Anti-Amyloid-β Monoclonal Antibodies for Alzheimer’s Disease: Pitfalls and Promise

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ABSTRACT
The majority of putative disease-modifying treatments in development for Alzheimer’s disease are directed against the amyloid-β (Aβ) peptide. Among the anti-Aβ therapeutic approaches, the most extensively developed is immunotherapy—specifically, passive immunization through administration of exogenous monoclonal antibodies (mAbs). Although testing of mAbs has been fraught with failure and confusing results, the experience gained from these trials has provided important clues for better treatments. This review summarizes the experience to date with anti-Aβ mAbs to enter clinical trials for Alzheimer’s disease and examines the evidence for clinical efficacy and the major problems with safety—i.e., amyloid-related imaging abnormalities. As mAbs differ considerably with regard to their epitopes and the conformations of Aβ that they recognize (monomers, oligomers, protofibrils, fibrils), the consequences of targeting different species are also considered. An often-cited explanation for the failure of anti-Aβ mAb trials is that they are set too late in the disease process. New trials are indeed evaluating treatments at prodromal and preclinical stages. We should expect to see additional studies of presymptomatic Alzheimer’s disease to join the ongoing prevention trials, for which mAbs continue to serve as the mainstay.

Keywords: Alzheimer’s disease, Amyloid-β, Amyloid-β oligomers, Amyloid-related imaging abnormalities, Immunotherapy, Monoclonal antibodies

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The amyloid hypothesis of Alzheimer’s disease (AD) holds that the accumulation of the amyloid-β (Aβ) peptide leads to synaptic dysfunction, neurodegeneration, and ultimately symptoms (1). The vast majority of potential disease-modifying treatments developed in recent years are directed against Aβ, including inhibitors of the synthetic enzymes gamma-secretase and beta-secretase, and Aβ aggregation inhibitors. However, the most elaborated anti-Aβ approach is immunotherapy, including both active vaccines to stimulate the immune system to produce its own antibodies and passive immunization through the administration of exogenous antibodies.

The advantage of active immunotherapy is long-term antibody production from short-term drug administration at limited cost. Conversely, immune response may be inconsistent or lacking, especially in older individuals, and adverse reactions—if immunologically based—may also be long-lasting. Initial experience with active vaccines was marred by an ill-fated trial of AN1792 (full-length Aβ[42] with QS-21 adjuvant) that was halted following the occurrence of T cell–mediated meningoencephalitis in 6% of treated participants (2). Second-generation vaccines such as ACC-001 (3–5) and CAD106 (6,7) have sought to generate anti-Aβ antibodies restricted to the N-terminus, while avoiding T cell epitopes at the C-terminus (8,9). CAD106 is the only vaccine to advance to phase 3 trials and has been selected for the Alzheimer Prevention Initiative APOE ε4 homozygote study (https://clinicaltrials.gov; Identifier: NCT02565511) (10).

In contrast to active vaccination, passive immunization has the advantages of ensuring consistent antibody titers and allowing control of adverse events by stopping treatment. The major drawbacks of monoclonal antibodies (mAbs) are the need for repeated administrations and the associated cost of production (11). Over the past approximately 15 years several mAbs have been engineered to bind and clear Aβ (Table 1) and have advanced to human trials (Table 2). Although the testing of mAbs has been fraught with failure and confusing results, the experience gained from these trials has provided important clues to enable the development of better treatments.

BAPINEUZUMAB
Bapineuzumab (AAB-001; Pfizer Inc., New York, NY, and Janssen Pharmaceuticals, Inc., Raritan, NJ), a humanized immunoglobulin (Ig) G1 anti-Aβ mAb, binds the five N-terminal residues and clears both fibrillar and soluble Aβ. In 2000, Bard et al. (12) reported that in PDAPP transgenic mice, 3D6 (the murine precursor of bapineuzumab) entered the brain, decorated plaques, and induced the Fc receptor–mediated microglial phagocytosis of Aβ deposits.

