with appropriate dose and level to a certain extent to avoid the side effects and shows the advantages of good targeting ability with low effective doses. However, due to its poor stability and low permeability efficiency of BBB, nanomaterials for delivering siBACE1 are also being developed to propose solutions. A glycosylated modified nanocarrier was designed to utilize the glucose transporter-1 (GLUT1) to promote the BBB penetration efficiency of siRNA of BACE1.⁸³ The stability was further enhanced by fluorinating the modified carrier. The obtained Gal-NP@siRNA nanodrugs had a 4.9-fold longer half-life in the blood than free siRNA, exhibiting excellent stability. Meanwhile, the brain accumulation of Gal-NP@Cy5-siRNA nanoparticles was 5.8-fold higher than the group without glycosylation modifications (NP@Cy5-siRNA), proving that improving BBB permeability through GLUT1 is a worthy method. In addition to vehicle modification, choosing the appropriate delivery method is also one of the ways to improve the efficiency of brain targeting and accumulation. Nasal administration is a method enabling bypass BBB and owns strong compliance, suitable for multiple administration of AD with a long course of disease. Gao et al designed an Aleuria aurantia lectin (AAL)-modified nanoparticle that could bind to fucose residue and locate at the nasal olfactory epithelium, and further enhance the ability to target A β lesion regions of the brain by modifying KLVFF (Figure 3E).⁷⁴ The nanocarrier contains rapamycin capable of inducing microglial autophagy activation and siRNAs that silence the BACE1 gene. After nasal administration, AAL-modified nanoparticles accumulated in large quantities in the brain within 2 h (Figure 3F and G), indicating that AAL modification can improve nose-to-brain transport and promote brain-targeted delivery.

Strategies to Inhibit A β in Advance

Immunotherapy against A β includes active immunotherapy represented by vaccines, and passive immunotherapy which has been detailed in the “Immunotherapies with anti-A β monoclonal antibodies in AD patients” section. In contrast to passive immunotherapy, vaccination facilitates the endogenous production of therapeutically effective antibodies without monthly re-administration, which makes vaccination more affordable and compliant. These superiorities address many of these limitations and highlight the need for vaccines against pathologically specific targets in populations bearing the risk of AD.

In 1999, human full-length A β peptides mixing with adjuvants were first injected into transgenic mice and were found to produce antibodies, completely preventing plaque formation.⁸⁴ As for the old mice, the A β deposition was observed significant reduction. As a result, A β active immunotherapy officially entered the stage of an AD treatment strategy. AN-1792 is the first vaccine reaching clinical trial, it showed A β plaque clearance in phase I trials.⁸⁵ However, the meningitis that happened in phase II hindered the further trials, but it did not stop the researchers from exploring further. The CAD106 developed by Novartis consists of truncated A β fragments (A β _{1–6}) to avoid T-cell activation.⁸⁶ No signs of meningitis within the 52-week treatment of the Phase I trial demonstrated the success of this strategy in improving vaccine safety. The UB-311 vaccine designed by United Biomedical was optimized by linking the A β _{1–14} epitope to the helper T cell peptide epitope.⁸⁷ This strategy was able to induce a B cell response to enhance immunogenicity while avoiding the inflammatory response of T cells. Although no A β -targeting vaccine has yet been approved for marketing, several candidates are undergoing clinical trials.

In addition to the most widely studied A β target, the N-terminally truncated A β peptide species harboring a pyroglutamate at position 3 (pE3-A β) was also focused by latest studies as a new promising target. It is derived from the subspecies pE3-A β which is a major component of intracerebral toxic plaques. Similar to A β _{1–42}, pE3-A β begins to accumulate much prior to AD symptoms occurring, demonstrating its potential as a target for vaccine development.⁸⁸ Recently, Ghochikyan et al decorated a C-terminal azide modifying pyroglutamated aa 3–11 peptide to the MultiTEP

carrier protein utilizing copper-free click chemistry. This vaccine successfully reduced both pE3-A β and full-length A β in 5 $\times$ FAD mice.⁸⁹ FDA approval of the aducanumab has greatly promoted the research and development of other AD immunotherapies. Efforts still needed to be made to overcome the obstacles, including side effect events, target selection, and delivery platform, with the aim of developing more effective, even customized immunotherapies.

