Challenges in a Biological Definition of Alzheimer Disease

Jemma Hazan, MBBS, MRCPsych, iBSc, Kathy Y. Liu, Harry Costello, MBBS, MSc, MRCPsych, BSc, Jeremy D. Isaacs, MA, MBBS, MRCP, PhD, Madhav Thambisetty, MD, PhD, and Robert Howard, MD

Neurology® 2024;103:e209884. doi:10.1212/WNL.0000000000209884

Correspondence

Dr. Hazan j.hazan@ucl.ac.uk

Abstract

It has been suggested that the diagnostic landscape of Alzheimer disease (AD) is undergoing a profound transformation, marked by a shift toward a biomarker-based approach, as proposed by the Revised Criteria for Diagnosis and Staging of Alzheimer's Disease. These criteria advocate for diagnosing AD solely on biomarkers, without requiring clinical symptoms. This article explores the drivers behind this transition, primarily influenced by the Food and Drug Administration's approval of amyloid-lowering treatments. We evaluate the proposed criteria, which allow for an AD diagnosis based on amyloid "A" or phosphorylated tau "T1" positivity through surrogate amyloid PET imaging, CSF, or plasma biomarkers, and consider the arguments for and against their use. The merits of the new criteria include a clearer definition of AD, which is currently used interchangeably to refer to both the presence of neuropathology and the clinical syndrome. We argue that a purely biological definition risks a category error and emphasize the need for longitudinal data to establish the lifetime risk of dementia in amyloid-positive and tau-positive individuals. We also caution against limiting the scope of biomarker-based AD diagnosis to amyloid and tau alone. In conclusion, we recommend that the criteria remain within the research domain for the present while advocating for the considered adoption of plasma biomarkers in clinical practice.

RELATED ARTICLE

Editorial

Alzheimer Disease Is a Specific Disorder Defined by Neuropathology Detectable During Life Page e209995

Introduction

Recent years have seen rapid evolution in Alzheimer disease (AD) research. In 2021, the Food and Drug Administration (FDA) approved aducanumab to treat patients with Alzheimer disease, and in 2023, it approved lecanemab for a similar indication. That same year, at the Alzheimer's Association International Conference, the phase III clinical trial data for donanemab and data on the utility and adoption of a range of AD plasma biomarkers were presented. At that meeting, the National Institute on Aging and the Alzheimer's Association (NIA-AA) convened a working group that presented a draft proposal for revised criteria for the diagnosis of AD. Because the National Institute on Aging now serves on an advisory capacity, its cosponsorship came under scrutiny because it was seen to extend beyond its research scope. The recently published 2024 AA criteria are known as Revised Criteria for Diagnosis and Staging of Alzheimer's Disease: Alzheimer's Association Workgroup. 6-8

The NIA-AA initially published AD diagnostic guidelines in 2011. These characterized an AD disease continuum, from preclinical AD to mild cognitive impairment (MCI) due to AD to AD dementia. Around the time when these guidelines were published, in vivo biomarkers of AD (amyloid and tau) PET and CSF (amyloid and tau measures) were developed, and these allowed an antemortem diagnosis of AD neuropathologic change. As a result, the 2018 NIA-AA research criteria proposed the amyloid (A)/tau (T)/neurodegeneration(N) system, with an A+T+ biomarker profile required for a biological definition of AD. 10

From the University College London (J.H., K.Y.L., H.C., R.H.); Neuroscience & Cell Biology Research Institute (J.D.I.), St George's, University of London, United Kingdom; and Clinical and Translational Neuroscience Section (M.T.), National Institutes of Health, Bethesda, MD; Laboratory of Behavioral Neuroscience (M.T.), National Institute on Aging, Baltimore, MD.

Go to Neurology.org/N for full disclosures. Funding information and disclosures deemed relevant by the authors, if any, are provided at the end of the article.

The Article Processing Charge was funded by UCL.