Bapineuzumab was the first mAb to enter human testing after termination of the AN1792 trial. In a phase 1 single ascending dose trial, 0.5, 1.5, or 5 mg/kg of bapineuzumab was generally safe and well tolerated in 30 participants with mild to moderate AD (13). However, 3 of 10 participants in the
highest dose group developed magnetic resonance imaging (MRI) abnormalities consistent with vasogenic edema, all of which later resolved. Two participants were asymptomatic, and one experienced mild, transient confusion. These events prompted the Alzheimer’s Association Research Roundtable to convene a Workgroup in July 2010, which coined the term amyloid-related imaging abnormalities (ARIA) to refer to MRI signal alterations associated with Aβ-modifying therapies—specifically, ARIA-E to denote vasogenic edema/effusions and ARIA-H to indicate microhemorrhage and hemosiderosis (14). The subsequent phase 2 trial studied intravenous bapineuzumab (0.15, 0.5, 1.0, or 2.0 mg/kg) administered every 13 weeks for 78 weeks in mild to moderate AD (15). No significant treatment differences were found for the primary efficacy end points, Alzheimer’s Disease Assessment Scale—Cognitive Subscale [ADAS-Cog11] (16,17) or Disability Assessment for Dementia (18), but prespecified exploratory analyses showed potential treatment differences for subjects who completed the study and APOE ε4 noncarriers. A parallel phase 2 study with [11C]-Pittsburgh compound B (11C-PiB) and positron emission tomography (PET) in 28 participants revealed some clearance of fibrillar Aβ (19). A retrospective review by two neuroradiologists of MRI scans from the phase 2 studies revealed that 36% of APOE ε4 carriers only, as treatment groups differed in change in brain Aβ burden by 11C-PiB-PET (22) and cerebrospinal fluid (CSF) phosphorylated tau concentrations. Negative baseline 11C-PiB-PET scans were found in 36% of APOE ε4 noncarriers, suggesting the necessity of incorporating biomarker evidence of disease into eligibility criteria in future trials.

**SOLANEZUMAB**

Solanezumab (LY2062430; Eli Lilly and Company, Indianapolis, IN), a humanized IgG1 mAb, binds the mid-domain of Aβ (residues 16–26) and increases clearance of monomers (23). Studies in transgenic PDAPP mice demonstrated that m266 (the murine precursor of solanezumab) reduced brain Aβ burden without binding Aβ deposits (24,25), opening the possibility of targeting the soluble pool of Aβ. Phase 1 and 2 studies of solanezumab revealed evidence of target engagement by dose-dependent increases in plasma and CSF total Aβ(26,27). In the phase 2 study of mild to moderate AD, 12 weeks of solanezumab treatment yielded a dose-dependent increase in CSF-free Aβ42, suggesting a shift in equilibria sufficient to mobilize Aβ42 from plaques (27).

The first phase 3 studies—EXPEDITION 1 and EXPEDITION 2—were 18-month trials of solanezumab 400 mg versus placebo (administered intravenously every 4 weeks) in 1012 and 1040 participants with mild to moderate AD (28). The original co-primary outcomes in both studies were the ADAS-Cog11 and Alzheimer’s Disease Cooperative Study–Activities of Daily Living (ADCS-ADL) (29). After analysis of data from EXPEDITION 1, the primary outcome for EXPEDITION 2 was revised to the ADAS-Cog14 in the mild AD subgroup (28). Solanezumab did not demonstrate significant benefit for the primary outcomes in either study but showed a favorable safety profile, as the incidence of ARIA-E was 0.9% with solanezumab and 0.4% with placebo. A prespecified subgroup analysis of pooled data from EXPEDITION 1 and EXPEDITION 2.