The Promotion of A β Clearance

Pathological results have shown that the impaired efficiency of A β clearance rate is the main inducement of a late-onset form of AD (LOAD) which accounts for a great proportion of total AD cases, indicating that improvement of A β elimination rate is another promising method to alleviate AD pathology.¹⁰ Recent evidence has revealed that intracerebral A β clearance is not only limited to the brain but consists of both central and peripheral pathways.⁹⁰ The central way mainly includes glial cell-mediated catabolism and A β -degrading enzymes. Glial cell incorporates microglia and astrocyte. The former is a sort of innate immune cell in charge of cerebral microenvironment monitoring.⁹¹ Microglia are also a secreted source of some A β -degrading enzymes, such as insulin-degrading enzyme (IDE) and matrix metalloprotease-9 (MMP-9).⁹² Astrocyte is responsible for cell-to-cell communication and part of the nutrients that neurons require. The enzyme pathway requires many kinds of A β -degrading enzymes (ADEs), and a large number of ADEs have been discovered, including neprilysin (NEP), IDE, MMP-9, and so on. These enzymes can degrade full-length A β into smaller peptides with less toxicity, which are then cleared by glial cells or efflux out of the brain.⁹³ As for the peripheral approach, the vasculature undertakes a critical role in ferrying cerebral-derived A β to peripheral organs (primarily the liver and kidney) to realize final degradation.⁹⁴

As the disease progresses with age and progression of AD, both pathways of clearance are affected to varying degrees, resulting in a decrease in clearance. Therefore, accelerating the clearance of A β is also a major strategy for AD treatment.

Nanomedicine Strategies of A β Central Clearance

During brain development, microglia play the role of clearing excess synapses to maintain neuronal homeostasis, while toxic A β plaque would induce the state of microglia from quiescent condition to reactive condition.⁹⁵ Proinflammatory cytokine released by hyperactivated microglia promotes astrocyte activation and contributes to A β -damaged neurons, also impairing synaptic function, and hindering cell communication and growth. Microglial regulation is, therefore, one of the attractive approaches for central A β clearance.

Previous studies have shown that a critical signal transformer, nuclear factor-kappa B (NF- κ B) proteins, has a key role in microglia inflammation regulation.⁹⁶ Once the NF- κ B pathway is activated in microglia, it stimulates the secretion of ROS and pro-inflammatory cytokines, which in turn exert secondary neurotoxic effects. The team of Ding et al exploited an apoA-I-based HDL as an A β nano scavenger to achieve central clearance by microglia phagocytosis in the brain.⁹⁷ Meanwhile, the natural compound curcumin is used to inhibit the NF- κ B pathway of microglia and reduce the production of pro-inflammatory cytokines. These two complement each other to enable the continuous functioning of microglia (Figure 4A). After immunofluorescence labeling of microglia (Iba1 and CD86) and A β deposition, it can be seen in Figure 4B that pHDL/Cur-siBACE1 preparation could enhance microglial infiltration near plaques, in contrast to the lack of microglia next to plaques in AD mice. It was indicated that after treatment, the dysfunction of microglia phagocytosis A β was effectively reversed, and curcumin successfully regulated microglial function and improved A β clearance with silencing BACE1 (Figure 4C). In addition to inflammation, autophagy, and mitochondrial function regulation are also important strategies for microglia to target AD therapy. Among them, the Akt/mTOR/HIF-1 α signaling pathway has been shown to be an important pathway to regulate microglial phenotypic transition and autophagy. Figure 4D Glutathione (GSH)-functionalized gold nanocage was designed and loaded with immunosuppressant fingolimod hydrochloride (FTY720) to form mTOR-mediated immunometabolic reprogramming nanoparticles (GAF NPs).⁹⁸ Through CD86/CD206 double staining (Figure 4E), it was evident that GAF NPs were able to reduce the green fluorescence of the hippocampus and cortex representing CD86 positivity, indicating that the phenotype of microglia in the brain changed from M1 to M2, which is more conducive to A β clearance.