 $See the \ Highlighted \ Changes \ supplement \ at \ Neurology. or \textit{g}, showing \ the \ changes \ made \ in \ this \ updated \ version.$

This is an open access article distributed under the terms of the Creative Commons Attribution License 4.0 (CC BY), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Glossary

 $A\beta$ = amyloid-beta; AD = Alzheimer disease; FDA = Food and Drug Administration; MCI = mild cognitive impairment; NIA-AA = National Institute on Aging and the Alzheimer's Association.

The 2024 AA criteria propose the adoption of a purely biologically based construction of AD to inform clinical practice and research.⁵ Based on these criteria, a diagnosis of AD can now be made based on either amyloid "A" or phosphorylated tau "T₁" positivity using surrogate amyloid PET imaging or CSF or plasma biomarkers (p-tau217). The "N" classification was removed from the core biomarker criteria. Crucially, although the criteria highlight the value of clinical judgement for the diagnosis, the biological AD definition does not require the presence of clinical symptoms. The criteria do not advocate for the screening of preclinical disease, although those with subjective cognitive concerns may be tested. A diagnosis of AD, which can be made in the absence of clinical symptoms, will have significant implications for clinical practice and patient care. In addition, 2024 draft US FDA guidance proposes that a change in a surrogate biomarker concentration alone can be accepted as evidence of efficacy in the licensing of treatments for asymptomatic preclinical AD individuals. 11 This could mean that a biomarker result could be used to both diagnose AD (2024 AA criteria) and as evidence of treatment response (2024 FDA draft criteria), in an asymptomatic individual.

Proposed Reasoning for Updating the Criteria

The 2024 AA criteria can be seen as an inevitable outcome of FDA approval of amyloid-lowering treatments based at least partly on surrogate biomarker endpoints and the proposed sole use of biomarker concentrations as indicators of treatment efficacy in asymptomatic trial participants. 11-13 The criteria distinguish between an asymptomatic AD "disease" phase and an "illness" stage, when symptoms are evident. Proponents of the criteria assert that this approach aligns with other areas of medicine, particularly oncology, where surrogate biomarkers are used to screen and diagnose disease before symptom development. With the advent of treatments targeting AD pathology, the new definition seeks to conceptually align the diagnosis of AD with an integrated set of biological markers and clinical stages that begin in the presymptomatic state. This approach aims to improve diagnostic precision, identify eligible participants for clinical trials (including those without symptoms), and target and monitor treatment responses.

Arguments for and Against the Proposed Draft Criteria

One advantage of a purely biological definition is that it clearly conceptualizes AD as a distinct pathologic process,

separate from the clinical manifestations of the various diseases that cause cognitive decline. This strictly biological definition may help to avoid the confusion caused by the term "Alzheimer disease," which can refer to both the underlying biology and the dementia syndrome. If abnormal amyloid and tau accumulation are risk factors for developing Alzheimer dementia, it would be important to identify and mitigate this at the earliest possible timepoint. This could be comparable with identifying and treating cerebrovascular disease risk factors, such as hypertension. Ongoing studies such as AHEAD 3-45 Study (BAN2401-G000-303) may provide valuable insights into treatment outcomes in this asymptomatic amyloid-positive population.¹⁴ However, secondary prevention studies may not have a long enough period of follow-up or adequate sample size to provide meaningful information on efficacy in dementia risk reduction. 15 A consensus biological definition of AD could set the stage for earlier and more precise identification of clinical trial participants. The second advantage is that biomarker use in clinical trial recruitment has helped to increase diagnostic validity within those trials. 16 Patients and carers could use such information to facilitate earlier care planning and inform decisions to enter presymptomatic treatment trials.