### Table 1. Monoclonal Antibodies Bind Different Epitopes and Conformations of Amyloid-β

| Antibody       | Manufacturer                          | Origin    | Subclass | Epitope    | Conformations Recognized | Monomer | Oligomer | Fibril | ARIA-E |
|----------------|---------------------------------------|-----------|----------|------------|--------------------------|---------|----------|--------|--------|
| Bapineuzumab   | Pfizer Inc./Janssen Pharmaceuticals, Inc. | Humanized | IgG1     | AA 1–5     | Yes                      | Yes     | Yes      | High   |        |
| Solanezumab    | Eli Lilly and Company                 | Humanized | IgG1     | AA 16–26   | Yes                      | No      | No       | Low    |        |
| Gantenerumab   | Hoffman-La Roche                      | Human     | IgG1     | AA 3–12, 18–27 | Weak                | Yes     | Yes      | High (?) |        |
| Crenuzumab     | Genentech, Inc.                       | Humanized | IgG4     | AA 13–24   | Yes                      | Yes     | Yes      | Low    |        |
| Ponezumab      | Pfizer Inc.                           | Humanized | IgG2     | AA 30–40   | Yes                      | No      | No       | None   |        |
| BAN2401        | BioArctic Neuroscience, AB/Eisai Co., Ltd. | Humanized | IgG1     | Protoberin | —                      | —       | —        | —      | —      |
| Aducanumab     | Biogen, Inc.                          | Human     | IgG1     | AA 3–6     | No                      | Yes     | Yes      | High   |        |

Epitope, Conformations Recognized, and ARIA-E are explained further in the text. Dashes indicate absence of information. AA, amino acid; ARIA-E, amyloid-related imaging abnormalities—edema; Ig, immunoglobulin.
| Drug          | Publication                   | Phase | Sample | Participants | Age, Years | Dose                              | Duration, Weeks | Efficacy                        | ARIA-E                                                                 | Biomarkers                                                                 |
|--------------|-------------------------------|-------|--------|--------------|------------|----------------------------------|----------------|--------------------------------|--------------------------------------------------------------------------|---------------------------------------------------------------------------|
| Bapineuzumab | Salloway et al., 2009 (15)    | 2     | 234    | Mild-moderate AD | 50-85     | 0.15, 0.5, 1, 2 mg/kg IV every 3 months | 78             | Failed primary end points       | 17%, retrospective analysis                                              | No effect on CSF $A\beta_{42}$, tau, or p-tau                          |
| Bapineuzumab | Rinne et al., 2010 (19)       | 2     | 28     | Mild-moderate AD | 50-80     | 0.5, 1, 2 mg/kg IV every 3 months  | 78             | Failed primary end points       | Retrospective analysis combined with Salloway, 2009                      | $|$Cortical $^{11}$C-PiB compared with baseline and placebo                |
| Bapineuzumab | Salloway et al., 2014 (21)    | 3     | 2204   | Mild-moderate AD | 50-88     | 0.5, 1, 2 mg/kg IV every 3 months  | 78             | Failed primary end points       | 15.3% of $APOE\varepsilon_4$ carriers, 4.2%, 9.4%, and 14.2% of three dose groups in noncarriers | $|$Cortical $^{11}$C-PiB and $|$CSF p-tau in $APOE\varepsilon_4$       |
| Solanezumab  | Farlow et al., 2012 (27)      | 2     | 52     | Mild-moderate AD | >50       | 100, 400, 1600 mg/ month IV       | 52             | No cases                       | $|$A$\beta_{40}$ and $|$A$\beta_{42}$ in CSF                            |
| Solanezumab  | Doody et al., 2014 (28)       | 3     | 2052   | Mild-moderate AD | >55       | 400 mg IV every month             | 78             | Failed primary end points       | 0.9% solanezumab vs. 0.4% placebo                                       | No effect on brain $A\beta$ (PET); $|$A$\beta_{40}$ and $|$A$\beta_{42}$ in CSF |
| Solanezumab  | Completed                    | 3     | 2129   | Mild AD, $A\beta^+$ | 55-90     | 400 mg IV every month             | 78             | Failed primary end point        | No effect on brain $A\beta$ or tau (PET)                               | $|$Cortical $^{11}$C-PiB compared with baseline                         |
| Gantenerumab | Ostrowitzki et al., 2012 (34) | 1     | 18     | Mild-moderate AD | 50-90     | 60, 200 mg IV every 4 weeks       | 24             | 2/6 participants on 200-mg dose | $|$Cortical $^{11}$C-PiB compared with baseline                         |
| Gantenerumab | Ongoing                      | 2/3   | 799    | Prodromal AD, $A\beta^+$ | 50-85     | 105 or 225 mg SC every 4 weeks    | 104            | Nonsignificant benefit          |                                        |                                                                 |
| Crenezumab   | Cummings et al., in press (38)| 2     | 431    | Mild-moderate AD | 50-80     | 300 mg SC every 2 weeks, 15 mg/kg IV every 4 weeks | 68             | Failed primary end points       | 1 case, $APOE\varepsilon_4$ homozygote                                 | $|$CSF $A\beta_{42}$                                                     |
| Crenezumab   | Completed                    | 2     | 91     | Mild-moderate AD | 50-80     | 300 mg SC every 2 weeks, 15 mg/kg IV every 4 weeks | 68             | Failed primary end points       | No effect on brain $A\beta$ (PET); $|$A$\beta$ in CSF                   |
| Crenezumab   | Ongoing                      | 3     | Mild-prodromal AD, $A\beta^+$ | 50-85     | 100                                               |                     |                                             |                                                                 |
| BAN2401      | Ongoing                      | 2     | Mild-prodromal AD, $A\beta^+$ | 50-90     | 2.5, 5, 10 mg/kg IV every 2 weeks, 5, 10 mg/kg IV every 4 weeks | 78             | Failed primary end points       | No cases                                                               | $|$CSF $A\beta_{42}$                                                     |
| Ponezumab    | Landen et al., 2013 (44)     | 1     | Mild-moderate AD | >50       | 10 mg/kg IV                             | 52             | Failed primary end points       | No cases                                                               | $|$CSF $A\beta_{42}$                                                     |
| Aducanumab   | Sevigny et al., 2016 (50)    | 1     | 165    | Mild-prodromal AD, $A\beta^+$ | 50-90     | 1, 3, 6, 10 mg/kg IV every 4 weeks | 54             | Exploratory; $|$decline in CDR (10 mg/kg) and MMSE (3, 10 mg/kg)       | 3%, 6%, 37%, 41% of four dose groups                                    | $|$Cortical $^{[18F]}$-florbetapir                                      |
| Aducanumab   | Ongoing                      | 3     | Mild-prodromal AD, $A\beta^+$ | 50-85     | 78                                           |                     |                                             |                                                                 |

