Alzheimer's disease



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In this Seminar, we highlight the main developments in the field of Alzheimer's disease. The most recent data indicate that, by 2050, the prevalence of dementia will double in Europe and triple worldwide, and that estimate is 3 times higher when based on a biological (rather than clinical) definition of Alzheimer's disease. The earliest phase of Alzheimer's disease (cellular phase) happens in parallel with accumulating amyloid β , inducing the spread of tau pathology. The risk of Alzheimer's disease is 60–80% dependent on heritable factors, with more than 40 Alzheimer's disease-associated genetic risk loci already identified, of which the *APOE* alleles have the strongest association with the disease. Novel biomarkers include PET scans and plasma assays for amyloid β and phosphorylated tau, which show great promise for clinical and research use. Multidomain lifestyle-based prevention trials suggest cognitive benefits in participants with increased risk of dementia. Lifestyle factors do not directly affect Alzheimer's disease pathology, but can still contribute to a positive outcome in individuals with Alzheimer's disease. Promising pharmacological treatments are poised at advanced stages of clinical trials and include anti-amyloid β , anti-tau, and anti-inflammatory strategies.

Introduction

Alzheimer's disease is the main cause of dementia and is quickly becoming one of the most expensive, lethal, and burdening diseases of this century. Since the Seminar published in 2016,² important developments have taken place in the understanding of the underlying pathology, the recognition of multiple causative and protective genes, the identification of new blood-based and imaging biomarkers, and the first cautious signals of positive effects of disease-modifying treatments and lifestyle interventions. The aim of this new Seminar is to provide the reader with up to date insight into the field of Alzheimer's disease.

Clinical signs and symptoms

Three cases, in panel 1 (see also figure 1), illustrate the clinical spectrum of Alzheimer's disease. Case A highlights Alzheimer's disease that is determined genetically, as per the ongoing global initiatives of the Dominantly Inherited Alzheimer Network and Alzheimer Prevention Initiative and their associated clinical trials. Case B represents a language variant of Alzheimer's disease, usually occurring at a younger age (under 70 years), illustrating the difficulty in recognising Alzheimer's disease in those for whom memory problems are not the first and most prominent feature. Case C is a typical amnestic variant, more commonly seen in patients older than 70 years, illustrating the growing population affected by Alzheimer's disease and dementia: older individuals often living alone, and increasingly dependent on others for care.

Diagnostic criteria: from clinical, to clinical and biological, to biological

The diagnosis of Alzheimer's disease has gone from a purely pathological one, in the days of Alois Alzheimer (1864–1915), to a clinical, exclusionary approach in 1984. The clinical diagnosis was based on the criteria defined by the National Institute of Neurological and Communicative Disorders and Stroke and the Alzheimer's Disease and Related Disorders Association,³ via a combined clinical and biological approach developed by the International

Working Group^{4,5} and subsequent efforts by the National Institute on Aging and the Alzheimer's Association working groups,⁶ incorporating biomarkers to make the categorisation of Alzheimer's disease purely biological.⁷ Initially, the diagnosis of Alzheimer's disease was restricted to the stage of dementia, a clinical syndrome characterised by substantial progressive cognitive impairment affecting several domains, or neurobehavioral symptoms of enough severity to cause evident functional impact on daily life. A person with dementia is no longer fully independent, and this loss of independence is the primary feature differentiating dementia from mild cognitive impairment.⁸

Given the developments in the biomarker field and the desire to make them usable in a diagnostic setting, Jack and colleagues grouped the biomarkers into A (amyloid), T (phosphorylated tau), and N (neurodegeneration, measured by total tau where applicable): the ATN framework (appendix p 1). In this research framework, the diagnosis of Alzheimer's disease is defined by the presence of amyloid β and phosphorylated tau. The

Search strategy and selection criteria

Between Dec 1, 2019, and Sept 1, 2020, we searched the Cochrane Library for articles published exclusively in English during 2010–15, PubMed for articles published during 2016–20, and Embase for articles published during 2016–20. We used the search term "Alzheimer's disease" in combination with the following: "pathology", "imaging", "diagnosis", "therapy", "trials", "epidemiology", "CSF", "genetics", and "biomarkers". We largely selected publications from the past 5 years, and especially focused on changes that occurred after the publication of the previous Seminar in 2016.¹ We also searched the reference lists of articles identified by this search strategy and selected those that were judged relevant. Review articles and book chapters are cited to provide readers with references for more details than this Seminar can include.

Lancet 2021; 397: 1577-90

Published Online March 2, 2021 https://doi.org/10.1016/ S0140-6736(20)32205-4

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See Online for appendix

Panel 1: Case vignettes

Mrs A, aged 42 years, a successful manager of an IT company, presents at the Alzheimer Centre Amsterdam because of self-perceived memory loss and loss of oversight and multitasking abilities. She recognises these complaints all too well because her mother had Alzheimer's disease for 5 years, until her death at the age of 47 years. Two of her four brothers also had Alzheimer's disease, and had been tested and found to be carriers of a PSEN1 mutation. Although she has not been tested herself, she always felt she would be a carrier and subsequently chose not to have children. She asked for a full evaluation because she wanted to have the option of participating in a clinical trial programme. Her Mini-Mental State Exam score was 27/30 and her Montreal Cognitive Assessment score was 24/30. Given her age, these scores suggest mild memory and executive disturbances, which were confirmed by neuropsychological testing. A brain MRI showed no abnormalities. Cerebrospinal fluid biomarker values were 750 pg/mL for amyloid \(\beta 42, 335 pg/mL \) for tau, and 35 pg/mL for phosphorylated tau 181, all in the abnormal range. Serum neurofilament light chain value was 25 pg/mL, which is abnormal for her age, according to in-house defined reference curves. APOE status was $\varepsilon 3/\varepsilon 4$. All these biomarker values indicate the presence of Alzheimer's disease pathology and onset in a clinically mildly affected patient. Genetic testing confirmed the presence of the same PSEN1 mutation carried by her brothers. She was informed about the diagnosis, followed up at 6 month intervals at the centre, and put on the list for a clinical trial within the Dominantly Inherited Alzheimer Network Trials Unit programme. She informed her colleagues at work and agreed to have regular meetings with the company's physician. Mr B, aged 62 years, is a high school teacher who presented to the neurologist with gradually progressive difficulty finding words and understanding sentences, and slight memory loss. He had visited another neurologist because of suspicion of a

presence of amyloid β (regardless of the presence of phosphorylated tau and neurodegeneration) is termed Alzheimer's pathological change, basing the research diagnosis of Alzheimer's disease on biomarker evidence only. Clinical stages can range from cognitively normal to mild cognitive impairment and dementia, stressing the continuum of Alzheimer's disease, which spans a period of years. The ATN framework underpins the importance of amyloid β and tau as the defining characteristics of Alzheimer's disease, consequently proposing that Alzheimer's disease can be diagnosed by biomarkers only, and definitively distinguishing between

vascular event, but a brain MRI showed no abnormalities.

and the Montreal Cognitive Assessment score was 24/30,

routine neurological and laboratory investigations.

On examination, his Mini-Mental State Exam score was 25/30

both within normal range for his age, with normal findings at

Neuropsychological and detailed language assessment revealed

a decrease in fluency, naming, and repetition of long sentences.

Review of the MRI showed slight asymmetry of the temporal lobes, with grade 2 hippocampal atrophy on the left side and grade 1 hippocampal atrophy on the right side, without any other abnormalities (figure 1B). Because of his young age, and his and his family's desire to obtain a firm diagnosis to plan ahead and make proper adjustments to his working life, an amyloid-PET scan was done and showed diffuse cortical uptake of the ligand (figure 1A). As part of a research project, a tau-PET scan was done and showed left-temporal abnormal tau deposition (figure 1C). A diagnosis of logopenic variant of Alzheimer's disease was made. Lifestyle advice was given and regular visits to a speech therapist were offered. Given the diagnosis and the perceived grim future, as well as the high demands of his job on his language skills, he decided to take sick leave from his job.

Mrs C, aged 78 years, lives independently on her own after being widowed 6 years ago. She was known to her general practitioner with controlled hypertension and moderate heart failure, for which she takes medication. Her son lives abroad and her daughter lives 100 km away. Both have demanding jobs and young children. During telephone and Skype calls, her children noticed increasing forgetfulness and one of the neighbours had recently informed the daughter that her mother mixed up the days, forgot to eat, and was not able to take good care of herself anymore. The daughter accompanied her mother to the Alzheimer Centre Amsterdam on referral by the general practitioner, who had initially dismissed the worries of the daughter. On examination by a geriatrician, she was found to be malnourished and underweight. The Mini-Mental State Exam score was 17/30 and a brief neuropsychological test battery showed scores below the norm for memory and executive function. Her score on the Amsterdam Instrumental Activities of Daily Living test³ was 58, indicating severe impairment. An MRI showed a medial temporal atrophy score of 2 bilaterally, and moderate to severe white matter changes (Fazekas score 2). A diagnosis of mild to moderate dementia due to Alzheimer's disease with some vascular contribution was made, and a case manager was assigned to organise and supervise care to have her stay at home as long as possible. Vascular risk factors were checked and cholinesterase inhibitor therapy was started.

the concepts of Alzheimer's disease and dementia (figure 2).

Despite the critique that other key causes of dementia, in particular vascular disease, were omitted, 9 the authors of the ATN framework argued that dementia has multiple underlying pathologies, of which Alzheimer's disease is one, but Alzheimer's disease is defined by the presence of amyloid β and tau (acknowledging that many other pathologies can also be present in these patients). 10 The large number of ATN categories, combined with the fact that other pathologies are not evaluated in the scheme, makes the ATN approach not

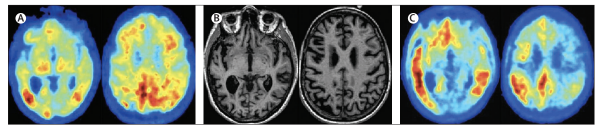


Figure 1: Imaging findings of a case similar to patient B's case in panel 1

(A) Amyloid Pittsburgh compound B-PET scan showing amyloid deposition predominantly in the posterior cingulate region. (B) T1-weighted MRI images showing generalised cortical atrophy, left to right. (C) Tau-PET image using AV1451 tracer, showing left-sided inferotemporal lobe, parietal, and mild posterior cingulate deposition of tau. Image courtesy of Rik Ossenkoppele and Gil Rabinovici.

yet suitable for clinical practice." In addition, there are operational limitations to defining A, T, and N positivity or negativity, such as some biomarkers not having established cutoff points, and different biomarkers being combined in one category. Although the AT or ATN approach is the cornerstone of current trials of disease-modifying interventions in Alzheimer's disease, clinical diagnosis still rests on the criteria set by the National Institute on Aging in 2011.