Despite these putative advantages of using a biological definition of AD in clinical trials and clinical care, several questions arise regarding the widespread use of the proposed AA criteria. First, does the biological definition of AD fall into the trap of a category error? A category error is made when we assign a problem to a category that is not appropriate for solving it. An example of a category trap is the use of blood glucose concentrations to define type II diabetes.¹⁷ Because biomarkers such as blood glucose level are continuously distributed variables, it is difficult to establish a clear threshold that distinguishes between healthy and potentially harmful levels that warrant a diagnosis and subsequent intervention. Defining such thresholds can prove elusive, particularly when considering the heterogeneity of the populations, and contributory environmental factors that affect the risk of developing complications of diabetes. Establishing thresholds introduce an intermediate or "grey zone" for biomarker cut points, which require clinical interpretation, integrating the result in the context of the history, examination, and clinical judgment. By equating pathologic changes with the disease itself and relying on rigid biomarker causal pathways, using fixed cut points, we would move away from appreciating the complex interplay of etiologies that drive the dementia process.18

Further work is required to establish the lifetime risk of dementia in amyloid-positive and tau-positive individuals, particularly amyloid-beta (Aβ) biomarker positivity is not always deterministically associated with a dementia outcome 19 and tau is more strongly associated with cognitive status and neurodegeneration than amyloid. 18 Although the presence of increased brain amyloid is necessary for the propagation of tau beyond the medial temporal lobe, we know that AD Aß biomarker positivity alone is not sufficient. 20,21 This may be due to a substantial lag before tauopathy spreads beyond the medial temporal lobe. For example, young-onset dysexecutive AD is characterized by a high tau load during the MCI phase and will progress faster than late-onset limbic-predominant AD.²² Consequently, as late-onset limbic-predominant AD typically happens later in life, many people with elevated amyloid will die of other causes before this progression takes place.

There is a danger of narrowing of the scope of biomarkerbased AD diagnosis with a focus purely on the amyloid and tau hypotheses. However, these processes are not fully elucidated with questions remaining over the causal relationship between A β or tau and AD.²³ This could detract from other important contributory pathologic mechanisms and that dementia in most older people is characterized by copathology, including TDP-43, Lewy bodies, and cerebrovascular changes.^{24,25} Amyloid and tau imaging have revealed the constraints of solely focusing on plaque and tangle pathology. This limitation has led to the recognition of other contributors, such as limbic TDP-43 proteinopathy.²⁶ The heterogeneous nature of late onset AD with both a long natural history of the disease and a variation in presentation may mean that the specificity of these biomarkers for AD dementia is uncertain.

Technological progression should not automatically be assumed to be beneficial for the health and well-being of populations. Are we risking overmedicalization and introducing iatrogenic harm? The balance between risks and potential benefits of a biological AD diagnosis will differ between AD severity stages, for example, compared with patients with 'early' symptomatic AD, the potential clinical benefits of amyloid-lowering agents in asymptomatic individuals are even more uncertain, mainly owing to limitations in interpreting trial outcomes in the context of a long natural disease course. ^{15,27}

The prevalence of amyloid positivity in asymptomatic individuals aged 80–89 years is around 40%, and the same study estimated that 22% of all adults aged older than 50 years will be amyloid-positive. ²⁸ This means that 26 million people in the United States alone may be amyloid-positive. According to the new criteria, such individuals can now be classified as having AD. Because the predictive accuracy of amyloid positivity for dementia is uncertain, ¹⁹ diagnosing them with AD will introduce unintended harms including distress for patients and carers. Who will benefit from the

potentially significant expansion in the number of people who qualify for a drug treatment? Focusing resources on amyloid-lowering and tau-lowering treatments will divert funds from already underfunded effective evidence-based psychosocial interventions and social support for patients and caregivers. ^{29,30} We do not yet have effective treatments that can arrest the disease process before it causes symptoms; if we did, we would support these biologically defined criteria