Emerging evidence has demonstrated that triggering receptors expressed on myeloid cell 2 (TREM2) deficient microglia tend to gather around A β deposition, and TREM2.⁹⁹ The interrelationship between AD pathogenesis and

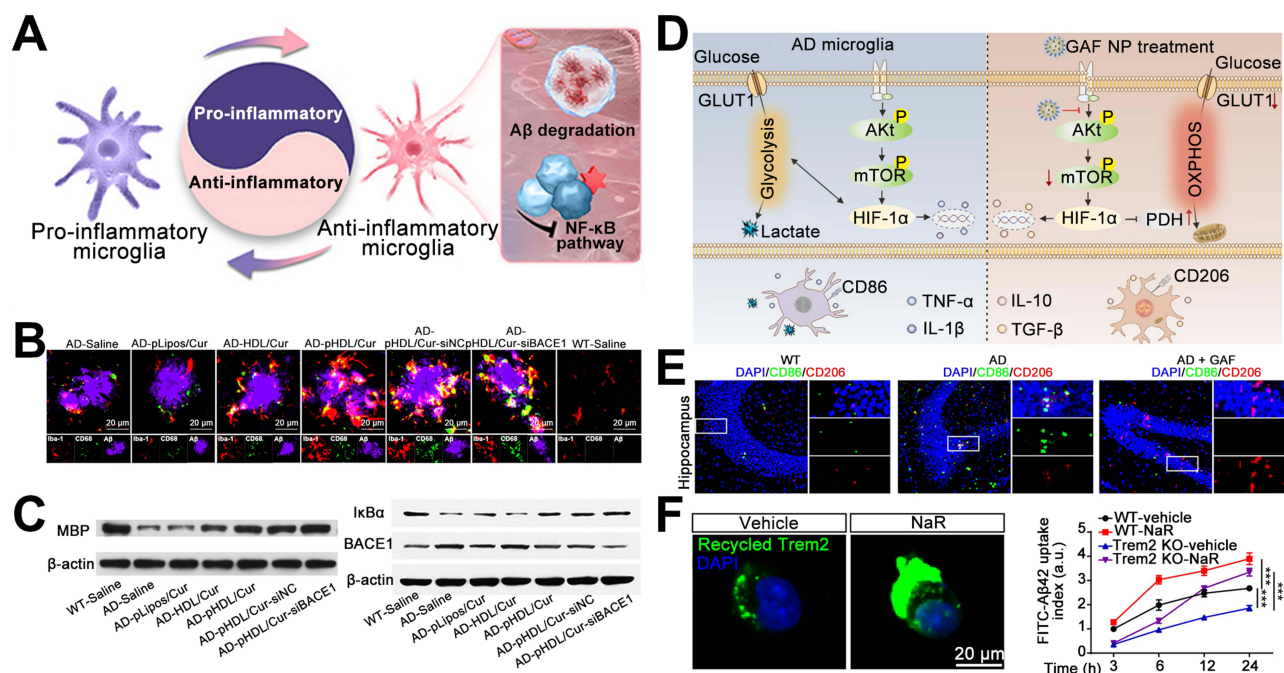


Figure 4 Strategies of A β central clearance via microglia. **(A)** The normalization of microglial dysfunction by blocking the NF- κ B pathway to accelerate A β degradation. **(B and C)** pHDL/Cur-siBACE1 reduced microglia activation by inhibiting the NF- κ B signal and decreased the A β deposition. Reproduced with permission from Zhang H, Jiang W, Zhao Y, et al. Lipoprotein-Inspired Nanoscavenger for the Three-Pronged Modulation of Microglia-Derived Neuroinflammation in Alzheimer's Disease Therapy. *Nano Lett.* 2022;22:2450–2460. Copyright © 2022, American Chemical Society.⁹⁷ **(D)** The illustration of how GAF NPs act on the Akt/mTOR/HIF-1 α signaling pathway. **(E)** GAF NP treatment mobilized microglial polarization from the M1 to M2 in AD mouse brain. Reproduced with permission from Yang F, Zhao D, Cheng M et al mTOR-Mediated Immunometabolic Reprogramming Nanomodulators Enable Sensitive Switching of Energy Deprivation-Induced Microglial Polarization for Alzheimer's Disease Management. *ACS Nano.* 2023. Copyright © 2023 American Chemical Society.⁹⁸ **(F)** NaR treatment significantly increased the recycling of TREM2 in microglial cells, and NaR-mediated A β phagocytosis and clearance are not affected by microglial phagocytosis receptors. *** $p < 0.001$. Reproduced from Pan R-Y, Ma J, Kong X-X, et al. Sodium rutin ameliorates Alzheimer's disease-like pathology by enhancing microglial amyloid- β clearance. *Sci Adv.* 2019;5:eaau6328. Copyright © 2019 The Authors, some rights reserved; exclusive licensee American Association for the Advancement of Science. No claim to original US Government Works. Distributed under a Creative Commons Attribution NonCommercial License 4.0 (CC BY-NC).¹⁰⁰