AD, Alzheimer’s disease; $A\beta^+$, positive for amyloid-$\beta$ biomarker (PET or CSF); $APOE\varepsilon_4^+$, positive for $APOE\varepsilon_4$; ARIA-E, amyloid-related imaging abnormalities–edema; CDR, Clinical Dementia Rating; CSF, cerebrospinal fluid; IV, intravenous; MMSE, Mini-Mental State Examination; PET, positron emission tomography; p-tau, phosphorylated tau; $^{11}$C-PiB, $^{[11}$C]-Pittsburgh compound B; SC, subcutaneous.
showed that in participants with mild AD, there was a 34% slowing of decline for the ADAS-Cog14 and 18% for the instrumental items of the ADCS-ADL (ADCS-iADL) (23). Therefore, a third phase 3 trial, EXPEDITION 3 (NCT01900665), restricted to mild-stage AD, was launched in July 2013. Owing to the high rate of cases negative for the Aβ biomarker in EXPEDITION 1 and EXPEDITION 2, the EXPEDITION 3 trial required PET showing positive Aβ for eligibility.

In December 2016 at the Clinical Trials on Alzheimer’s Disease meeting, the negative results of EXPEDITION 3 were presented (30). In 2129 participants with mild AD (confirmed by positive Aβ on PET), solanezumab provided a nonsignificant 11% slowing of decline on the primary outcome, the ADAS-Cog14. This effect size was smaller than in the pooled subgroup analysis from EXPEDITION 1 and EXPEDITION 2. Several secondary outcomes favored solanezumab, including the Clinical Dementia Rating Sum of Boxes (CDR-SB), Mini-Mental State Examination (MMSE), and ADCS-iADL; however, these analyses were not corrected for multiplicity. Solanezumab had no effect on Aβ and tau PET biomarkers. Based on the results of EXPEDITION 3, the development of solanezumab for dementia was discontinued.