Looking Toward the Future

Defining the presence of AD purely on biomarker evidence of amyloid and tau positivity does not reflect the complexity of the dementia construct, and we are concerned that such an approach may cause more harm than good. Such an approach has value in research settings, where it can be used to frame hypotheses, but it is not yet appropriate for use in clinical practice. We do not propose rejecting the criteria in its entirety. The incorporation of plasma biomarkers may have diagnostic utility for diagnosing people with symptomatic AD.4 However, it is important that the criteria incorporate the full range of pathologic mechanisms that contribute to the AD dementia process. A biologically reductionist approach cannot capture the multiple processes involved in Alzheimer dementia, including the role of psychological and social factors. Rather than eliminating these from our understanding of dementia, we must embrace complexity. This will ensure that we do not invest false therapeutic certainty in the dominant hypotheses of the day. To date, there is no evidence that targeting tau is associated with clinical benefit in people with AD. Indeed, the modest clinical outcome data from A\beta therapy trials show the danger of assuming that risk factors or biomarkers for a disease are the disease.^{2,18}

The proposed narrowly defined biological model of sporadic AD is currently insufficient, unrelated to etiologic complexity and a long pathologic process, and overshadows other biopsychosocial models³¹ and risk-modification strategies.³² It is important to maintain a clear distinction between the definition of AD pathology and the clinical syndrome of AD dementia. If there really was utility in defining an illness on the basis of a minimal set of features that are (1) present in all patients and (2) potentially modifiable, then the most propitious definition of AD dementia would be as a disorder of social function because it is the quality and quantity of social support that can determine a patient's quality of life and speed of disease progression.

Study Funding

The authors report no targeted funding.

Disclosure

K. Liu was funded by the UK Medical Research Council (MR/S021418/1); H. Costello was supported by a Wellcome

Trust Clinical Training Fellowship; R. Howard was supported by University College London Hospitals' National Institute for Health Research (NIHR) Biomedical Research Centre. All other authors report no relevant disclosures. Go to Neurology.org/N for full disclosures.

Publication History

Received by *Neurology* April 19, 2024. Accepted in final form July 31, 2024. Submitted and externally peer reviewed. The handling editor was Editor-in-Chief José Merino, MD, MPhil, FAAN.

Appendix Authors		
Name	Location	Contribution
Jemma Hazan, MBBS, MRCPsych, iBSc	University College London, United Kingdom	Drafting/revision of the manuscript for content, including medical writing for content; study concept or design
Kathy Y. Liu	University College London, United Kingdom	Drafting/revision of the manuscript for content, including medical writing for content; study concept or design
Harry Costello, MBBS, MSc, MRCPsych, BSc	University College London, United Kingdom	Drafting/revision of the manuscript for content, including medical writing for content; study concept or design
Jeremy D. Isaacs, MA, MBBS, MRCP, PhD	Neuroscience and Cell Biology Research Institute, St George's, University of London, United Kingdom	Drafting/revision of the manuscript for content, including medical writing for content; study concept or design
Madhav Thambisetty, MD, PhD	Clinical and Translational Neuroscience Section, National Institutes of Health, Bethesda, MD; Laboratory of Behavioral Neuroscience, National Institute on Aging, Baltimore, MD	Drafting/revision of the manuscript for content, including medical writing for content; study concept or design
Robert Howard, MD	University College London, United Kingdom	Drafting/revision of the manuscript for content, including medical writing for content; study concept or design