The ATN framework clearly paves the way for a diagnosis before the stage of Alzheimer's disease-associated dementia, and it makes individualised risk-profiling for patients with mild cognitive impairment feasible.13 However, a clinical encounter study evaluating doctor-patient communication in memory clinics showed that clinicians are reluctant to share specific prognostic information with patients with mild cognitive impairment.14 In the context of predementia diagnosis, subjective cognitive decline is even more challenging. A recent Personal View provides a clinical characterisation of subjective cognitive decline, and attempts to provide clinicians with guidance on how to deal with this decline (which might or might not be attributable to underlying Alzheimer's disease).15 At a group level, ATN biomarkers clearly predict incident dementia in subjective cognitive decline, but individualised risk modelling remains challenging. 16,17 In a Delphi study to identify topics most relevant to discuss in the diagnostic process, patients and caregivers indicated that they value precise and specific information, even when it does not provide complete certainty.18 Tools to support decision making and communication about Alzheimer's disease diagnosis, such as ADappt,19 are urgently needed.

Epidemiology

Incidence and prevalence

In 2018, Alzheimer's Disease International estimated a dementia prevalence of about 50 million people worldwide, projected to triple in 2050, with two-thirds living in low-income and middle-income countries.²⁰ The most recent data estimate that dementia prevalence in Europe will double by 2050.¹ Accumulating evidence suggests that the incidence of dementia is declining in high-income countries,²¹ although evidence for a decline in prevalence is less convincing.²²

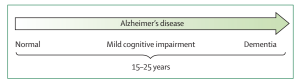


Figure 2: Alzheimer's disease is a continuum

The arrow points to the continuum of Alzheimer's disease, stretching over a period of 15–25 years, in which Alzheimer's disease pathology can be present without any symptoms via a stage of mild cognitive impairment leading up to overt dementia, illustrating that dementia is the end result of a long-time presence of Alzheimer's disease pathology. Not every patient will necessarily follow this path by definition. Note: between normal and mild cognitive impairment, patients can experience subjective complaints, but not all complaints are early signs of dementia and the predictive value of having complaints for dementia is unknown.

Mortality

The relatively stable prevalence despite decreasing incidence could be explained by a long disease duration, although studies on mortality do not support this notion. A US-based study evaluating survival after a dementia diagnosis in almost 60 000 individuals reported survival times of 3-4 years.23 In an European, memory clinicbased cohort, median survival time was 6 years after a diagnosis of Alzheimer's disease dementia (median 6.2 years [range 6.0-6.5]).²⁴ This estimate coincides with a multicentre study that provided estimates of duration not only of the dementia stage, but also of the prodromal (mild cognitive impairment) and of preclinical disease stage of Alzheimer's disease.25 For an individual aged 70 years, duration estimates are 10 years for the preclinical stage, 4 years for the prodromal stage, and 6 years for the dementia stage of Alzheimer's disease, totalling 20 years. A first attempt at estimating prevalence on the basis of a biological (rather than clinical) definition showed that, at the age of 85 years, the prevalence of biologically defined Alzheimer's disease is 3 times higher than that of clinically defined Alzheimer's disease.26

Risk factors for dementia and Alzheimer's disease

The strongest risk factors for Alzheimer's disease are advanced age (older than 65 years, although this is not a fixed definition) and carrying at least one $APOE\ \epsilon 4$ allele. Moreover, women are more likely to develop Alzheimer's disease than are men, especially after the age of 80 years. ²⁰

For **ADappt** see www.adappt. health Women are also more likely to have a higher tau load, despite having a similar amyloid β burden.^{28,29} In addition, cardiovascular risk factors and an unhealthy lifestyle have been associated with an increased risk of dementia. The Lancet Commission on Dementia Prevention estimated that 12 modifiable risk factors together account for roughly 40% of the worldwide risk of any type of dementia.30 These estimates illustrate that prevention by intervening on modifiable risk factors is of great relevance, even if most of the dementia burden cannot be prevented via a lifestyle-intervention approach. However, evidence suggests that vascular risk factors do not increase the risk of Alzheimer's disease pathology as measured by cerebrospinal fluid biomarkers or PET.31-33 This evidence implies that lifestyle and vascular risk factors contribute to dementia, but not via the Alzheimer's disease pathway.

Genetics

Causative and risk genes

Studies of twins showed that the risk of Alzheimer's disease is 60-80% dependent on heritable factors.34 The common APOE E4 allele explains a substantial part of, but does not completely account for the heritability of, Alzheimer's disease.35,36 Large genome-wide association studies have been done to identify novel genetic variants in Alzheimer's disease, the latest of which to date investigated about 150000 people with Alzheimer's disease and age-matched controls, and more than 300000 people with a proxy-phenotype Alzheimer's disease (parental history of Alzheimer's disease) and controls (no parental history of Alzheimer's disease), which increased the number of Alzheimer's diseaseassociated risk alleles to more than 40.37 However, although the common APOE E4 risk allele is associated with an estimated 3-4 times increased risk of Alzheimer's disease across different genome-wide association studies, other Alzheimer's disease risk alleles are associated with much smaller contributions to the total disease risk (odds ratio between 1.05 and 1.2; figure 3B).37

Based on the presence or absence of these risk alleles in the genome of an individual, a polygenic risk score can be calculated, which is currently able to distinguish between patients with Alzheimer's disease and controls with 75-85% accuracy.38,39 Although the bulk of this accuracy can be ascribed to the APOE E4 allele, the 40 or so other variants also collectively contribute substantially to Alzheimer's disease risk.27 Functional annotation of these risk loci indicate that, next to amyloid β metabolism, the modulation of the immune response, cholesterol, lipid dysfunction, endocytosis, and vascular factors play a role in the development of Alzheimer's disease. 40-45 Next-generation sequencing techniques have shown rare protein-damaging variants in the SORL1,46 ABCA7,47 and TREM2 genes. 48,49 These findings suggest that the intact protein products of these genes are essential in maintaining brain health (figure 3A).

Protective genes

The identification of risk-increasing genetic variants has fuelled the interest in the detection of protective genetic variants (figure 3C). Carriers of the protective APOE E2 allele have an estimated 2 times decreased lifetime risk of Alzheimer's disease compared with noncarriers,50 which translates into an exceptionally low likelihood of Alzheimer's disease for homozygous APOE ε2 allele carriers. 51 The discovery of the rare Ala673Thr Icelandic protective mutation of APP52 was associated with prolonged cognitive health. Similarly, compared with middle-aged individuals, a rare Pro522Arg amino acid change in the PLCG2 gene was associated with a near 2 times reduced risk of Alzheimer's disease⁵³ and other types of dementia, and with a 2.3 times increased chance of reaching 100 years in cognitive health.54,55 Genetic resilience was even reported in a person with a PSEN1 mutation who lived beyond the age of onset of symptoms common in her family, due to a homozygous rare protective variant in the APOE ε3 allele (Christchurch mutation). 56 Variants in the klotho longevity gene were associated with a similar effect.⁵⁷ Such protective genetic variants hold great promise in Alzheimer's disease research, as they might pinpoint mechanistic processes protecting cognitive health.

Pathophysiology

Basic scientists designate the preclinical phase of Alzheimer's disease as the cellular phase. Alterations in neurons, microglia, and astroglia drive the insidious progression of the disease before cognitive impairment is observed. ⁵⁸ Neuro-inflammation, ⁵⁹ alterations in the vessels, ^{60,61} ageing, ⁶² and dysfunction of the glymphatic system ⁶³ act upstream or in parallel to accumulating amyloid β in this cellular disease landscape. Amyloid β induces, via an unknown way, the spread of tau pathology, ⁶⁴ which is associated with the appearance of necroptosis markers in neurons displaying granulo-vacuolar degeneration. ⁶⁵

Single-cell transcriptome analysis has elucidated the microglia response.66 APOE and TREM2, two major Alzheimer's disease risk genes, are important parts of this response. 66-68 ApoE binds to amyloid β plaques, 69 and the Alzheimer's disease-associated genetic variants of TREM2 Arg47His, Arg62His, and Asp87Asn decrease binding of TREM2 to ApoE (figure 3).70 Several other proteins linked to genetic risk of Alzheimer's disease, such as SHIP1, CD2AP, RIN3, BIN1, PLCG2, CASS4, and PTKB2 act presumably downstream of ApoE and TREM2 signalmodulating endocytosis, motility, and phagocytosis in microglia (figure 4). CD33 acts in opposition to TREM2,77 and MS4A4A modulates the secretion of soluble TREM2 protein.78 The fact that so many Alzheimer's disease risk genes converge on microglial response pathways indicates their central role in the disease pathogenesis. However, further research is needed to elucidate whether the microglia response is to amyloid β plaques only,⁷⁶ or

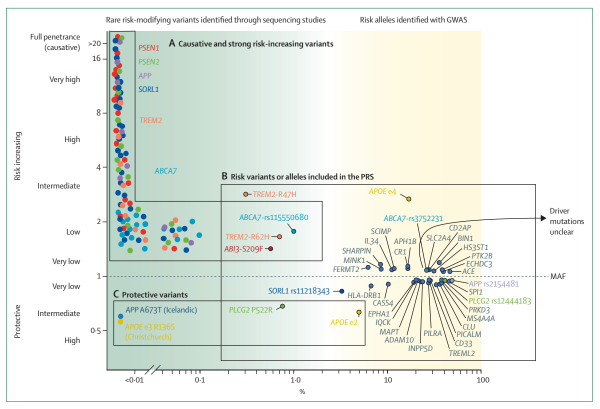


Figure 3: The genetic landscape of Alzheimer's disease

MAF (x-axis) is the frequency at which a non-reference (variant) allele occurs in the population. Variant carriers with OR=1 and non-carriers have the same odds of developing Alzheimer's disease, variants with OR >1 are associated with an increased risk of Alzheimer's disease, and variants with OR <1 are associated with a protective effect (y-axis). (A) Causative or strong risk increasing variants. A schematic representation of individual rare variants for which ORs cannot be estimated due to extreme variant rareness. Linkage studies in large pedigrees indicate that specific rare variants in PSEN1, PSEN2, and APP cause autosomal dominant Alzheimer's disease, in some cases with age at onsets as early as 40 years old. Note that not all variants in these three genes give rise to autosomal dominant Alzheimer's disease; some might be risk-modifiers or non-pathogenic. Further, evidence is accumulating that certain variants in the SORL1 gene are causative of Alzheimer's disease before the age of 70 years. The Alzheimer's disease-association of variants in the SORL1, ABCA7, and TREM2 genes was found in gene-based tests; carriers may come from small pedigrees with inheritance patterns of Alzheimer's disease suggestive of autosomal dominant inheritance. (B) GWAS hits are common (by convention, MAF >1%) variants that represent risk alleles that occur with significantly different frequency in patients with Alzheimer's disease and controls. Each variant is represented by the gene in which it occurs, or when the variant is non-coding, by the gene that maps closest to the variant (depicted in dark grey). (C) Protective variants are (very) rare variants suggested to confer resistance against age-associated or disease-associated risk factors of cognitive decline. GWAS=genome-wide association studies. MAF=minor allele frequency. OR=odds ratio. PRS=polygenic risk scores.

that it also mediates toxicity induced by tau pathology 79 or acts protectively against tau. 80

The contradictory effects of the microglia response partly reflect the limitations of mice models overexpressing tau for the study of Alzheimer's disease. It is possible that strong transgenic tau overexpression? induces an artificially strong neuroinflammatory response that is not seen in milder tau models. The use of mice models that do not overexpress tau, mouse-human chimeric mice, and one in-vitro models derived from human, induced pluripotent stem cells might help to explain the conflicting observations. Of note, all preclinical models are reductionistic in nature, implying that any conclusions towards therapeutic developments need to be made with caution.