The excellent safety profile of solanezumab and encouraging trends in the exploratory analyses in mild AD led to its inclusion in two secondary prevention trials, which are continuing in the hope that earlier intervention may yield more substantial benefit. The A4 study (NCT02008357) started in February 2014 and will enroll 1150 cognitively normal individuals 65–85 years of age who have positive Aβ on PET scans (31). The Dominantly Inherited Alzheimer Network (DIAN) selected solanezumab (and gantenerumab, described below) for its phase 2/3 trial in individuals at risk for and with early-stage autosomal-dominant AD (NCT01760005) (32).

GANTENERUMAB
Gantenerumab (Hoffman-La Roche, Basel, Switzerland), the first fully human IgG1 anti-Aβ mAb, binds a conformational epitope expressed on Aβ fibrils (33). This epitope encompasses both N-terminal (3–12) and central (18–27) amino acids of Aβ and thus requires that the peptide be folded with the midregion near the N-terminus. In PS2APP transgenic mice, gantenerumab significantly reduced Aβ plaques by recruiting microglia and prevented new plaque formation without altering plasma Aβ levels (33).

In phase 1 trials in mild to moderate AD, gantenerumab treatment, including up to seven intravenous infusions (60 or 200 mg) every 4 weeks, reduced brain Aβ burden as measured by 11C-PiB PET (34). Gantenerumab was generally safe and well tolerated, but two of six participants in the 200-mg group experienced ARIA-E (34).

In 2010, a phase 2 trial of gantenerumab was launched in 360 participants with prodromal AD and CSF evidence of Aβ deposition using doses of 105 mg or 225 mg administered subcutaneously every 4 weeks for 2 years. In 2012, the trial was expanded to a phase 2/3 registration trial of 799 participants (NCT01224106). Co-primary end points included CDR-SB and change in brain Aβ levels on [18F]-flortetapir PET. However, the trial was terminated in December 2014 following an interim futility analysis. At the Alzheimer’s Association International Conference meeting in July 2015, the study results were presented and revealed no significant treatment effects for CDR-SB or change in brain Aβ levels (35). However, post hoc subgroup analyses suggested that participants with fast progression—i.e., participants whose hippocampal volume and CDR-SB score declined most rapidly—may have benefited, especially individuals with higher serum levels of gantenerumab. The incidence of ARIA-E and ARIA-H ranged from 0.4% to 14%, increasing with gantenerumab dose and APOE ε4 status. These results were interpreted as supporting the continuation of gantenerumab trials using higher doses (35). Gantenerumab (along with solanezumab) is also being evaluated by DIAN in a phase 2/3 trial in individuals at risk for and with early-stage autosomal-dominant AD (NCT01760005) (32).

CRENEZUMAB
Crenezumab (MABT5102A; Genentech, Inc., South San Francisco, CA) was engineered on an IgG4 backbone to minimize the activation of Fc gamma receptors (36). In transgenic mice, it reduced the Fc gamma receptor–mediated activation of microglia and triggered less release of the proinflammatory cytokine tumor necrosis factor alpha, thought to contribute to neurotoxicity, as well as ARIA (36). Crenezumab prefers the mid-domain of the Aβ peptide (residues 13–24) (37) and binds multiple conformations of Aβ (monomers, oligomers, fibrils), with a 10-fold higher affinity for oligomers versus monomers (36,38). The epitope recognized by crenezumab overlaps that of solanezumab, explaining their observed cross-reactivity but not their different binding profiles for various species of Aβ (39). However, Uhtsch et al. (37) reported that crenezumab and solanezumab actually target slightly different epitopes (residues 13–24 vs. 16–26, respectively). The authors suggested that solanezumab-bound Aβ possesses an alpha-helical structure between residues 21 and 26, whereas crenezumab-bound Aβ has a random coil structure between residues 21 and 24. They further proposed that the alpha-helical epitope is present in monomeric Aβ but absent from aggregated species, potentially explaining solanezumab’s preference for monomers but crenezumab’s recognition of multiple species, including oligomers (37). Phase 1 studies in mild to moderate AD produced no cases of ARIA-E following single doses (0.3–10 mg/kg intravenously) or multiple ascending doses (0.5–5 mg/kg intravenously) of crenezumab (36), allowing higher doses in phase 2.