several TREM2 variants has been suggested to be vital for microglial activation. It was previously reported that the rutin derivative NaR was able to enhance the infiltration of microglia around A β plaques.¹⁰⁰ This is due to the enhancement of the number of phagocytosis receptors on the cell surface by NaR, specifically the expression and recycling of TREM2 (Figure 4F). To further investigate the potential regulatory pathway based on TREM2, Sheng et al proposed a hypothesis that TREM2 may have extracellular ligands and identified a group of lipoprotein particles with apoE as the main apolipoproteins.¹⁰¹ Results showed that, through lipid apoE interaction with TREM2, the extracellular A β clearance via microglial accelerated (Figure 4F). Due to the ambiguous relationships between inflammation and TREM2, it is still not clear how microglial lacking or overexpressing TREM2 is involved in phagocytosis in AD. Consequently, proper TREM2 function on AD needs to be clarified as a priority in future studies.

Astrocytes, a major member of intracerebral glial cell, contribute to synaptogenesis, maintenance of the neural circuit function, and formation of BBB.¹⁰² Astrocytes possess the ability to detect neurotransmitters and initiate a response by increasing intracellular calcium level. Additionally, they are capable of communicating with neurons through the release of gliotransmitters.¹⁰³ Evidence showed that the reactivity of astrocytes has a close relationship with increased A β production, thereby inducing A β -triggered tau pathology.¹⁰⁴ Hence, regulating the dysfunction of astrocytes has become a potential therapeutic avenue. Considering the participation of astrocytes in hippocampal neuronal abnormal activity, the earliest mark observed in both humans and mouse models, A β -dependent transient receptor potential ankyrin 1 (TRPA1) channels capable of activating astrocytes have been studied.¹⁰⁵ After specifically blocking the pharmacology of TRPA1, the activities of both astrocytes and neurons were successfully normalized, thus preventing memory decline. In addition, AD therapeutic strategy was also developed based on astrocyte–microglia cross-talk.¹⁰⁶ The microglia modulated by astrocytic interleukin-3 (IL-3) exhibited more effective A β clearance, limiting the progression of AD.