References

- Tolar M, Abushakra S, Hey JA, Porsteinsson A, Sabbagh M. Aducanumab, gantenerumab, BAN2401, and ALZ-801—the first wave of amyloid-targeting drugs for Alzheimer's disease with potential for near term approval. Alzheimers Res Ther. 2020; 12:95. doi:10.1186/s13195-020-00663-w
- Sims JR, Zimmer JA, Evans CD, et al. Donanemab in early symptomatic Alzheimer disease: the TRAILBLAZER-ALZ 2 randomized clinical trial. *JAMA*. 2023;330(6): 512-527. doi:10.1001/jama.2023.13239
- Van Dyck CH, Swanson CJ, Aisen P, et al. Lecanemab in early Alzheimer's disease. N Engl J Med. 2023;388(1):9-21. doi:10.1056/NEJMoa2212948
- Hansson O, Edelmayer RM, Boxer AL, et al. The Alzheimer's Association appropriate use recommendations for blood biomarkers in Alzheimer's disease. Alzheimers Dement. 2022;18(12):2669-2686. doi:10.1002/alz.12756
- NIA-AA. Revised Criteria for Diagnosis and Staging of Alzheimer's Disease: Alzheimer's Association Workgroup. 2023. Accessed December 14, 2023. alz.org/media/Documents/ scientific-conferences/Clinical-Criteria-for-Staging-and-Diagnosis-for-Public-Comment-Draft-2.pdf?_gl=1*exh9z4*_ga*MjASMDIzNzcSMy4xNzAwMjQyMDgy*_ga_ QSFTKCEH7C*MTcwMTEyMjkSMC40LjEuMTcwMTEyMzIyOS42MC4wLjA*_ ga_9JTEWVX24V*MTcwMTEyMjkSMC40LjEuMTcwMTEyMzIyOS42MC4wLjA