Although cellular pathology has become central in the study of Alzheimer's disease, great progress has also been made in understanding the preceding biochemical phase of the disease (in ATN terms, before A positivity [presence of amyloid β]). Thanks to cryo-electron microscopy, amyloid β ⁸⁵ and tau fibrils are now known in finer detail. 66 Cryo-electron microscopy has also allowed full insight into how presenilins, the catalytic subunits of γ -secretases, interact with APP⁸⁷ and Notch substrates. 68 Complemented by functional studies on purified γ -secretase complexes, 71 it is now understood that clinical mutations in presenilins destabilise the γ -secretase—APP interactions, leading to premature release of longer, aggregation-prone amyloid β peptides. These insights support the development of new therapeutic approaches to tackle amyloid β in Alzheimer's disease.

The role of amyloid β in the disease cascade needs to be reintegrated with concepts of resilience and susceptibility. To this end, the cellular responses of neurons, astroglia, microglia, pericytes, and endothelial cells, which are largely defined by the genetic makeup of a patient, will

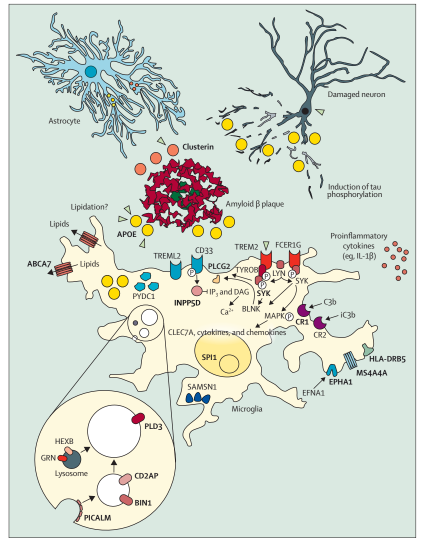


Figure 4: The cellular phase of Alzheimer's disease

Although amyloid plaques (red, middle of the figure) and tau phosphorylation and tangles (neurons, top right corner) are still considered the defining features of Alzheimer's disease, the focus of research has been widened from neurons to the response of other cell populations in the disease. The microglia-mediated inflammation, known for decades to be present in Alzheimer's disease, has finally taken centre-stage in functional research on the pathogenesis of the disease. Many of the risk-genes protein products (bold and capitals) identified in Alzheimer's disease (figure 3) are expressed and have functions in microglia. These genes become upregulated when microglia are exposed to amyloid plaques and many of the Alzheimer's disease risk genes are enriched in the disease-associated microglia response that characterises this cell state. The genes involved in this response and moderately positive in genome-wide association studies are indicated as well. Adapted from Sierksma et al., by permission of EMBO Molecular Medicine.

determine whether and how long a brain affected by amyloid pathology will continue to function normally. 58,76 Once homoeostasis collapses, Alzheimer's disease manifests itself clinically. Where and when tau influences this cellular phase is one of the most interesting questions for the field.

Apart from the core signature biochemical amyloid and tau pathology and the microglia response, which defines Alzheimer's disease, it is clear that vasculature, ⁸⁹ the blood–brain barrier, ⁸⁹ the glymphatic ⁶³ and other clearance

systems of the brain, 61 the peripheral immune system, 90 and potentially the gastrointestinal microbiome 91 affect the clinical development of the disease. Vascular pathology also affects blood–brain barrier integrity. 89,92 Leakage of the blood–brain barrier causes dementia independently from amyloid β and tau pathology, especially in $APOE\ \epsilon 4$ carriers. 92

Biomarkers

The biological definition of Alzheimer's disease is operationalised by the use of ATN biomarkers (appendix p 1).

Imaging biomarkers

Established markers: MRI, ¹⁸fluorodeoxyglucose (¹⁸FDG)-PET, and amyloid-PET

The three best validated neuroimaging biomarkers for Alzheimer's disease are medial temporal lobe atrophy on MRI and posterior cingulate and temporoparietal hypometabolism on $^{18}\text{FDG-PET}$ as measures of neurodegeneration, and cortical amyloid β deposition on amyloid-PET imaging. A five-phase strategic roadmap showed that the three biomarkers have almost achieved analytical and clinical validity (phases 1 to 3), although evidence for their clinical utility (phases 4 and 5) is considered insufficient. 93

Large prospective studies could provide answers regarding the clinical impact and utility of amyloid β imaging. The ABIDE study showed that amyloid β imaging improved diagnostic accuracy and confidence in a memory clinic setting with relatively young patients (under the age of 70 years). 94 The IDEAS study, carried out in individuals aged 65 years and older, showed that amyloid-PET imaging affected clinical diagnosis and diagnostic confidence in about 60% of patients with mild cognitive impairment or dementia. 95

Uncertainty regarding the order of tests hinders the widespread implementation of these imaging biomarkers. An interdisciplinary group of experts recently concluded that, although MRI is always recommended as the necessary first step after clinical evaluation, the decision on necessity and choice of the next biomarker test depends on the specific clinical profile and the individual diagnostic question. Amyloid-PET is most useful to rule out Alzheimer's disease, whereas 18FDG-PET has value for the differential diagnosis of neurodegenerative diseases, prediction of short-term clinical outcome, and staging of extent and localisation of neurodegenerative processes. Such algorithms can also be used to support clinicians in the choice of whether or not to do an additional diagnostic test.

Finally, consideration of regional (instead of global) cortical amyloid β deposition could allow detection of the earliest amyloid β stages (in temporobasal and frontomedial areas) with much higher sensitivity.⁷³

Ταυ-ΡΕΤ

Tau-PET ligands allow the in-vivo characterisation of tau tracer retention, consistent with Braak stages.⁷⁴ In

contrast to amyloid β deposition, tau-PET binding topography correlates with cognitive deficits, 97 is specific to the different Alzheimer's disease clinical phenotypes,75 and is predictive of subsequent rates of cognitive decline% and atrophy.99 Tau-PET is a powerful biomarker for differential diagnosis between Alzheimer's diseasetauopathy and other neurodegenerative tauopathies.100 Finally, longitudinal tau-PET studies highlight the sensitivity of this technique to track the progression of the disease, 101 and the spread of tau along brain networks, consistent with neuron-to-neuron propagation. 102 Tau-PET also helps to better understand the role of tau and its interaction with amyloid β. Preliminary data suggest that amyloid β might both accelerate tau accumulation¹⁰³ and allow the spread of tau outside of the medial temporal lobe.104

In May, 2020, the tau tracer flortaucipir was approved for clinical use by the US Food and Drug Administration. For tau-PET to enter clinical practice, methodological refinement is needed; for example, off-target and non-specific binding and analysis procedures are issues to be resolved. Descond-generation tracers that seem to have better signal-to-noise ratio, less off-target binding, and lower non-specific binding than first-generation tracers have been developed. Descond-generation tracers have been developed.

Other imaging modalities

Developments in PET ligands targeting SV2A imaging have opened new avenues to explore brain synaptic density. This progress is of particular interest in Alzheimer's disease, with preliminary reports of decreased SV2A binding in the hippocampus in patients with mild cognitive impairment or Alzheimer's disease. Further development of PET markers for neuroinflammation, α -synuclein, TDP43, and neurotransmitter systems are also eagerly awaited. Better use of multimodal neuroimaging is needed, including through the development of dual-phase amyloid-tau-PET imaging, hybrid PET-MR imaging, and artificial intelligence.

Fluid biomarkers

Amyloid β, phosphorylated tau, and neurodegeneration can also be ascertained via body fluid biomarkers (appendix p 1), greatly facilitated by the development of automated platforms for the analysis of amyloid β1–42, phosphorylated tau 181, and total tau. 109-111 Through extensive global collaboration (panel 2), reference methods and materials have been developed116 and assay outcomes between providers of cerebrospinal fluid biomarker assays for Alzheimer's disease have been aligned.72 Standardised operating procedures for cerebrospinal fluid collection and analysis 117,118 have also been developed, and a qualitycontrol programme for monitoring consistency in analysis of the results has been firmly established. 118,119 All these endeavors are directed at generating global, uniform cutoff points to define if a patient's profile is Alzheimer's disease-like.

Panel 2: Fluid biomarker consortia relevant to the field of Alzheimer's disease

Global Biomarker Standardization Consortium of the Alzheimer's Association 112

Aims to achieve consensus on the best ways to standardise and validate biomarker tests for use in global clinical practices.

Society for CSF Analysis and Clinical Neurochemistry¹¹³

Aims to exchange high-level international scientific experience, to facilitate the incorporation of cerebrospinal fluid diagnostics into clinical practice, and to give advice on the inclusion of cerebrospinal fluid analysis into clinical quidelines.

Foundation for the National Institutes of Health Biomarkers Consortium¹¹⁴

Aims to bring together the expertise and resources of various partners to rapidly identify, develop, and qualify potential high-impact biomarkers, particularly to enable improvements in drug development, clinical care, and regulatory decision making.

International Federation of Clinical Chemistry and Laboratory Medicine Working Group CSF Proteins¹¹⁵
Aims to develop certified reference material and reference methods for amyloid β42 or amyloid β40 and tau in

Cerebrospinal fluid markers

cerebrospinal fluid.

Aside from the established cerebrospinal fluid biomarkers amyloid β1-42, amyloid β1-40, phosphorylated tau 181, and total tau, some new developments can be reported. Markers reflecting axonal damage and synaptic dysfunction are relevant in light of synaptic pathology being present early in the disease course, and of its relation with functional outcomes and cognitive decline. Several of these biomarkers are emerging (eg, neurogranin, SNAP25, synaptotagmins, and the neuronal calcium sensing protein VLP1). 120-124 Of these, neurogranin seems the most promising, given its specificity for Alzheimer's disease and its increase in early stages. YKL40 (CHI3L1), a microglia and astrocyte biomarker and a promising marker to monitor treatment effect, is especially increased in frontotemporal dementia and (to a lesser extent) in Alzheimer's disease. 122,125 Soluble TREM2 is interesting because of its previously mentioned link to genetics. Increases in serum concentrations of TREM2 are observed at a group level independently of the presence of the mutation, and concentrations appear to have a bimodal course along the Alzheimer's disease spectrum.126

Some non-protein biomarkers are worth mentioning. Initial remarkable results on a plasma metabolomics profile¹²⁷ were replicated,¹²⁸⁻¹³⁰ although with different profiles. An important issue for the metabolome is the absence of specificity to a disease process and the subtlety of changes.