Nanomedicine Strategies of A β Peripheral Clearance

Recent studies have demonstrated the rapid flow of A β from the brain to the periphery, and the physiological breakdown of brain-derived A β in the peripheral system has been revealed in both humans and mice, providing a novel perspective for understanding the pathogenesis of AD and developing therapeutic approaches. The equilibrium between central and peripheral A β is maintained through continuous exchange,^{94,107} primarily involving drainage across the blood-brain barrier and cerebrospinal fluid (CSF)-interstitial fluid (ISF) convection via the lymphatic system.¹⁰⁸ Specifically, cerebral A β is eliminated by low-density lipoprotein receptor-related protein-1 (LRP1), which is located in brain endothelium and mural cells, serving as a transport mechanism for A β across the blood-brain barrier. Additionally, soluble LRP1 (sLRP1) acts as a ligand in peripheral circulation to prevent free plasma A β from re-entering the brain.¹⁰⁹ Building upon this pathway, Qu et al have developed a biomimetic nanozyme named Cu_xO@EM-K wrapped with erythrocyte membrane from 3 $\times$ FAD mice and A β -specific sequence KLVFF, working synergistically with erythrocyte membrane to selectively recognize A β in blood.¹¹⁰ The Cu_xO core with superior multiple antioxidant enzyme-like activities plays a key role in stabilizing the outer erythrocyte membrane shell maintaining membrane integrity by relieving A β -induced membrane oxidative injury to enable prolonged blood circulation. The well-designed Cu_xO@EM-K successfully captures circulating A β followed by its elimination through liver metabolism. This rapid clearance of peripheral A β facilitates significant efflux of the central A β towards peripheral tissues due to the “sink in” phenomenon resulting in a reduced burden of A β accumulation within the brain. The genius of this design comes from the fully expanded application of inorganic materials such as copper nanoparticles in the treatment of AD, whereas this range of materials was previously thought to cause difficult metabolism and potential security issues in the treatment of central system diseases (Figure 5A and B). The highly safe peripheral circulation undoubtedly provides a more reliable therapeutic environment for such materials and expands the boundaries of anti-AD nanomedicine. In another study, Hyeon et al designed an extracorporeal A β cleansing system, where multifunctional magnetite/ceria nanoparticle assemblies are used to capture and remove A β from flowing blood in a dialysis-like manner. A β antibody-conjugated magnetite/ceria nanoparticles are introduced into the extracorporeal bloodstream circuit for specific capture of A β peptides and achieve facile magnetic separation by applying an external magnetic field. The treated blood then returns to the body after the magnetic separation of the nanoparticles and their bound A β peptides.¹¹¹ 5 $\times$ FAD transgenic mice were used to demonstrate the blood A β cleansing treatment and showed that 71% of A β peptides in the blood on average were removed by the cleansing treatment resulting in a 77% decrease of A β plaques in cerebral cortices compared with the no-treatment group (Figure 5C–E). Both strategies are passive A β drainage using endogenous A β concentration gradient difference. This kind of treatment itself is affected by the course of AD disease. Specifically, A β in AD lesions is more in the form of aggregates that are difficult to flow, and the expression level of key affecting receptors is changed due to the lesions of the blood-brain barrier. Therefore, the application of this kind of passive effecting strategy in AD disease may be limited. In another recent study, Kim et al developed a chemical dissociator of brain plaques to drive the dissociation of A β aggregates in AD brains which release a substantial portion of soluble, smaller A β species to achieve outflow of dissociated amyloid burden from brain to blood (Figure 5F and G).¹¹² This encouraging attempt provides an effective train of thought and tool for overcoming pathogenic A β effect impairment in AD progress, and further work is expected to expand this class of drugs from single dissociator to active drainage in further study.

In addition to BBB-mediated A β efflux, the exchange between CSF and ISF plays a crucial role in removing soluble A β . Although the details of this pathway are not yet fully understood, this progress largely relies on the transportation of the glia-lymphatic (glymphatic) system and the draining of meningeal lymphatic fluid into deep cervical lymph nodes. The pia membrane penetrates the brain parenchyma with blood vessels (Virchow-Robin space),¹¹³ allowing cerebrospinal fluid to flow from the subarachnoid space into the perivascular space and exchange with interstitial fluid in the brain. The polarized aquaporin protein-4 (AQP4) on astrocyte foot processes drives the flow of solutes in the brain parenchyma toward the venous/perivascular Virchow-Robin space.¹¹⁴ The waste products then further flow back into the cerebrospinal fluid and are either recycled through the meningeal lymphatic canal or discharged into the systemic blood circulation through the arachnoid granules. This newly recognized pathway has been shown to be closely related to the metabolism of various pathogenic toxic proteins such as A β and tau. On the basis of these findings, there have been preliminary studies on the application of recombinant vascular endothelial growth factor-C (VEGF-C) and adeno-associated virus expression of VEGF-C in the treatment of AD by directly regulating lymphogenesis and lymphatic vessel function to promote lymphatic