- The American Geriatrics Society. American Geriatrics Society Response—Draft NIA-AA Revised Clinical Criteria for Alzheimers Disease. 2023. Accessed June 9, 2024. americangeriatrics.org/sites/default/files/inline-files/AGS%20Comments%20on% 20Draft%20NIA-AA%20Revised%20Clinical%20Criteria%20for%20Alzheimer's% 20Disease%20(8%2016%2023)%20FINAL 0.pdf
- Gleckman H. NIH Steps Back From Development of New Alzheimer's Diagnostic Standards. Forbes; 2023.
- Jack CR Jr, Andrews JS, Beach TG, et al. Revised criteria for diagnosis and staging of Alzheimer's disease: Alzheimer's association workgroup. Alzheimers Dement. 2024; 20(8):5143-5169. doi:10.1002/alz.13859
- Jack CR Jr, Albert MS, Knopman DS, et al. Introduction to the recommendations from the National Institute on Aging-Alzheimer's Association workgroups on diagnostic guidelines for Alzheimer's disease. Alzheimers Dement. 2011;7(3):257-262. doi:10.1016/j.jalz.2011.03.004
- Jack CR Jr, Bennett DA, Blennow K, et al. NIA-AA research framework: toward a biological definition of Alzheimer's disease. Alzheimers Dement. 2018;14(4):535-562. doi:10.1016/j.jalz.2018.02.018
- U.S. Department of Health and Human Services, Food and Drug Administration, Center for Drug Evaluation and Research (CDER), Center for Biologics Evaluation and Research (CBER). Early Alzheimer's Disease: Developing Drugs for Treatment Guidance for Industry. 2024. Accessed March 28, 2024. fda.gov/regulatory-information/search-fdaguidance-documents/early-alzheimers-disease-developing-drugs-treatment
- Liu KY, Schneider LS, Howard R. The need to show minimum clinically important differences in Alzheimer's disease trials. *Lancet Psychiatry*. 2021;8(11):1013-1016. doi: 10.1016/S2215-0366(21)00197-8
- Dunn B, Stein P, Cavazzoni P. Approval of aducanumab for Alzheimer disease—the FDA's perspective. JAMA Intern Med. 2021;181(10):1276-1278. doi:10.1001/jamainternmed.2021.4607
- Rafii MS, Sperling RA, Donohue MC, et al. The AHEAD 3-45 study: design of a prevention trial for Alzheimer's disease. Alzheimers Dement. 2023;19(4):1227-1233. doi:10.1002/alz.12748
- Liu KY, Thambisetty M, Howard R. How can secondary dementia prevention trials of Alzheimer's disease be clinically meaningful? Alzheimers Dement. 2023;19(3): 1073-1085. doi:10.1002/alz.12788
- Salloway S, Sperling R, Fox NC, et al. Two phase 3 trials of bapineuzumab in mild-to-moderate Alzheimer's disease. N Engl J Med. 2014;370(4):322-333. doi:10.1056/neimoa1304839
- Gale EA. Is type 2 diabetes a category error? Lancet. 2013;381(9881):1956-1957. doi: 10.1016/S0140-6736(12)62207-7
- Korczyn AD, Grinberg LT. Is Alzheimer disease a disease? Nat Rev Neurol. 2024; 20(4):245-251. doi:10.1038/s41582-024-00940-4
- Brookmeyer R, Abdalla N. Estimation of lifetime risks of Alzheimer's disease dementia using biomarkers for preclinical disease. Alzheimers Dement. 2018;14(8):981-988. doi: 10.1016/j.jalz.2018.03.005
- Pooler AM, Polydoro M, Maury EA, et al. Amyloid accelerates tau propagation and toxicity in a model of early Alzheimer's disease. Acta neuropathologica Commun. 2015; 3:14-11. doi:10.1186/s40478-015-0199-x
- Dubois B, von Arnim CA, Burnie N, Bozeat S, Cummings J. Biomarkers in Alzheimer's disease: role in early and differential diagnosis and recognition of atypical variants. Alzheimers Res Ther. 2023;15(1):175. doi:10.1186/s13195-023-01314-6
- Frontzkowski L, Ewers M, Brendel M, et al. Earlier Alzheimer's disease onset is associated with tau pathology in brain hub regions and facilitated tau spreading. Nat Commun. 2022;13(1):4899. doi:10.1038/s41467-022-32592-7
- Neve RL, Robakis NK. Alzheimer's disease: a re-examination of the amyloid hypothesis. Trends Neurosci. 1998;21(1):15-19. doi:10.1016/s0166-2236(97)01168-5
- Govindpani K, McNamara LG, Smith NR, et al. Vascular dysfunction in Alzheimer's disease: a prelude to the pathological process or a consequence of it? J Clin Med. 2019; 8(5):651. doi:10.3390/jcm8050651
- Bayram E, Shan G, Cummings JL. Associations between comorbid TDP-43, Lewy body pathology, and neuropsychiatric symptoms in Alzheimer's disease. J Alzheimers Dis. 2019;69(4):953-961. doi:10.3233/JAD-181285
- Nelson PT, Dickson DW, Trojanowski JQ, et al. Limbic-predominant age-related TDP-43 encephalopathy (LATE): consensus working group report. *Brain*. 2019; 142(6):1503-1527. doi:10.1093/brain/awz099
- Liu KY, Walsh S, Brayne C, Merrick R, Richard E, Howard R. Evaluation of clinical benefits of treatments for Alzheimer's disease. *Lancet Healthy Longev.* 2023;4(11): e645-e651. doi:10.1016/S2666-7568(23)00193-9
- Roberts RO, Aakre JA, Kremers WK, et al. Prevalence and outcomes of amyloid positivity among persons without dementia in a longitudinal, population-based setting. JAMA Neurol. 2018;75(8):970-979. doi:10.1001/jamaneurol.2018.0629
- McDermott O, Charlesworth G, Hogervorst E, et al. Psychosocial interventions for people with dementia: a synthesis of systematic reviews. *Aging Ment Health*. 2019; 23(4):393-403. doi:10.1080/13607863.2017.1423031
- Keogh F, Pierse T, Challis D, O'Shea E. Resource allocation across the dementia continuum: a mixed methods study examining decision making on optimal dementia care among health and social care professionals. BMC Health Serv Res. 2021;21:243. doi:10.1186/s12913-021-06230-9
- Spector A, Orrell M. Using a biopsychosocial model of dementia as a tool to guide clinical practice. Int Psychogeriatr. 2010;22(6):957-965. doi:10.1017/S1041610210000840
- Livingston G, Huntley J, Sommerlad A, et al. Dementia prevention, intervention, and care: 2020 report of the Lancet Commission. *Lancet*. 2020;396(10248):413-446. doi: 10.1016/S0140-6736(20)30367-6