Serum and plasma biomarkers

Ultrasensitive technologies enable the accurate measurement of CNS proteins in blood. A poignant example is neurofilament light, a major axonal cytoskeleton protein that is a cross-disease biomarker of neurodegeneration.¹³¹ Levels of neurofilament light are increased in blood similarly as in cerebrospinal fluid, making clinical implementation of this marker feasible. In the dementia spectrum, neurofilament light has particular promise in the diagnosis of frontotemporal dementia,¹³² which makes it potentially useful in monitoring treatment response.¹³³⁻¹³⁵

Also encouraging are the consistent and converging reports showing that reductions in plasma amyloid β concentrations in Alzheimer's disease can be sensitively measured by immunoprecipitation combined with mass spectrometry, or microfluidics and other advanced technologies, such as Simoa, immunoreduction, and protein amide bond analysis. 136-142 Results of current collaborative investigations will show which technology provides the best sensitivity for different purposes (eg, screening, stratification, and effect monitoring) and holds the strongest promise for implementation in highthroughput analysis, which is needed when drugs become available and prescreening and monitoring of amyloid \(\beta \) changes becomes relevant. As for phosphorylated tau, three recent papers show strong evidence of plasma phosphorylated tau 181 and 217 as diagnostic biomarkers for Alzheimer's disease versus other dementias, and for identification of both amyloid β and phosphorylated tau pathology via PET.143-145

The exciting and rapid developments in plasma-based assays hold promise for prescreening in research (reducing the need for, and associated costs with, lumbar punctures and PET scans), and also, once properly validated, for diagnostic purposes in clinical practice. 146

For the World Wide FINGERS network see www.alz.org/ wwfingers

Treatment options

Non-pharmacological

Evidence for lifestyle changes

In 2019, WHO released the first guidelines for reduction of risk of cognitive decline and dementia. ¹⁴⁷ The guidelines acknowledge that, for some factors (eg, physical activity, diet, overweight or obesity, tobacco and alcohol use, hypertension, and diabetes), recommendations can be provided, although with different degrees of certainty. Some limitations in the current evidence include the scarcity of harmonisation (eg, exposure definition) and of long-term, randomised controlled trials, and little evidence from low-income and middle-income countries, where the prevalence of dementia is increasing rapidly.

The SPRINT-MIND trial reported that intensive blood pressure control (goal <120 mm Hg) is more effective in reducing the risk of cognitive impairment than standard blood pressure control (goal <140 mm Hg). 148 These results further highlight the idea that what is good for the heart is good for the brain, although the question of

the optimal therapeutic target remains, especially for individuals older than 70 years.

Multidomain interventions to prevent cognitive decline and dementia

Previous single-intervention failures stress the crucial need for a multimodal preventive approach that has been successful in the cardiovascular and diabetes prevention fields.149 The Finnish FINGER study was the first largescale, long-term, randomised controlled trial showing that a multidomain lifestyle-based intervention can reduce the risk of cognitive impairment among individuals at risk. 150,151 FINGER combined healthy balanced nutrition, physical exercise, cognitive training and social activities, and vascular and metabolic risk management. The trial showed benefits on cognition, even in people with genetic susceptibility to Alzheimer's disease. The French MAPT trial 1552 tested the association of a lifestyle intervention with omega-3 fatty acids supplements, and the Dutch PreDIVA trial¹⁵³ focused on the pharmacological management of vascular and metabolic risk factors. Both trials were negative for the primary outcomes, although subgroup analyses suggested cognitive benefits in subpopulations of participants with increased risk of dementia. In a substudy using amyloid-PET in the MAPT trial, 154 lifestyle intervention alone or in combination with omega-3 fatty acids was associated with improved primary cognitive outcome in people with positive amyloid β status. This finding highlights that even when lifestyle factors do not directly affect Alzheimer's disease pathology, they can still contribute to a positive outcome in individuals with Alzheimer's disease (appendix pp 2–4).

Future directions: from complexity to precision prevention

In 2020, more than 25 countries joined the World Wide FINGERS network, which aims to adapt, test, and optimise the FINGER model in different geographical, cultural, and economic settings. In many ongoing World Wide FINGERS trials, substudies are focusing on biomarkers (eg, the US-POINTER substudies with MRI, amyloid-PET and tau-PET), which will further clarify the role of various biomarkers and mechanisms among individuals at risk.¹⁵⁵ The Multimodal Prevention Trial for Alzheimer's Disease evaluates the feasibility of the FINGER multidomain lifestyle intervention in patients with prodromal Alzheimer's disease. This trial is an example of potential future studies in which pharmacological and non-pharmacological preventive strategies can be tested in combination. The study is testing the feasibility of a multidomain intervention, combined with a medical food product that showed promising results after 2 years of treatment in a large randomised controlled trial in patients with prodromal Alzheimer's disease,156 and sustained positive effects on clinical dementia rating and hippocampal volume after 3 years.¹⁵⁷ This type of study is necessary to identify the prevention potential on an individual basis, ultimately enabling a future of personalised medicine for Alzheimer's disease, in which multimodal interventions can be based on individually tailored combinations of lifestyle and drugs.

Pharmacological

Cognitive enhancing treatments for Alzheimer's disease Approved treatments that encompass the standard of care for many patients with Alzheimer's disease include cholinesterase inhibitors and the N-methyl-D-aspartate receptor antagonist memantine. No other symptomatic cognitive enhancing agent has been approved globally since the Seminar in 2016.¹ Three programmes assessing the utility of 5-HT₆ receptor antagonists for cognitive improvement have shown that this pathway is not a viable therapeutic approach for cognition.¹

Drugs to treat neuropsychiatric symptoms of Alzheimer's disease Progress is being made in developing psychotropic interventions specific for Alzheimer's disease or for dementia. Pimavanserin is a 5-HT_{2A} receptor inverse agonist that was assessed in a basket trial for dementia-related psychosis, which included patients with psychosis in the setting of Alzheimer's disease, Parkinson's disease with dementia, dementia with Lewy bodies, frontotemporal degeneration spectrum disorders, and vascular dementia.¹⁵⁹ The trial was stopped early for success, and pimavanserin will be submitted to the US Food and Drug Administration as a therapy for dementia-related psychosis.

Agitation is a common problem in dementia, occurring in up to 70% of patients with Alzheimer's disease in the course of their illness. ¹⁶⁰ Recent trials have been supportive of treatment with brexpiprazole (an atypical antipsychotic), citalopram (a selective serotonin reuptake inhibitor), and nabilone (a cannabinoid). ¹⁶¹ These studies suggest that appropriate interventions can reduce agitation. Ongoing trials are assessing the efficacy of brexpiprazole, escitalopram, prazosin, dextromethorphan plus quinidine, and dextromethorphan plus bupropion for agitation related to Alzheimer's disease.

Sleep and night-time behavioural disturbances disrupt the lives of patients and caregivers. A trial of suvorexant showed significant increases in total sleep time and decreased awakening after falling asleep. Suvorexant is a dual orexin antagonist approved for insomnia, and the authorised prescribing information now includes clinical trial and adverse event information regarding the use of the agent to treat insomnia in Alzheimer's disease. Lemborexant, another dual orexin antagonist, is in a trial for irregular sleep-wake rhythm disorder in patients with Alzheimer's disease.

Disease-modifying therapies for Alzheimer's disease

Most of the Alzheimer's disease drug-development pipeline is devoted to disease-modifying therapies (appendix pp 5–9). ^{64,163} These agents are in secondary prevention trials in individuals with preclinical, prodromal or mild, or moderate-to-severe Alzheimer's disease.

Amyloid β is the most common target of drug development programmes in phase 2 and phase 3. Growing evidence suggests that by removing amyloid β oligomers (soluble aggregates of amyloid β) and plaques (insoluble extracellular aggregates of fibrillar amyloid β) with monoclonal antibodies, disease progression can be slowed. 164 Aducanumab, BAN2401, and gantenerumab all reduce amyloid β plaques. 165 These agents also reduce phosphorylated tau, neurogranin, and neurofilament light in the cerebrospinal fluid; observations that suggest that removal of amyloid β is associated with downstream effects on tau pathology and neurodegeneration. In each case, ambiguities in the clinical trials remain to be resolved. No therapeutic agents have yet been approved by regulatory authorities, and phase 3 clinical trials (NCT04241068, NCT03443973, NCT04339413, NCT04592341, NCT03444870, NCT03887455, and NCT04468659) are ongoing. A recent phase 2 trial of donanemab suggests that this antibody directed against the pyroglutamate-modified form of amyloid β (an oligomer of amyloid β pE3 and amyloid β 42) has promise as an amyloid-targeted treatment. 166 New phase 3 trials of donanemab (NCT04437511 and NCT04640077) have since been initiated.

Amyloid β vaccines are being tested in active immunotherapy trials and are a promising area for Alzheimer's disease therapeutics. BACE1 and BACE2 inhibitors were a promising class of Alzheimer's disease-modifying therapies that markedly reduce concentrations of cerebrospinal fluid amyloid β . Several of these trials have been stopped because of an acceleration of deterioration in cognition, elevated liver enzymes, or futility. ¹⁶⁷ Because many trials were stopped early on, it remains unclear whether longer treatments would have exerted beneficial effects. Further development of this class of agents is unlikely, unless major new insights into their safety and efficacy are achieved.

Tau biology is providing another repertoire of potentially important targets for disease-modifying therapies. Les Several monoclonal antibodies targeting different epitopes are in trials. The monoclonal antibodies are intended to engage extracellular tau as it spreads from cell to cell. Small molecules targeting tau aggregation and neurofibrillary tangle formation are being assessed. All these approaches come with potential side-effects and experts in the field should seriously think about risks to benefits and more complex trials, with better dose finding and measurements of therapeutic target engagement. Otherwise, it is probable that tau-targeted trials will end in premature futility analyses, with little additional learning as to why these trials are negative and what can be improved.

Neuroinflammation is recognised as a major component of the pathology of Alzheimer's disease, contributing to disease progression and neurodegeneration. Oligomannate was approved in China in 2019, after a phase 3 trial conducted in the same country showed cognitive improvement. ¹⁶⁹ This agent is hypothesised to be efficacious, on

the basis of non-clinical observations, in reducing brain inflammation in patients with Alzheimer's disease through its effect on the gut microbiome, reducing dysbiosis, restoring normal gut bacterial composition, and decreasing peripheral inflammatory cell populations, which can contribute to central inflammation. A global trial (NCT04520412) is planned to determine the extent to which these effects can be reproduced in other populations.

Various other mechanisms are being targeted in Alzheimer's disease drug development programmes (appendix pp 9–12). Infections are also hypothesised to contribute to Alzheimer's disease onset or progression, and agents that target bacteria or viruses are in clinical trials for Alzheimer's disease. Neuroprotection is essential for successful disease-modification, and some agents target neuroprotection directly through growth factors, mitochondrial function, or other mechanisms, in an effort to slow disease progression.