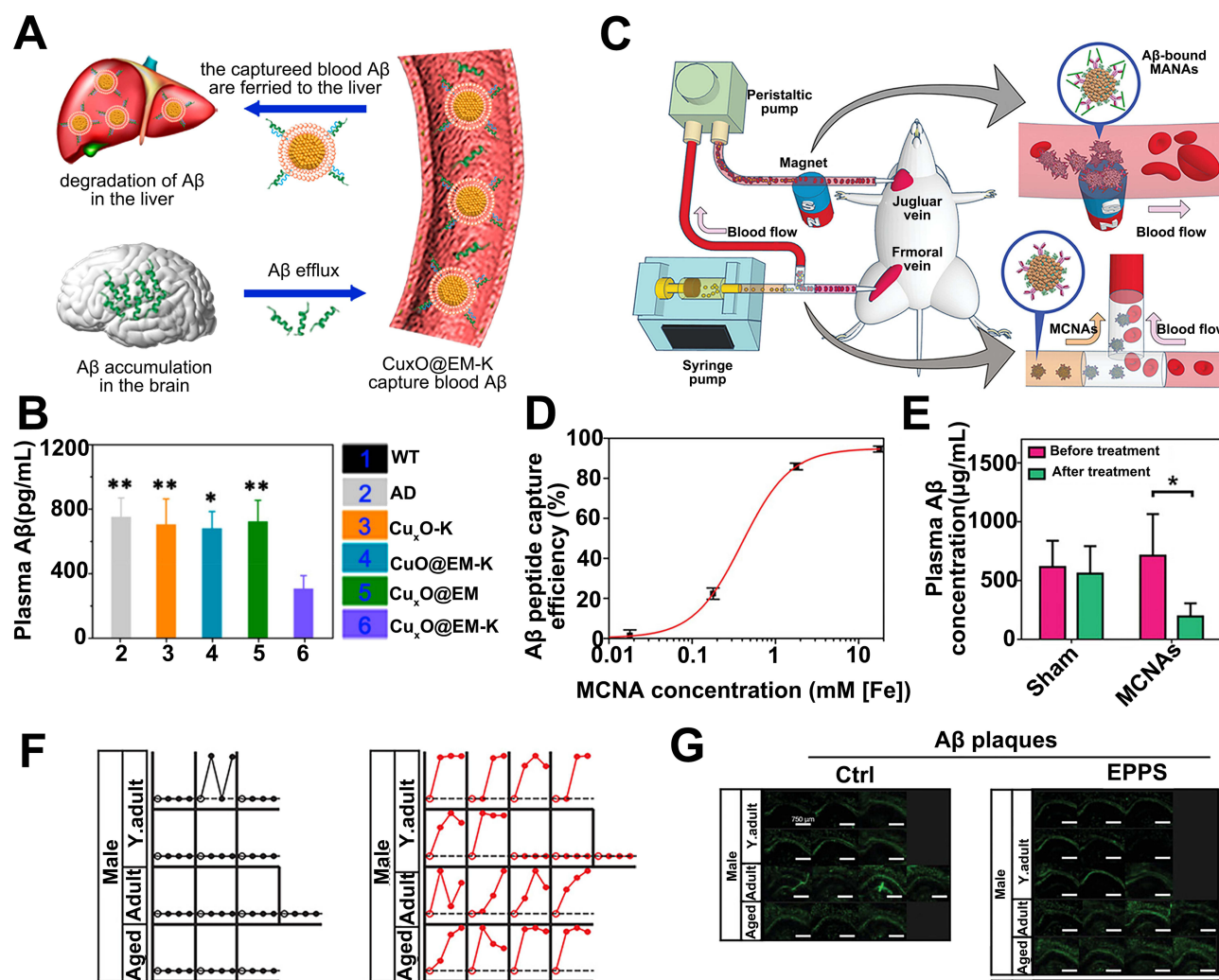


Figure 5 Strategies of Aβ peripheral clearance. (A) Schematic illustration of Aβ captures in the blood by Cu_xO@EM-K, followed by elimination of Aβ bound to Cu_xO@EM-K by the liver. Subsequently, clearance of peripheral Aβ facilitates a large efflux of Aβ from the brain into the blood through the sink effect, leading to the reduction of brain Aβ burden. (B) Blood Aβ levels were measured after treatment with Cu_xO@EM-K. *p < 0.05 and **p < 0.01. Reproduced with permission from Ma M, Liu Z, Gao N, et al. Self-Protecting Biomimetic Nanozyme for Selective and Synergistic Clearance of Peripheral Amyloid-β in an Alzheimer's Disease Model. *J Am Chem Soc.* 2020;142:21702–21711.¹¹⁰ Copyright 2020 American Chemical Society. (C) Schematic illustration of Aβ capture and magnetic separation by magnetic nanoparticle. (D) MCNA concentration-dependent Aβ peptide capture efficiency. (E) Plasma Aβ concentrations were measured before and after the Aβ cleansing treatment with MCNAs. *p < 0.05. Reproduced with permission from Kim D, Kwon HJ & Hyeon T. Magnetite/Ceria Nanoparticle Assemblies for Extracorporeal Cleansing of Amyloid-β in Alzheimer's Disease. *Adv Mater.* 2019;31:e1807965.¹¹¹ © 2019 WILEY-VCH Verlag GmbH & Co. KGaA, Weinheim (F) Individually normalized plasma Aβ tracking data of blood samples collected at weeks 5, 8, and 12 for each mouse in all groups using basal plasma Aβ and maximal plasma Aβ levels of individuals (Control, black; EPPS, red; Y. adult, Young-adult). (G) ThS stained Aβ plaques in the brain tissue of each mouse. Reproduced with permission from Lee D, Kim HV, Kim HY, et al. Chemical-Driven Outflow of Dissociated Amyloid Burden from Brain to Blood. *Adv Sci (Weinh).* 2022;9:e2104542.¹¹² Copyright 2022 John Wiley & Sons, Inc.