The major phase 2 trial (NCT01343966) enrolled 431 participants with mild to moderate AD who received either low-dose SC crenezumab 300 mg or placebo biweekly (n = 184) or high-dose intravenous crenezumab 15 mg/kg or placebo every 4 weeks (n = 247) for 68 weeks (38,40). No significant treatment benefits were observed for the primary (ADAS-Cog12 and CDR-SB) or secondary outcomes at either dose. However, in a post hoc subgroup analysis of the high-dose cohort, crenezumab treatment was observed to attenuate decline on the ADAS-Cog12 in the mildest subgroup (MMSE 22–26). A parallel 91-participant biomarker study reported no treatment effects on brain fibrillar Aβ by PET, but CSF Aβ rose slightly with treatment (41,42). Adverse events were balanced between treatment groups, and only one case of ARIA-E was reported in an APOE ε4 homozygote receiving the high dose.
These data were interpreted as supporting the testing of crenezumab at even higher doses in prodromal to mild AD (confirmed by positive Ab on PET). A phase 3 study (NCT02670083) is ongoing in participants with prodromal to mild AD (MMSE 22–30) using a higher dose of crenezumab (38). Participants are randomly assigned to receive intravenous crenezumab or placebo every 4 weeks for 100 weeks, and the primary outcome measure is the CDR-SB. Crenezumab is also being evaluated in a secondary prevention paradigm as part of an Alzheimer Prevention Initiative trial of 300 cognitively normal presenilin 1 carriers from the world’s largest early-onset AD kindred in Antioquia, Colombia (NCT01998841)(10).

PONEZUMAB
Ponezumab (PF-04360365; Pfizer Inc.), a humanized IgG2 mAb, targets the C-terminus of Ab40 (residues 30–40) (43). Compared with IgG1, IgG2 antibodies have a lower propensity to induce immune effector function (44). A number of phase 1 trials tested safety, pharmacokinetics, and pharmacodynamics of ponezumab in mild to moderate AD (44–46). These trials pointed to a favorable safety profile without evidence of ARIA, but the antibody was poorly detectable in CSF. Two subsequent phase 2 studies revealed no clinical efficacy, and development of ponezumab for AD was discontinued (44).

BAN2401
BAN2401 (BioArctic Neuroscience AB, Stockholm, Sweden, and Eisai Co., Ltd., Tokyo, Japan), a humanized IgG1 mAb, selectively binds and clears soluble Ab protofibrils. It was derived from the E22G Arctic mutation in the amyloid precursor protein and has been shown to reduce Ab protofibrils in the brain and CSF of Tg-ArcSwe mice (47). In a phase 1/2a study using single and multiple ascending intravenous doses (48), BAN2401 was well tolerated with no cases of ARIA-E. A phase 2b 18-month trial testing five different intravenous doses was launched in January 2013 in prodromal or mild AD (confirmed by positive Ab on PET) (NCT01767311)(49).

ADUCANUMAB
Aducanumab (BIIB037; Biogen, Inc., Cambridge, MA), a fully human IgG1 mAb, selectively reacts with Ab aggregates, including soluble oligomers and insoluble fibrils (50). It binds the N-terminus (residues 3–6) and recognizes a conformational epitope present on aggregated species of Ab but absent from monomers. Aducanumab was developed by screening libraries of memory B cells from healthy elderly individuals for reactivity against aggregated Ab. In Tg2576 mice, an analog of aducanumab was shown to cross the blood-brain barrier, bind parenchymal Ab, and reduce soluble and insoluble Ab in a dose-dependent manner (50).

A phase 1b clinical trial has been completed in which participants with prodromal or mild AD and Ab-positive PET scans who received 1 year of monthly intravenous infusions of aducanumab (1, 3, 6, or 10 mg/kg) evidenced reduced brain fibrillar Ab in a dose- and time-dependent manner (Figure 1) (50). The phase 1b study was not powered for efficacy; however, exploratory analysis of clinical assessments demonstrated dose-dependent slowing of progression at 1 year. CDR-SB scores declined less with aducanumab treatment, with the greatest slowing at 3 and 10 mg/kg.