The Dominantly Inherited Alzheimer Network Treatment Unit is an adaptive prevention trial platform, assessing multiple agents simultaneously in individuals with autosomal dominant Alzheimer's disease. To A recent report presented at the 2020 Alzheimer's and Parkinson's Diseases Conference showed that, in a small sample of mutation carriers, neither solanezumab nor gantenerumab affected clinical outcomes compared with placebo. To Gantenerumab, but not solanezumab, positively affected biomarker outcomes.

An overview of the Alzheimer's disease-modifying treatments pipeline shows that several agents have clinical or biomarker benefits, and confirmatory trials are being pursued. Some agents have been submitted to the US Food and Drug Administration or the European Medicines Agency for regulatory review. The development of improved trial designs, a larger repertoire of biomarkers reporting on a wider variety of cell processes, improved outcome measures, and better analytical approaches, along with improving insight into the biology of Alzheimer's disease, support the optimism in the field that the emergence of important new therapies for Alzheimer's disease might be imminent.

Conclusions

In the past 5 years, substantial progress has been made into understanding the pathophysiology and genetic basis of Alzheimer's disease. The amyloid β cascade hypothesis has been modified by a more thorough understanding of the cellular, preclinical, phase of Alzheimer's disease. Genetic studies have moved from pinpointing three causal genes and one risk gene to identifying a plethora of genes that can be put into a polygenic risk score for Alzheimer's disease. The developments in biomarker diagnosis have led to a complete rethinking of how to label Alzheimer's disease outside of and before clinical symptomatology, enabling the enrolment of patients in research in a much earlier phase of the disease, particularly now that blood biomarkers seem to be within

reach. Further refinement of the diagnostic classification and pathological underpinnings of the disease will be made by molecular imaging, allowing visualisation of copathology and regional protein aggregation. Following these developments will be insights in risk reduction, primary and secondary prevention, non-pharmacological and pharmacological approaches, ultimately given in parallel and at a much earlier timepoint than has been trialled before. If the field keeps up this pace, very early identification and multimodal treatment of patients can become a reality.

Contributors

PS devised the paper and invited the other authors to provide parts of the text according to their expertise. PS and WMVdF collated and edited the contributions and did additional literature searches. All authors contributed parts of the paper, reviewed the full paper, and gave final approval for submission.

Declaration of interests

PS reports research support from Alzheimer Nederland, Dioraphte, Stichting VUmc Fonds, and Stichting Alzheimer & Neuropsychiatrie; is a consultant for EIP Pharma, Vivoryon, Toyama Fuji Film, AC Immune, Axon Neuroscience, GemVax, Medavante, Novartis Cardiology, and Green Valley; is co-editor-in-chief of Alzheimer's Research & Therapy; and is a part-time managing partner of the Life Science Partners Dementia Fund since Oct 1, 2020 (after acceptance of this Seminar). BDS reports research support from Janssen Pharmaceutica, European Research Council, various academic research sponsors, Methusalem Grant of the Flemish Government, Medical Research Council, Alzheimer Association (USA), Alzheimer Research UK, and Alzheimer Society (UK), BDS is a consultant for Eisai Pharmaceuticals and holds several patents (appendix p 13). MK reports research support from Academy of Finland, Swedish Research Council, Joint Program of Neurodegenerative Disorders, IMI, Knut and Alice Wallenberg Foundation, Center for Innovative Medicine, Stiftelsen Stockholms Sjukhem, Konung Gustaf V:s och Drottning Victorias Frimurarestiftelse, Alzheimer's Research and Prevention Foundation, Alzheimerfonden, Hjärnfonden, Region Stockholm, and Leif Lundblad Foundation grants. MK is part of a guidelines development group in WHO, a Governance Committee member of the Global Council on Brain Health, and is on the advisory board for Combinostics, Roche, and Biogen. HH reports research support from ZonMw, NWO, EU-JPND, Alzheimer Nederland, and Stichting Dioraphte. GC reports research support from European Union Horizon 2020 programme, INSERM, Fondation d'entreprise MMA des Entrepreneurs du Futur, Fondation Alzheimer, Programme Hospitalier de Recherche Clinique, Région Normandie, Association France Alzheimer et maladies apparentées, and Fondation Vaincre Alzheimer, CET reports research support from European Commission (Marie Curie International Training Network, JPND), Health Holland, the Dutch Research Council (ZonMw), The Weston Brain Institute, and Alzheimer Nederland; and has the following industrial collaborations: collaboration contract with ADx Neurosciences, contract research or received grants from Probiodrug, AC Immune, Biogen-Esai, CogRx, Toyama, Janssen Prevention Center, Boehringer, AxonNeurosciences, Fujirebio, EIP farma, PeopleBio, and Roche, and is on the advisory board of Roche. JC has provided consultation to Acadia, Actinogen, AgeneBio, Alkahest, Alzheon, Annovis, Avanir, Axsome, Biogen, Cassava, Cerecin, Cerevel, Cognoptix, Cortexyme, EIP Pharma, Eisai, Foresight, GemVax, Green Valley, Grifols, Hisun, Karuna, Nutricia, Orion, Otsuka, ReMYND, Resverlogix, Roche, Samumed, Samus Therapeutics, Third Rock, Signant Health, Sunovion, Suven, and United Neuroscience pharmaceutical and assessment companies. JC receives support from Keep Memory Alive, COBRE grant #P20GM109025, TRC-PAD #R01AG053798, and DIAGNOSE CTE #U01NS093334. JC owns the copyright of the Neuropsychiatric Inventory. WMVdF reports research support from ZonMw, NWO, EU-FP7, EU-JPND, Alzheimer Nederland, CardioVascular Onderzoek Nederland, Health~Holland, Topsector Life Sciences & Health, Stichting Dioraphte, Gieskes-Strijbis Fonds, Stichting Equilibrio, Pasman Stichting, Fujifilm, Boehringer Ingelheim, Life-MI, AVID, Janssen Stellar, Combinostics.

WMVdF is chair of Pasman Stichting. WMVdF has performed contract research for Biogen and received speaker fees from Biogen and Roche.

Acknowledgments

Research at the Alzheimer Center Amsterdam is part of the neurodegeneration research programme of Amsterdam Neuroscience. The Alzheimer Centre Amsterdam is supported by Stichting Alzheimer Nederland and Stichting VUmc Fonds. The chair of Wiesje van der Flier is supported by the Pasman Stichting. MK and PS are recipients of the JPND-funded Euro-FINGERS project (#ZonMW 733051102).

References

- 1 Alzheimer Europe. Dementia in Europe Yearbook 2019: estimating the prevalence of dementia in Europe. 2020. https://www.alzheimereurope.org/content/download/195515/1457520/file/FINAL%20 05707%20Alzheimer%20Europe%20yearbook%202019.pdf (accessed Jan 24, 2021).
- 2 Scheltens P, Blennow K, Breteler MM, et al. Alzheimer's disease. Lancet 2016; 388: 505–17.
- 3 Dubois B, Feldman HH, Jacova C, et al. Research criteria for the diagnosis of Alzheimer's disease: revising the NINCDS-ADRDA criteria. *Lancet Neurol* 2007; 6: 734–46.
- Dubois B, Feldman HH, Jacova C, et al. Revising the definition of Alzheimer's disease: a new lexicon. *Lancet Neurol* 2010; 9: 1118–27.
- McKhann G, Drachman D, Folstein M, Katzman R, Price D, Stadlan EM. Clinical diagnosis of Alzheimer's disease: report of the NINCDS-ADRDA Work Group under the auspices of Department of Health and Human Services Task Force on Alzheimer's Disease. Neurology 1984; 34: 939–44.
- 6 McKhann GM, Knopman DS, Chertkow H, et al. The diagnosis of dementia due to Alzheimer's disease: recommendations from the National Institute on Aging-Alzheimer's Association workgroups on diagnostic guidelines for Alzheimer's disease. Alzheimers Dement 2011; 7: 263–69.
- 7 Knopman DS, Petersen RC, Jack CR Jr. A brief history of "Alzheimer disease": multiple meanings separated by a common name. Neurology 2019; 92: 1053–59.
- 8 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: 535–62.
- 9 Sweeney MD, Montagne A, Sagare AP, et al. Vascular dysfunction—the disregarded partner of Alzheimer's disease. Alzheimers Dement 2019; 15: 158–67.
- 10 Jack CR Jr, Holtzman DM, Sperling R. Dementia is not synonymous with Alzheimer's disease. Sci Transl Med 2019; 11: eaav0511.
- Altomare D, de Wilde A, Ossenkoppele R, et al. Applying the ATN scheme in a memory clinic population: the ABIDE project. Neurology 2019; 93: e1635–46.
- 12 Cummings J, Lee G, Ritter A, Sabbagh M, Zhong K. Alzheimer's disease drug development pipeline: 2019. Alzheimers Dement (NY) 2019; 5: 272–93.
- 13 van Maurik IS, Vos SJ, Bos I, et al. Biomarker-based prognosis for people with mild cognitive impairment (ABIDE): a modelling study. *Lancet Neurol* 2019; 18: 1034–44.
- 14 Visser LNC, van Maurik IS, Bouwman FH, et al. Clinicians' communication with patients receiving a mild cognitive impairment diagnosis: the ABIDE project. PLoS One 2020; 15: e0227282.
- 15 Jessen F, Amariglio RE, Buckley RF, et al. The characterisation of subjective cognitive decline. *Lancet Neurol* 2020; 19: 271–78.
- van Maurik IS, Slot RER, Verfaillie SCJ, et al. Personalized risk for clinical progression in cognitively normal subjects—the ABIDE project. Alzheimers Res Ther 2019; 11: 33.
- 17 Ebenau JL, Timmers T, Wesselman LMP, et al. ATN classification and clinical progression in subjective cognitive decline: The SCIENCe project. *Neurology* 2020; 95: e46–58.
- 18 Fruijtier AD, Visser LNC, van Maurik IS, et al. ABIDE Delphi study: topics to discuss in diagnostic consultations in memory clinics. Alzheimers Res Ther 2019; 11: 77.
- 19 van Maurik IS, Visser LN, Pel-Littel RE, et al. Development and usability of ADappt: web-based tool to support clinicians, patients, and caregivers in the diagnosis of mild cognitive impairment and Alzheimer disease. JMIR Form Res 2019; 3: e13417.