drainage and the excretion of toxic metabolites.¹¹⁵ In conclusion, the lymphatic-targeting system may be a viable therapeutic strategy for AD treatment; however, research on the lymphatic system in central nervous system diseases faces challenges, such as difficulty in capturing the anatomical structure and the lack of appropriate means to objectively evaluate the internal drainage function. Therefore, the use of novel imaging technology and nanomedicine platforms to investigate fluid homeostasis and substance drainage in the brain can provide a more intuitive understanding of the anatomy and function of the central lymphatic system, which would make a significant contribution to AD treatment.

Conclusion

Alzheimer's disease (AD) has become one of the most significant health problems worldwide. With the recent emergence of two Aβ monoclonal antibodies approved by the FDA, the treatment of AD has made progress in overcoming the lack

of available disease-modified therapies. Unquestionably, these encouraging developments have confirmed the correlation between A β clearance and cognitive improvement in clinical settings. However, adverse clinical responses also indicate that relying solely on A β clearance is flawed. On one hand, the rapid disintegration of A β plaques leads to difficulties in transporting and clearing a large amount of soluble A β . Therefore, any therapeutic strategy targeting A β must consider stimulating the internal metabolic system to clear A β plaques quickly and safely while also avoiding adverse reactions in the central nervous system.¹¹⁶ On the other hand, these antibodies can only improve mild cognitive impairment. This is because advanced stages of the disease have already caused significant neurological damage and other symptoms, making it necessary for many AD patients to undergo multidimensional drug combinations.^{117,118}