The most common adverse effects were ARIA, which occurred at higher levels than in any previous anti-Ab mAb
study. ARIA-E was observed at some point during the trial in no participants in the placebo group compared with 1 (3%), 2 (6%), 11 (37%), and 13 (41%) participants receiving 1, 3, 6, and 10 mg/kg aducanumab, with increased incidence in APOE ε4 carriers. Of the 27 participants who developed ARIA-E, 15 (56%) continued treatment (50).

Based on the promising interim analysis of the phase 1b study, in August 2015, Biogen launched two identical 18-month pivotal phase 3 studies to evaluate the efficacy of monthly doses of aducanumab in slowing cognitive and functional impairment as measured by the CDR-SB. Each trial is enrolling 1350 participants with prodromal or mild AD, as confirmed by Aβ-positive PET (NCT02477800 and NCT02484547).

**Efficacy and Safety of mAbs**

To date, no mAb targeting Aβ has demonstrated significant efficacy. A meta-analysis of immunotherapies by Penninkilampi et al. (51) broadly found no significant treatment differences for typical primary outcome measures, such as the ADAS-Cog, ADCS-ADL, or CDR-SB. However, as noted by the authors, the divergence of mechanisms might be used for considering these agents individually. Solanezumab and crenezumab—both targeting mid-domain Aβ epitopes—have evidenced some post hoc trends for treatment effects in mild-stage AD (23,28,38,40). Solanezumab has completed phase 3 testing in mild-stage AD without meeting its endpoints; however, it continues in preclinical AD trials (31,32). Canumab, which demonstrated substantial reductions in brain amyloid, has been tolerated with aducanumab and gantenerumab, a similar mAb, in an early-phase study, accompanied by slowing of clinical decline at higher doses (50). These results have also provided encouragement for gantenerumab, a similar N-terminal antibody, to continue trials using higher doses (53).

Overall, the safety and tolerability profile of mAbs targeting Aβ has been acceptable. The aforementioned meta-analysis found no difference between pooled treatment and placebo groups in the incidence of adverse events, serious adverse events, and death (51). ARIA, the most concerning safety issue, occurs with N-terminal mAbs that clear fibrillar Aβ—bapineuzumab, gantenerumab, and aducanumab. ARIA-E is strongly associated with drug dose and APOE ε4 status but is also generally (approximately 78%) asymptomatic and self-limiting (20,21) and may not require temporary suspension of treatment (20). Serious complications are rare and must be balanced against the alternative outcome of untreated AD. The same frequency of ARIA events that was dose limiting in early trials of bapineuzumab and gantenerumab has more recently—perhaps fortuitously—been tolerated with aducanumab and associated with possible clinical benefit.

**Targets of Anti-Aβ mAbs**

The lack of efficacy thus far with anti-Aβ mAbs may bolster the case against the amyloid hypothesis of AD (52). However, encouraging results with some antibodies make it equally difficult to dismiss this hypothesis altogether. Converging evidence over the past 2 decades has suggested that the most neurotoxic species of Aβ is the soluble oligomer (1,53), which has emerged as the central target for disease-modifying treatments, including mAbs. Moreover, transgenic mouse models have suggested that therapeutic interventions reducing fibrillar Aβ at the cost of augmenting soluble species could actually be harmful (54), although mAbs that target fibrils may also target oligomers. In this regard, the clearance of fibrillar Aβ on a PET scan is perhaps not an essential goal of treatment but may occur as an epiphenomenon to the clearance of oligomers.

As reviewed by Montoliu-Gaya and Villegas (8), mAbs directed against the N-terminus of Aβ may be most effective in clearing the toxic aggregated species of Aβ. Transgenic mouse models have demonstrated that these antibodies inhibit Aβ aggregation and disaggregate preexisting Aβ fibrils (12,55,56). However, as described by Lu et al. (57), using seeded fibril growth from brain extract and data from solid-state nuclear magnetic resonance and electron microscopy, Aβ40 monomers aggregate in oligomers and fibrils with multiples of three units, in which N-termini are exposed, whereas hydrophobic C-termini are inaccessible to antibodies (8). If a similar structure held true for Aβ42, mAbs targeting the N-terminus would likely be most efficient in clearing Aβ oligomers. The success of N-terminal antibodies in clearing aggregated Aβ may also be related to microglial activation and phagocytosis, which is hypothesized to be a common feature of bapineuzumab, gantenerumab, and aducanumab (12,34,50,58).