- 20 Alzheimer's Disease International. World Alzheimer Report 2018. The state of the art of dementia research: new frontiers. September, 2018. https://www.alzint.org/u/ WorldAlzheimerReport2018.pdf (accessed Sept 9, 2020).
- 21 Wu YT, Beiser AS, Breteler MMB, et al. The changing prevalence and incidence of dementia over time—current evidence. Nat Rev Neurol 2017; 13: 327–39.
- 22 Prince M, Ali GC, Guerchet M, Prina AM, Albanese E, Wu YT. Recent global trends in the prevalence and incidence of dementia, and survival with dementia. Alzheimers Res Ther 2016; 8: 23.
- 23 Mayeda ER, Glymour MM, Quesenberry CP, Johnson JK, Perez-Stable EJ, Whitmer RA. Survival after dementia diagnosis in five racial/ethnic groups. Alzheimers Dement 2017; 13: 761–69.
- 24 Rhodius-Meester HFM, Tijms BM, Lemstra AW, et al. Survival in memory clinic cohort is short, even in young-onset dementia. J Neurol Neurosurg Psychiatry 2019; 90: 726–28.
- 25 Vermunt L, Sikkes SAM, Hout AVD, et al. Duration of preclinical, prodromal, and dementia stages of Alzheimer's disease in relation to age, sex, and APOE genotype. Alzheimer Dementia 2019; 15: 888–98.
- 26 Jack CR Jr, Therneau TM, Weigand SD, et al. Prevalence of biologically vs clinically defined Alzheimer spectrum entities using the National Institute on Aging-Alzheimer's Association research framework. JAMA Neurol 2019; 76: 1174–83.
- 27 van der Lee SJ, Wolters FJ, Ikram MK, et al. The effect of APOE and other common genetic variants on the onset of Alzheimer's disease and dementia: a community-based cohort study. *Lancet Neurology* 2018: 17: 434–44.
- 28 Buckley RF, Mormino EC, Rabin JS, et al. Sex differences in the association of global amyloid and regional tau deposition measured by positron emission tomography in clinically normal older adults. JAMA Neurol 2019; 76: 542–51.
- 29 Mofrad RB, Tijms BM, Scheltens P, et al. Sex differences in CSF biomarkers vary by Alzheimer's disease stage and APOE ε4 genotype. Neurology 2020; 95: e2378–88.
- 30 Livingston G, Huntley J, Sommerlad A, et al. Dementia prevention, intervention, and care: 2020 report of the *Lancet* Commission. *Lancet* 2020; 396: 413–46.
- 31 Lane CA, Barnes J, Nicholas JM, et al. Associations between vascular risk across adulthood and brain pathology in late life: evidence from a British birth cohort. JAMA Neurol 2019: 77: 175–83.
- 32 Gottesman RF, Mosley TH, Knopman DS, et al. Association of intracranial atherosclerotic disease with brain beta-amyloid deposition: secondary analysis of the ARIC study. JAMA Neurol 2019; 77: 350–57.
- 33 Gustavsson AM, van Westen D, Stomrud E, Engstrom G, Nagga K, Hansson O. Midlife atherosclerosis and development of Alzheimer or vascular dementia. Ann Neurol 2020; 87: 52–62.
- 34 Gatz M, Reynolds CA, Fratiglioni L, et al. Role of genes and environments for explaining Alzheimer disease. Arch Gen Psychiatry 2006: 63: 168–74.
- 35 Bellenguez C, Charbonnier C, Grenier-Boley B, et al. Contribution to Alzheimer's disease risk of rare variants in TREM2, SORL1, and ABCA7 in 1779 cases and 1273 controls. Neurobiol Aging 2017; 59: e1–9
- 36 Paudel HK, Ridge PG, Mukherjee S, Crane PK, Kauwe JSK. Alzheimer's disease: analyzing the missing heritability. PLoS One 2013; 8: e79771.
- 37 Jansen IE, Savage JE, Watanabe K, et al. Genome-wide metaanalysis identifies new loci and functional pathways influencing Alzheimer's disease risk. Nat Genet 2019; 51: 404–13.
- 38 Escott-Price V, Sims R, Bannister C, et al. Common polygenic variation enhances risk prediction for Alzheimer's disease. *Brain* 2015; 138: 3673–84.
- 39 Escott-Price V, Myers AJ, Huentelman M, Hardy J. Polygenic risk score analysis of pathologically confirmed Alzheimer disease. Ann Neurol 2017: 82: 311–14.
- 40 Di Marco LY, Venneri A, Farkas E, Evans PC, Marzo A, Frangi AF. Vascular dysfunction in the pathogenesis of Alzheimer's disease—a review of endothelium-mediated mechanisms and ensuing vicious circles. *Neurobiol Dis* 2015; 82: 593–606.
- 41 Bennett RE, Robbins AB, Hu M, et al. Tau induces blood vessel abnormalities and angiogenesis-related gene expression in P301L transgenic mice and human Alzheimer's disease. Proc Natl Acad Sci USA 2018; 115: E1289–98.

- 42 Verheijen J, Sleegers K. Understanding Alzheimer disease at the interface between genetics and transcriptomics. *Trends Genet* 2018; 34: 434–47
- 43 Naj AC, Schellenberg GD. Genomic variants, genes, and pathways of Alzheimer's disease: an overview. Am J Med Genet B Neuropsychiatr Genet 2017; 174: 5–26.
- 44 Hardy J, Bogdanovic N, Winblad B, et al. Pathways to Alzheimer's disease. J Intern Med 2014; 275: 296–303.
- 45 Van Cauwenberghe C, Van Broeckhoven C, Sleegers K. The genetic landscape of Alzheimer disease: clinical implications and perspectives. Genet Med 2016; 18: 421–30.
- 46 Holstege H, van der Lee SJ, Hulsman M, et al. Characterization of pathogenic SORL1 genetic variants for association with Alzheimer's disease: a clinical interpretation strategy. Eur J Hum Genet 2017; 25: 973–81
- 47 Cuyvers E, De Roeck A, Van den Bossche T, et al. Mutations in ABCA7 in a Belgian cohort of Alzheimer's disease patients: a targeted resequencing study. *Lancet Neurol* 2015; 14: 814–22.
- 48 Jonsson T, Stefansson H, Steinberg S, et al. Variant of TREM2 associated with the risk of Alzheimer's disease. N Engl J Med 2013; 368: 107–16.
- 49 Guerreiro R, Wojtas A, Bras J, et al. TREM2 variants in Alzheimer's disease. N Engl J Med 2013; 368: 117–27.
- 50 Genin E, Hannequin D, Wallon D, et al. APOE and Alzheimer disease: a major gene with semi-dominant inheritance. Mol Psychiatry 2011; 16: 903–07.
- 51 Reiman EM, Arboleda-Velasquez JF, Quiroz YT, et al. Exceptionally low likelihood of Alzheimer's dementia in APOE2 homozygotes from a 5,000-person neuropathological study. Nat Commun 2020; 11: 667.
- 52 Jonsson T, Atwal JK, Steinberg S, et al. A mutation in APP protects against Alzheimer's disease and age-related cognitive decline. Nature 2012: 488: 96–99
- 53 Sims R, van der Lee SJ, Naj AC, et al. Rare coding variants in PLCG2, ABI3, and TREM2 implicate microglial-mediated innate immunity in Alzheimer's disease. Nat Genets 2017; 49: 1373–84.
- 54 van der Lee SJ, Conway OJ, Jansen I, et al. A nonsynonymous mutation in *PLCG2* reduces the risk of Alzheimer's disease, dementia with Lewy bodies and frontotemporal dementia, and increases the likelihood of longevity. *Acta Neuropathol* 2019; 138: 237–50
- 55 Beker N, Sikkes SAM, Hulsman M, et al. Longitudinal maintenance of cognitive health in centenarians in the 100-plus study. JAMA Netw Open 2020; 3: e200094.
- 56 Arboleda-Velasquez JF, Lopera F, O'Hare M, et al. Resistance to autosomal dominant Alzheimer's disease in an APOE3 Christchurch homozygote: a case report. Nat Med 2019; 25: 1680–83.
- 57 Belloy ME, Napolioni V, Han SS, Le Guen Y, Greicius MD, Alzheimer's Disease Neuroimaging Initiative. Association of klothovs heterozygosity with risk of Alzheimer disease in individuals who carry APOE4. JAMA Neurol 2020; 77: 849–62.
- 58 De Strooper B, Karran E. The cellular phase of Alzheimer's disease. Cell 2016; 164: 603–15.
- 59 Venegas C, Kumar S, Franklin BS, et al. Microglia-derived ASC specks cross-seed amyloid-beta in Alzheimer's disease. *Nature* 2017; 552: 355–61.
- 60 Sweeney MD, Sagare AP, Zlokovic BV. Blood-brain barrier breakdown in Alzheimer disease and other neurodegenerative disorders. Nat Rev Neurol 2018; 14: 133–50.
- 61 Da Mesquita S, Louveau A, Vaccari A, et al. Functional aspects of meningeal lymphatics in ageing and Alzheimer's disease. *Nature* 2018: 560: 185–91.
- 62 Lu T, Aron L, Zullo J, et al. REST and stress resistance in ageing and Alzheimer's disease. *Nature* 2014; 507: 448–54.
- 63 Plog BA, Nedergaard M. The glymphatic system in central nervous system health and disease: past, present, and future. Annu Rev Pathol 2018; 13: 379–94.
- 64 Long JM, Holtzman DM. Alzheimer disease: an update on pathobiology and treatment strategies. Cell 2019; 179: 312–39.
- 65 Koper MJ, Van Schoor E, Ospitalieri S, et al. Necrosome complex detected in granulovacuolar degeneration is associated with neuronal loss in Alzheimer's disease. Acta Neuropathol 2020; 139: 463–84.