In the research context, the highly controllable and intelligent advantages of nanomedicine provide ample means to address the complex symptoms of AD. However, it must be noted that the prerequisite for the clinical transformation of nanomedicine, that is, the appropriate selection of its research objects, is undoubtedly crucial. At present, most AD animal models can only simulate part of AD lesions, and animal models that fully conform to human AD lesions have not yet come out.¹¹⁹ However, with the development of molecular neurobiology, more effective targets and crosstalk relationships between targets have been discovered,¹²⁰ which makes it difficult to propose a gold standard for AD disease models. Therefore, it is important to select the appropriate animal model according to the pathological process to be studied. In the recent study of the relationship between meningeal lymph and A β metabolism, for example, this lesion was difficult to detect in aggressive 5xFAD mice, although this animal model has been widely used in other A β pathological studies.⁶³ This is precisely because of the inconsistency between the period of the onset of A β pathology and the pathological changes of physiological structure. In view of these considerations, we reviewed the animal models of AD characterized by A β deposition, and attempted to guide the selection of subsequent preclinical animal studies of nanomedical drugs through A β deposition level, deposition initiation time and cascade lesion characteristics. In subsequent drug development, more rational program design will inevitably be accompanied by the selection of reliable animal models, and even the pharmacodynamic research will be extended to the emerging human-derived induced pluripotent stem cell models, so as to more accurately reflect the development of disease in the human body.

Based on this premise, we summarized the design strategy of nanomedicine based on A β in the whole life cycle of A β , and interpreted the intervention strategy for the whole life cycle of A β from the perspective of inhibiting A β production (including inhibition of secretase and active immunoprophylaxis of A β production) and promoting A β clearance. Different from previous reviews that focused more on functional nanocarrier design, we describe the potential adverse reactions of excessive A β clearance and microglial inflammation, and propose a strategy of synergistic A β clearance and burden reduction through microglial polarization regulation. With the clarification of the metabolic pathways of A β inside and outside the brain, especially the breakthrough of central metabolic pathways such as meningeal lymphatic vessels, the metabolism of A β has become a systematic process, emphasizing the integrity of pathological intervention in AD.¹¹⁵ Therefore, the next generation of A β nanomedical drug development may pay more attention to the regulation of its transport process in multiple sites throughout the body by taking advantage of systematic elimination with greater potential.¹²¹ In summary, this paper aims to guide the development of nanomedicine from the perspective of AD pathology and promote the development of A β -based synergistic therapies.

Abbreviations

AD, Alzheimer's disease; SP, senile plaques; A β , amyloid-beta; NFTs, neurofibrillary tangles; FDA, Food and Drug Administration; AChE, acetylcholine esterase; NMDA, N-methyl-D-aspartic acid; APP, amyloid precursor protein; ARIA, amyloid-related imaging abnormalities; BBB, blood-brain barrier; mAbs, A β monoclonal antibodies; CDR-SB, Clinical Dementia Rating Scale-Sum of Boxes; MCI, mild cognitive impairment; CTAD, Clinical Trials on Alzheimer's Disease; iADRS, Integrated Alzheimer's Disease Rating Scale; PS1, Presenilin 1; BACE1, beta-site amyloid precursor protein cleaving enzyme 1; HE, hydroxyethylene; HTS, high-throughput screening; siRNA, small interfering RNA; GLUT1, glucose transporter-1; AAL, aleuria aurantia lectin; pE3-A β , A β peptide species harboring a pyroglutamate at position 3; LOAD, late-onset form of AD; IDE, insulin-degrading enzyme; MMP-9, matrix metalloprotease-9; ADEs, A β -degrading enzymes; NEP, neprilysin; NF-KB, nuclear factor-kappa B; TREM2, triggering receptor expressed on myeloid cells 2; CSF, cerebrospinal fluid; ISF, interstitial fluid; LRP1, low-density lipoprotein receptor-related protein-1; AQP4, polarized aquaporin protein-4; VEGF-C, vascular endothelial growth factor-C.

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Disclosure

The authors declare that they have no conflicts of interest.

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