Aducanumab (Aduhelm) was recently approved by the US Food and Drug Administration (FDA) for the treatment of Alzheimer disease (AD). The approval of aducanumab is controversial. We address why we support this decision and welcome the availability of aducanumab as an important therapy for patients with early AD.

Building on a phase 1B study that informed dose decisions and demonstrated amyloid lowering by aducanumab, Biogen launched 2 phase 3 trials: EMERGE and ENGAGE.1 These were identically designed global trials with 1,600 amyloid-positive participants with early AD in each trial. At trial initiation, participants were randomized to high dose, low dose, or placebo. A dose increase from 6 mg/kg to 10 mg/kg was made for APOE4 gene carriers midway through the trial; EMERGE participants had higher doses for longer periods as the EMERGE trial began later than the ENGAGE trial and patients had more time on the 10 mg/kg dose. An interim efficacy analysis showed no foreseeable drug–placebo difference, and both trials were stopped. Blinded data continued to accrue and the final data set at week 78 demonstrated that EMERGE met its primary outcome with a 22% drug–placebo difference in slowing on the Clinical Dementia Rating sum of boxes (CDR-SB) in the high-dose group (\(p = 0.01\)). Secondary outcomes on standard clinical trial instruments showed significant drug–placebo differences in favor of aducanumab with greater decline in the placebo group than in the aducanumab group in the course of the trial: the AD Assessment Scale–cognitive subscale showed a 27% drug–placebo difference (\(p = 0.009\)), the AD Cooperative Study activities of daily living mild cognitive impairment scale demonstrated a 40% drug–placebo difference (\(p = 0.0006\)), and the Mini-Mental State Examination exhibited an 18% difference (\(p = 0.04\)). Functional measures are used to determine the clinical meaningfulness of a treatment response; a 40% reduction in functional decline is statistically significant and clinically important. Amyloid lowering was robust and statistically significant in both dose groups. CSF biomarker analyses revealed significant dose-related reductions in hyperphosphorylated tau in the active treatment groups. Pooled EMERGE and ENGAGE data for tau PET showed increases in the placebo group and less increase or decreases in the high-dose group in several cortical regions. The ENGAGE trial showed no drug–placebo difference for the primary and secondary clinical outcomes in the final data set. There was significant amyloid lowering in both dose arms. Of ENGAGE participants who had completed 14 months of treatment at the 10 mg/kg dose (the dose recommended in the package insert), there was a 27% slowing on the CDR-SB comparable to the slowing observed in EMERGE. The principal adverse events occurring in both trials were amyloid-related imaging abnormalities occurring in 34% of those in the EMERGE high-dose group and 35.5% in the ENGAGE high-dose group and leading to 6.6% and 7.3% discontinuations in EMERGE and ENGAGE, respectively.

The FDA approved aducanumab using the accelerated approval pathway that allows marketing of a drug based on a biomarker considered reasonably likely to predict clinical benefit. Accelerated approval is accompanied by a requirement to generate additional efficacy and safety data in a postmarketing study. The biomarker considered likely to predict clinical benefit was plaque lowering demonstrated by amyloid PET.
The body of evidence supporting use of aducanumab in patients with early AD includes consistent amyloid reduction on PET imaging, slowing of decline on primary and secondary outcome measures in EMERGE, encouraging effects on biomarkers consistent with effect on biological processes linked to disease progression, and preliminary observations of slowing of cognitive decline in other trials of plaque-lowering antibodies (e.g., donanemab). The absence of disease-modifying therapies for AD supports making the therapy available on the basis of these data. Further study is required to confirm clinical benefits and to identify individuals most likely to respond.

Implementation of aducanumab will make new demands on the health care system, including clinician vigilance for patients with early AD; availability of amyloid PET, lumbar puncture, and MRI resources; and access to infusion centers. We recommend that the patients treated with aducanumab have features similar to those included in the EMERGE/ENGAGE trials: early symptomatic AD with evidence of brain amyloid. Progress in the development of a blood test for AD diagnosis would accelerate the integration of aducanumab into the diagnostic/therapeutic workflow. Successful implementation of aducanumab will require collaboration among primary care clinicians and specialists and a commitment to equity of access.

Aducanumab opens a new era in AD research and care and provides a foundation for development of progressively better therapies.

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**References**


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