- 66 Keren-Shaul H, Spinrad A, Weiner A, et al. A unique microglia type associated with restricting development of Alzheimer's disease. Cell 2017: 169: 1276–90.
- Farhizkar S, Arzberger T, Brendel M, et al. Loss of TREM2 function increases amyloid seeding but reduces plaque-associated ApoE. Nat Neurosci 2019; 22: 191–204.
- 68 Sala Frigerio C, Wolfs L, Fattorelli N, et al. The major risk factors for Alzheimer's disease: age, sex, and genes modulate the microglia response to Aβ plaques. Cell Rep 2019; 27: 1293–306.
- 69 Hansen DV, Hanson JE, Sheng M. Microglia in Alzheimer's disease. J Cell Biol 2017; 217: 459–72.
- 70 Yeh FL, Wang Y, Tom I, Gonzalez LC, Sheng M. TREM2 binds to apolipoproteins, including APOE and CLU/APOJ, and thereby facilitates uptake of amyloid-beta by microglia. *Neuron* 2016; 91: 328–40.
- 71 Szaruga M, Munteanu B, Lismont S, et al. Alzheimer's-causing mutations shift Aβ length by destabilizing γ-secretase-Aβn interactions. *Cell* 2017; 170: 443–56.
- 72 Kuhlmann J, Andreasson U, Pannee J, et al. CSF Abeta1–42 an excellent but complicated Alzheimer's biomarker—a route to standardisation. Clin Chim Acta 2017; 467: 27–33.
- 73 Grothe MJ, Barthel H, Sepulcre J, et al. In vivo staging of regional amyloid deposition. *Neurology* 2017; 89: 2031–38.
- 74 Cho H, Choi JY, Hwang MS, et al. In vivo cortical spreading pattern of tau and amyloid in the Alzheimer disease spectrum. Ann Neurol 2016; 80: 247–58.
- 75 Ossenkoppele R, Schonhaut DR, Baker SL, et al. Tau, amyloid, and hypometabolism in a patient with posterior cortical atrophy. *Ann Neurol* 2015; 77: 338–42.
- 76 Sierksma A, Lu A, Mancuso R, et al. Novel Alzheimer risk genes determine the microglia response to amyloid-beta but not to TAU pathology. EMBO Mol Med 2020; 12: e10606.
- 77 Griciuc A, Patel S, Federico AN, et al. TREM2 acts downstream of CD33 in modulating microglial pathology in Alzheimer's disease. *Neuron* 2019; 103: 820–35.
- 78 Deming Y, Filipello F, Cignarella F, et al. The MS4A gene cluster is a key modulator of soluble TREM2 and Alzheimer's disease risk. Sci Transl Med 2019: 11: eaau2291.
- 79 Leyns CEG, Ulrich JD, Finn MB, et al. TREM2 deficiency attenuates neuroinflammation and protects against neurodegeneration in a mouse model of tauopathy. *Proc Natl Acad Sci USA* 2017; 114: 11524–29.
- Leyns CEG, Gratuze M, Narasimhan S, et al. TREM2 function impedes tau seeding in neuritic plaques. *Nat Neurosci* 2019; 22: 1217–22.
- 81 Sasaguri H, Nilsson P, Hashimoto S, et al. APP mouse models for Alzheimer's disease preclinical studies. EMBO J 2017; 36: 2473–87.
- 82 Mancuso R, Van Den Daele J, Fattorelli N, et al. Stem-cell-derived human microglia transplanted in mouse brain to study human disease. Nat Neurosci 2019; 22: 2111–16.
- 83 Hasselmann J, Coburn MA, England W, et al. Development of a chimeric model to study and manipulate human microglia in vivo. *Neuron* 2019; 103: 1016–33.
- 84 Park J, Wetzel I, Marriott I, et al. A 3D human triculture system modeling neurodegeneration and neuroinflammation in Alzheimer's disease. Nat Neurosci 2018; 21: 941–51.
- 85 Gremer L, Schölzel D, Schenk C, et al. Fibril structure of amyloid-beta (1–42) by cryo-electron microscopy. Science 2017; 358: 116–19.
- 86 Fitzpatrick AWP, Falcon B, He S, et al. Cryo-EM structures of tau filaments from Alzheimer's disease. *Nature* 2017; 547: 185–90.
- 87 Zhou R, Yang G, Guo X, Zhou Q, Lei J, Shi Y. Recognition of the amyloid precursor protein by human g-secretase. *Science* 2019; 363: eaaw0930.
- 88 Yang G, Zhou R, Zhou Q, et al. Structural basis of Notch recognition by human γ-secretase. Nature 2019; 565: 192–97.
- 89 Nation DA, Sweeney MD, Montagne A, et al. Blood-brain barrier breakdown is an early biomarker of human cognitive dysfunction. *Nat Med* 2019; 25: 270–76.
- 90 Gate D, Saligrama N, Leventhal O, et al. Clonally expanded CD8 T cells patrol the cerebrospinal fluid in Alzheimer's disease. Nature 2020; 577: 399–404.

- 91 Dodiya HB, Frith M, Sidebottom A, et al. Synergistic depletion of gut microbial consortia, but not individual antibiotics, reduces amyloidosis in APPPS1–21 Alzheimer's transgenic mice. Sci Rep 2020; 10: 8183.
- 92 Montagne A, Nation DA, Sagare AP, et al. APOE4 leads to bloodbrain barrier dysfunction predicting cognitive decline. Nature 2020; 581: 71–76.
- 93 Frisoni GB, Boccardi M, Barkhof F, et al. Strategic roadmap for an early diagnosis of Alzheimer's disease based on biomarkers. *Lancet Neurol* 2017; 16: 661–76.
- 94 de Wilde A, van der Flier WM, Pelkmans W, et al. Association of amyloid positron emission tomography with changes in diagnosis and patient treatment in an unselected memory clinic cohort: the ABIDE project. JAMA Neurol 2018; 75: 1062–70.
- 95 Rabinovici GD, Gatsonis C, Apgar C, et al. Association of amyloid positron emission tomography with subsequent change in clinical management among medicare beneficiaries with mild cognitive impairment or dementia. JAMA 2019; 321: 1286–94.
- 96 Rhodius-Meester HFM, van Maurik IS, Koikkalainen J, et al. Selection of memory clinic patients for CSF biomarker assessment can be restricted to a quarter of cases by using computerized decision support, without compromising diagnostic accuracy. PLoS One 2020; 15: e0226784.
- 97 Aschenbrenner AJ, Gordon BA, Benzinger TLS, Morris JC, Hassenstab JJ. Influence of tau PET, amyloid PET, and hippocampal volume on cognition in Alzheimer disease. *Neurology* 2018; 91: e859–66.
- 98 Brier MR, Gordon B, Friedrichsen K, et al. Tau and Aβ imaging, CSF measures, and cognition in Alzheimer's disease. Sci Transl Med 2016: 8: 338m66
- 99 La Joie R, Visani AV, Baker SL, et al. Prospective longitudinal atrophy in Alzheimer's disease correlates with the intensity and topography of baseline tau-PET. Sci Transl Med 2020; 12: eaau5732.
- 100 Ossenkoppele R, Rabinovici GD, Smith R, et al. Discriminative accuracy of [18F]flortaucipir positron emission tomography for Alzheimer disease vs other neurodegenerative disorders. JAMA 2018: 320: 1151–62.
- 101 Harrison TM, La Joie R, Maass A, et al. Longitudinal tau accumulation and atrophy in aging and Alzheimer disease. Ann Neurol 2019; 85: 229–40.
- 102 Franzmeier N, Neitzel J, Rubinski A, et al. Functional brain architecture is associated with the rate of tau accumulation in Alzheimer's disease. Nat Commun 2020; 11: 347.
- 103 Jack CR Jr, Wiste HJ, Schwarz CG, et al. Longitudinal tau PET in ageing and Alzheimer's disease. Brain 2018; 141: 1517–28.
- 104 Wang L, Benzinger TL, Su Y, et al. Evaluation of tau imaging in staging Alzheimer disease and revealing interactions between beta-amyloid and tauopathy. JAMA Neurol 2016; 73: 1070–77.
- 105 Villemagne VL, Dore V, Burnham SC, Masters CL, Rowe CC. Imaging tau and amyloid-beta proteinopathies in Alzheimer disease and other conditions. Nat Rev Neurol 2018; 14: 225–36.
- 106 Leuzy A, Chiotis K, Lemoine L, et al. Tau PET imaging in neurodegenerative tauopathies-still a challenge. Mol Psychiatry 2019; 24: 1112–34.
- 107 Cai Z, Li S, Matuskey D, Nabulsi N, Huang Y. PET imaging of synaptic density: a new tool for investigation of neuropsychiatric diseases. *Neurosci Lett* 2019; 691: 44–50.
- 108 Chen MK, Mecca AP, Naganawa M, et al. Assessing synaptic density in Alzheimer disease with synaptic vesicle glycoprotein 2a positron emission tomographic imaging. [AMA Neurol 2018; 75: 1215–24.
- 109 Blennow K, Shaw LM, Stomrud E, et al. Predicting clinical decline and conversion to Alzheimer's disease or dementia using novel Elecsys Aβ(1–42), pTau and tTau CSF immunoassays. Sci Rep 2019; 9: 19024.
- 110 Hansson O, Seibyl J, Stomrud E, et al. CSF biomarkers of Alzheimer's disease concord with amyloid-beta PET and predict clinical progression: a study of fully automated immunoassays in BioFINDER and ADNI cohorts. Alzheimers Dement 2018; 14: 1470–81.
- 111 Kaplow J, Vandijck M, Gray J, et al. Concordance of Lumipulse cerebrospinal fluid t-tau/Abeta42 ratio with amyloid PET status. Alzheimers Dement 2020; 16: 144–52.
- 112 Global Biomarker Standardization Consortium. https://www.alz. org/research/for_researchers/partnerships/gbsc (accessed Sept 9, 2020).

- 113 Society for CSF Analysis and Clinical Neurochemistry. https://h001. ssl-redirect.de/www.neurochem.info (accessed Sept 1, 2020).
- 114 FNIH Biomarkers Consortium. https://fnih.org/what-we-do/biomarkers-consortium (accessed Sept 1, 2020).
- 115 International Federation of Clinical Chemistry and Laboratory Medicine Working Group CSF Proteins. https://www.ifcc.org/ifcc-scientific-division/sd-working-groups/csf-proteins-wg-csf (accessed Sept 1, 2020).
- 116 Pannee J, Gobom J, Shaw LM, et al. Round robin test on quantification of amyloid-beta 1–42 in cerebrospinal fluid by mass spectrometry. Alzheimers Dement 2016; 12: 55–59.
- 117 Engelborghs S, Niemantsverdriet E, Struyfs H, et al. Consensus guidelines for lumbar puncture in patients with neurological diseases. Alzheimers Dement (Amst) 2017; 8: 111–26.
- 118 Hok AHYS, Willemse EAJ, Teunissen CE, Del Campo M. Guidelines for CSF processing and biobanking: impact on the identification and development of optimal CSF protein biomarkers. Methods Mol Biol 2019; 2044: 27–50.
- 119 Mattsson N, Andreasson U, Persson S, et al. CSF biomarker variability in the Alzheimer's Association quality control program. Alzheimers Dement 2013; 9: 251–61.
- 120 Galasko D, Xiao M, Xu D, et al. Synaptic biomarkers in CSF aid in diagnosis, correlate with cognition and predict progression in mild cognitive impairment and Alzheimer's disease. Alzheimers Dement (N Y) 2019; 5: 871–82.
- 121 Hoglund K, Schussler N, Kvartsberg H, et al. Cerebrospinal fluid neurogranin in an inducible mouse model of neurodegeneration: a translatable marker of synaptic degeneration. *Neurobiol Dis* 2020; 134: 104645.
- 122 Janelidze S, Hertze J, Zetterberg H, et al. Cerebrospinal fluid neurogranin and YKL-40 as biomarkers of Alzheimer's disease. *Ann Clin Transl Neurol* 2016; **3**: 12–20.
- 123 Duits FH, Brinkmalm G, Teunissen CE, et al. Synaptic proteins in CSF as potential novel biomarkers for prognosis in prodromal Alzheimer's disease. Alzheimers Res Ther 2018; 10: 5.
- 124 Schindler SE, Li Y, Todd KW, et al. Emerging cerebrospinal fluid biomarkers in autosomal dominant Alzheimer's disease. Alzheimers Dement 2019; 15: 655–65.
- 125 Teunissen CE, Elias N, Koel-Simmelink MJ, et al. Novel diagnostic cerebrospinal fluid biomarkers for pathologic subtypes of frontotemporal dementia identified by proteomics. Alzheimers Dement (Amst) 2016; 2: 86–94.
- 126 Suarez-Calvet M, Capell A, Araque Caballero MA, et al. CSF progranulin increases in the course of Alzheimer's disease and is associated with sTREM2, neurodegeneration and cognitive decline. EMBO Mol Med 2018; 10: e9712.
- 127 Mapstone M, Cheema AK, Fiandaca MS, et al. Plasma phospholipids identify antecedent memory impairment in older adults. Nat Med 2014; 20: 415–18.
- 128 de Leeuw FA, Peeters CFW, Kester MI, et al. Blood-based metabolic signatures in Alzheimer's disease. Alzheimers Dement (Amst) 2017; 8: 196–207.
- 129 Stamate D, Kim M, Proitsi P, et al. A metabolite-based machine learning approach to diagnose Alzheimer-type dementia in blood: results from the European Medical Information Framework for Alzheimer disease biomarker discovery cohort. Alzheimers Dement (N Y) 2019; 5: 933–38.
- 130 van der Lee SJ, Teunissen CE, Pool R, et al. Circulating metabolites and general cognitive ability and dementia: evidence from 11 cohort studies. Alzheimers Dement 2018; 14: 707–22.
- 131 Bridel C, van Wieringen WN, Zetterberg H, et al. Diagnostic value of cerebrospinal fluid neurofilament light protein in neurology: a systematic review and meta-analysis. *JAMA Neurol* 2019; 76: 1035–48.
- 132 Al Shweiki MR, Steinacker P, Oeckl P, et al. Neurofilament light chain as a blood biomarker to differentiate psychiatric disorders from behavioural variant frontotemporal dementia. J Psychiatr Res 2019; 113: 137–40.
- 133 Olsson B, Alberg L, Cullen NC, et al. NFL is a marker of treatment response in children with SMA treated with nusinersen. J Neurol 2019; 266: 2129–36.
- 134 Barro C, Benkert P, Disanto G, et al. Serum neurofilament as a predictor of disease worsening and brain and spinal cord atrophy in multiple sclerosis. *Brain* 2018; 141: 2382–91.