Thus far, a tight coupling has been observed between mAbs that target aggregated Aβ and the occurrence of ARIA. If ARIA-E is caused by increased trafficking to and clearance of fibrillar Aβ from cerebral vessels (20), mAbs could be designed with conformationally specific epitopes selective for soluble aggregated species (oligomers and protofibrils) and avoid ARIA-E. Alternatively, if ARIA-E has an inflammatory component (14), antibodies may be designed to avoid inflammation. In this regard, it is unclear whether the infrequency of ARIA-E with crenezumab is related to its IgG4 structure or its mid-domain epitope. Preclinical studies have suggested that it binds all forms of Aβ, including fibrils (36). However, more clinical testing is needed to see if it clears plaques. If, in fact, ARIA-E is more related to inflammation, single-chain variable fragments and other structures lacking the microglia-activating Fc fragment could emerge as promising therapies (59,60). They may offer an alternative, noninflammatory approach to the clearance of Aβ, potentially avoiding ARIA that occurs with complete antibodies.

**Mechanism of Aβ Clearance by mAbs: Brain Entry Versus Peripheral Sink**

Not fully resolved is whether brain entry of anti-Aβ mAbs is necessary, although many experts have attributed the failure of these agents to poor central nervous system penetration (only approximately 0.1% cross the blood-brain barrier) (11). Novel attempts to improve antibody penetration into brain have included targeting receptors on the blood-brain barrier to induce active transport of antibodies into the central nervous system or delivering the genes encoding antibodies and inducing expression in the subject (8).

The peripheral sink hypothesis of mAbs is based on transport of Aβ across the blood-brain barrier as well as an
equilibrium between Aβ in brain and periphery (61,62). By draining plasma Aβ, this equilibrium can be altered to leach Aβ from brain without any direct action of antibodies. Ponezumab exploited the peripheral sink effect—at least for plasma Aβ (40)—but failed to meet clinical end points. Solanezumab continues to test this hypothesis (62), which may still have hope if instituted in preclinical stages (A4 and DIAN-TU) (31,32).

**IMPORTANCE OF HIGHER DOSES**

The failure of anti-Aβ mAbs trials has raised questions about the need for higher doses. For solanezumab, the combination of insignificant efficacy and excellent safety begs the question of whether higher doses would have yielded significant effects for the primary outcomes (30) and whether these should still be considered for ongoing studies in preclinical AD (31,32). Similarly, the encouraging results with aducanumab pose a conundrum following disappointing results with other N-terminal antibodies—bapineuzumab and gantenerumab. Both antibodies share with aducanumab similar pharmacodynamic effects of fibrillar Aβ clearance on PET scans and ARIA-E, although at lower rates than aducanumab. Would higher doses of these drugs produce similar effects (35)?

**IMPORTANCE OF STAGE OF DISEASE**

An often-cited explanation for the failure of anti-Aβ immunotherapy trials is that they are set too late in the disease process (9,30). Obviously, earlier intervention with a disease-modifying treatment, including anti-Aβ mAbs, is advantageous. Less clear is whether early intervention is necessary for any treatment benefit—i.e., whether an Aβ cascade is initiated such that deterioration can no longer be slowed, or whether, in the setting of advanced Aβ deposition, modest Aβ clearance is simply irrelevant. If Aβ accumulation largely precedes cognitive impairment and is nearly complete by the dementia stage (63–65), later intervention with Aβ-lowering therapies may prove ineffective. Empirical trial evidence for this viewpoint is quite sparse and perhaps limited to the post hoc analyses from the phase 3 solanezumab trial (23) and the phase 2 crenezumab trial (38,40), suggesting clinical efficacy restricted to mild AD subgroups. Unquestionably, the need for higher doses. For solanezumab, the combination of whether higher doses would have yielded significant effects for the primary outcomes (A4 and DIAN-TU) (31,32).

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