- 135 Sormani MP, Haering DA, Kropshofer H, et al. Blood neurofilament light as a potential endpoint in phase 2 studies in MS. Ann Clin Transl Neurol 2019; 6: 1081–89.
- 136 Janelidze S, Stomrud E, Palmqvist S, et al. Plasma beta-amyloid in Alzheimer's disease and vascular disease. Sci Rep 2016; 6: 26801.
- 137 Ovod V, Ramsey KN, Mawuenyega KG, et al. Amyloid β concentrations and stable isotope labeling kinetics of human plasma specific to central nervous system amyloidosis. Alzheimers Dement 2017; 13: 841–49.
- 138 Schindler SE, Bollinger JG, Ovod V, et al. High-precision plasma beta-amyloid 42/40 predicts current and future brain amyloidosis. Neurology 2019; 93: e1647–59.
- 139 Verberk IMW, Slot RE, Verfaillie SCJ, et al. Plasma amyloid as prescreener for the earliest Alzheimer pathological changes. Ann Neurol 2018; 84: 648–58.
- 140 Palmqvist S, Janelidze S, Stomrud E, et al. Performance of fully automated plasma assays as screening tests for Alzheimer disease-related beta-amyloid status. *JAMA Neurol* 2019; 76: 1060–69.
- 141 Nabers A, Perna L, Lange J, et al. Amyloid blood biomarker detects Alzheimer's disease. EMBO Mol Med 2018; 10: e8763.
- 142 Teunissen CE, Chiu MJ, Yang CC, et al. Plasma amyloid-beta (Abeta42) correlates with cerebrospinal fluid abeta42 in Alzheimer's disease. J Alzheimers Dis 2018; 62: 1857–63.
- 143 Thijssen EH, La Joie R, Wolf A, et al. Diagnostic value of plasma phosphorylated tau181 in Alzheimer's disease and frontotemporal lobar degeneration. Nat Med 2020; 26: 387–97.
- 144 Janelidze S, Mattsson N, Palmqvist S, et al. Plasma p-tau181 in Alzheimer's disease: relationship to other biomarkers, differential diagnosis, neuropathology and longitudinal progression to Alzheimer's dementia. Nat Med 2020; 26: 379–86.
- 145 Palmqvist S, Janelidze S, Quiroz YT, et al. Discriminative accuracy of plasma phospho-tau217 for Alzheimer disease vs other neurodegenerative disorders. JAMA 2020; 324: 772–81.
- 146 Mattke S, Cho SK, Bittner T, Hlavka J, Hanson M. Blood-based biomarkers for Alzheimer's pathology and the diagnostic process for a disease-modifying treatment: projecting the impact on the cost and wait times. Alzheimer Dement 2020; 12: e12081.
- 147 WHO. Risk reduction of cognitive decline and dementia: WHO guidelines. 2019. Geneva: World Health Organization. https://www.who.int/mental_health/neurology/dementia/guidelines_risk_reduction/en/ (accessed Sept 1, 2020).
- 148 SPRINT MIND Investigators for the SPRINT Research Group, Williamson JD, Pajewski NM, et al. Effect of intensive vs standard blood pressure control on probable dementia: a randomized clinical trial. JAMA 2019; 321: 553–61.
- 149 Kivipelto M, Mangialasche F, Ngandu T. Lifestyle interventions to prevent cognitive impairment, dementia and Alzheimer disease. Nat Rev Neurol 2018; 14: 653–66.
- 150 Ngandu T, Lehtisalo J, Solomon A, et al. A 2 year multidomain intervention of diet, exercise, cognitive training, and vascular risk monitoring versus control to prevent cognitive decline in at-risk elderly people (FINGER): a randomised controlled trial. *Lancet* 2015; 385: 2255–63.
- 151 Solomon A, Turunen H, Ngandu T, et al. Effect of the apolipoprotein E genotype on cognitive change during a multidomain lifestyle intervention: a subgroup analysis of a randomized clinical trial. JAMA Neurol 2018; 75: 462–70.
- 152 Andrieu S, Guyonnet S, Coley N, et al. Effect of long-term omega 3 polyunsaturated fatty acid supplementation with or without multidomain intervention on cognitive function in elderly adults with memory complaints (MAPT): a randomised, placebocontrolled trial. *Lancet Neurol* 2017; 16: 377–89.
- 153 Moll van Charante EP, Richard E, Eurelings LS, et al. Effectiveness of a 6-year multidomain vascular care intervention to prevent dementia (preDIVA): a cluster-randomised controlled trial. *Lancet* 2016; 388: 797–805.

- 154 Delrieu J, Payoux P, Carrie I, et al. Multidomain intervention and/ or omega-3 in nondemented elderly subjects according to amyloid status. Alzheimers Dement 2019; 15: 1392–401.
- 155 Kivipelto M, Mangialasche F, Snyder HM, et al. World-Wide FINGERS Network: a global approach to risk reduction and prevention of dementia. Alzheimers Dement 2020; 16: 1078–94.
- 156 Soininen H, Solomon A, Visser PJ, et al. 24-month intervention with a specific multinutrient in people with prodromal Alzheimer's disease (LipiDiDiet): a randomised, double-blind, controlled trial. *Lancet Neurol* 2017; 16: 965–75.
- 157 Soininen H, Solomon A, Visser PJ, et al. 36-month LipiDiDiet multinutrient clinical trial in prodromal Alzheimer's disease. Alzheimer Dement 2020; published online Sept 13. https://doi.org/ 10.1002/alz.12172.
- 158 Atri A, Frolich L, Ballard C, et al. Effect of idalopirdine as adjunct to cholinesterase inhibitors on change in cognition in patients with Alzheimer disease: three randomized clinical trials. *JAMA* 2018; 319: 130–42.
- 159 Cummings J, Ballard C, Tariot P, et al. Pimavanserin: potential treatment for dementia-related psychosis. J Prev Alzheimers Dis 2018; 5: 253–58.
- 160 Halpern R, Seare J, Tong J, Hartry A, Olaoye A, Aigbogun MS. Using electronic health records to estimate the prevalence of agitation in Alzheimer disease/dementia. Int J Geriatr Psychiatry 2019; 34: 420–31.
- 161 Liu CS, Ruthirakuhan M, Chau SA, Herrmann N, Carvalho AF, Lanctot KL. Pharmacological management of agitation and aggression in Alzheimer's disease: a review of current and novel treatments. Curr Alzheimer Res 2016; 13: 1134–44.
- 162 Janto K, Prichard JR, Pusalavidyasagar S. An update on dual orexin receptor antagonists and their potential role in insomnia therapeutics. J Clin Sleep Med 2018; 14: 1399–408.
- 163 Cummings JC, Lee G, Ritter A, Sabbagh M, Zhong K. Alzheimer's disease drug development pipeline: 2020. Alzheimer Dement 2020; 6: e12050.
- 164 Sevigny J, Chiao P, Bussiere T, et al. The antibody aducanumab reduces $A\beta$ plaques in Alzheimer's disease. *Nature* 2016; **537**: 50–56.
- 165 van Dyck CH. Anti-amyloid-β monoclonal antibodies for Alzheimer's disease: pitfalls and promise. Bio Psychiatry 2017; 83: 311–19.
- 166 Lilly. News release: Lilly's donanemab slows clinical decline of Alzheimer's disease in positive phase 2 trial. Jan 11, 2021. https://investor.lilly.com/news-releases/news-release-details/lillys-donanemab-slows-clinical-decline-alzheimersdisease#:~:text=11%2C%202021%20%2FPRNewswire%2F%20 %2D%2D,results%20from%20Eli%20Lilly%20and (accessed Jan 24, 2021)
- 167 Egan MF, Kost J, Voss T, et al. Randomized trial of verubecestat for prodromal Alzheimer's disease. N Engl J Med 2019; 380: 1408–20.
- 68 Cummings J, Blennow K, Johnson K, et al. Anti-tau trials for Alzheimer's disease: a report from the EU/US/CTAD Task Force. *J Prev Alzheimers Dis* 2019; **6:** 157–63.
- 169 Wang X, Sun G, Feng T, et al. Sodium oligomannate therapeutically remodels gut microbiota and suppresses gut bacterial amino acids-shaped neuroinflammation to inhibit Alzheimer's disease progression. Cell Res 2019; 29: 787–803.
- 170 Bateman RJ, Benzinger TL, Berry S, et al. The DIAN-TU Next Generation Alzheimer's prevention trial: adaptive design and disease progression model. Alzheimers Dement 2017; 13: 8–19.
- 171 Alzforum. In DIAN-TU, Gantenerumab Brings Down Tau. By a Lot. Open Extension Planned. April 10, 2020. https://www.alzforum. org/news/conference-coverage/dian-tu-gantenerumab-brings-down-tau-lot-open-extension-planned (accessed Jan 26, 2021).
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