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The β -Secretase BACE1 in Alzheimer's Disease

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Authors

Hampel, Harald Vassar, Robert De Strooper, Bart <u>et al.</u>

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Harald Hampel, Robert Vassar, Bart De Strooper, John Hardy, Michael Willem, Neeraj Singh, John Zhou, Riqiang Yan, Eugeen Vanmechelen, Ann De Vos, Robert Nisticò, Massimo Corbo, Bruno Pietro Imbimbo, Johannes Streffer, Iryna Voytyuk, Maarten Timmers, Amir Abbas Tahami Monfared, Michael Irizarry, Bruce Albala, Akihiko Koyama, Naoto Watanabe, Teiji Kimura, Lisa Yarenis, Simone Lista, Lynn Kramer, and Andrea Vergallo

ABSTRACT

BACE1 (beta-site amyloid precursor protein cleaving enzyme 1) was initially cloned and characterized in 1999. It is required for the generation of all monomeric forms of amyloid- β (A β), including A β_{42} , which aggregates into bioactive conformational species and likely initiates toxicity in Alzheimer's disease (AD). BACE1 concentrations and rates of activity are increased in AD brains and body fluids, thereby supporting the hypothesis that BACE1 plays a critical role in AD pathophysiology. Therefore, BACE1 is a prime drug target for slowing down A β production in early AD. Besides the amyloidogenic pathway, BACE1 has other substrates that may be important for synaptic plasticity and synaptic homeostasis. Indeed, germline and adult conditional BACE1 knockout mice display complex neurological phenotypes. Despite BACE1 inhibitor clinical trials conducted so far being discontinued for futility or safety reasons, BACE1 remains a well-validated therapeutic target for AD. A safe and efficacious compound with high substrate selectivity as well as a more accurate dose regimen, patient population, and disease stage may yet be found. Further research should focus on the role of A β and BACE1 in physiological processes and key pathophysiological mechanisms of AD. The functions of BACE1 and the homologue BACE2, as well as the biology of A β in neurons and glia, deserve further investigation. Cellular and molecular studies of BACE1 and BACE2 knockout mice coupled with biomarker-based human research will help elucidate the biological functions of these important enzymes and identify their substrates and downstream effects. Such studies will have critical implications for BACE1 inhibition as a therapeutic approach for AD.

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BACE1 (beta-site amyloid precursor protein [APP] cleaving enzyme 1) is an aspartyl protease of the pepsin family and was discovered in 1999. BACE1 is a type I transmembrane protein, which makes it distinct from other peptidases of the same family, such as cathepsin D and E, which do not harbor a transmembrane domain (1–4). BACE1 is widely expressed in the brain, particularly in neurons, oligodendrocytes, and astrocytes, with particular abundance in various neuronal cell types (1–4). At the subcellular level, BACE1 localizes on the plasma membrane and in the endosomal compartments and was detected in healthy synaptic terminals and dystrophic neurites surrounding amyloid- β (A β) plaques (2,5,6). BACE1 is homologous with another membrane-bound secretase of the pepsin family, BACE2. The two proteases share 59% of their amino acid sequence and are composed of identical structural domain (1–4).

The active site of both secretases consists of 2 aspartic acid residues in their extracellular domains; they have 21-residue helical transmembrane domains and short cytoplasmic C-terminal domains (1–4). Three disulfide bonds help stabilize their tertiary structure, and the secretases are known to be glycosylated on several asparagine residues. Posttranslational

modifications are known on BACE1 secretase and are important for lipid raft localization, phosphorylation for cellular trafficking, and ubiquitination for degradation (see below).

BACE1 and BACE2 are expressed in the same cell types in the brain, only with BACE2 being much less abundant (7,8). However, BACE2 is thought to be more active in peripheral tissues, with particular functions in melanocytes (9) and pancreatic β cells (10).

BACE1: GENETIC AND EPIGENETIC

Genetic Mutations Affect BACE1 Cleavages

Mutations in the BACE1 gene have not yet been linked to Alzheimer's disease (AD) pathogenesis. However, mutations in APP near β -secretase sites are assumed to be either protective or causing early onset. The most prominent one is the Swedish mutation (K670M671 to N670L671 mutation at the cleavage P²-P¹ subsite), which increases processing of APP at the β site by 10- to 50-fold and causes early onset of AD (11).

Whole-genome sequencing studies identified a genetic variant of APP, significantly more frequent in Icelandic and

Scandinavian populations, that provides resilience against age-related brain A β deposition and AD (12,13).

Intriguingly, an Ala residue at the cleavage $P^{3\prime}$ subsite mutated to Thr (A673T) in APP confers an intrinsic biochemical resistance to cleavage by BACE1, resulting in less A β production overall and also in generation of peptides that are less prone to aggregation (12,13).

In particular, the A673T mutation is associated with a different BACE1 recognition motif at the position P2 (A673T), resulting in 20% to 30% lower soluble APP β (sAPP β) levels compared with control subjects (12,13). The A673T mutation appears to protect against AD and age-related cognitive decline as a result because of suppressed cleavage of APP at the β site, and carriers of the A673T variant in humans have about 28% lower levels of A β_{40} and A β_{42} in plasma compared with control subjects (12,13).

In addition, the E682K mutation in APP, located at the P^{1/} β site, favors A β production by suppressing the cleavage at the β' site to favor cleavage at the β site (12,13).

DNA Methylation

Epigenome-wide association studies on neuronal and glial cells sorted from postmortem brains of patients with AD and healthy donor subjects have revealed genes specific to neuronal cells, including APP, that undergo Braak stage-associated methylation changes (14). Interestingly, DNA methylation regulates BACE1 expression, likely owing to increased SP1 transcription factor binding to CpG sites on the BACE1 promoter region (15,16). More recently, a large cluster of significantly hypomethylated enhancers in the CpH sites were identified in prefrontal cortex neurons of individuals with severe AD pathology, and hypomethylation of these enhancers in the DSCAML1 gene likely upregulate BACE1 transcripts in AD (17). Similar DNA hypomethylation in the promoter region of APP enhances the expression of AD-related genes, including APP and PSEN1, leading to increased A β production (17,18).

DNA Acetylation

Although BACE1 expression and/or its activity have been extensively studied at both transcriptional and posttranslational levels, evidence of altered expression of BACE1 due to epigenetic acetylation remains weak. BACE1 messenger RNA levels are significantly increased in 3xTg mouse brains and in peripheral blood mononuclear cells from patients with AD but are much less elevated in mild cognitive impairment (MCI) compared with control subjects, and this increase is linked to H3 acetylation facilitating the accessibility of the BACE1 promoter (19). Decreasing acetylated H3 in the BACE1 promoter regions by galangin treatment in SH-SY5Y cells reduces the BACE1 messenger RNA level, likely related to upregulated endogenous HDAC1-mediated deacetylation (20).

Inhibition of histone acetyltransferase p300 by curcumin can also decrease acetylated H3 to reduce BACE1 transcripts (21). On the other hand, royal jelly peptides appear to regulate BACE1 expression through the control of HDAC1 (21). BACE1 messenger RNA levels are also regulated by diverse factors, including different transcription factors, that are summarized elsewhere (22,23).

Regulated BACE1 Expression by microRNA

Recently, the role of noncoding RNA, in particular microRNA, in regulating BACE1 has been gaining traction. Noncoding RNA, often referred to as microRNA, is 19 to 22 nucleotides long and often regulates gene and protein expression at the posttranscriptional level by binding to 3' untranslated region RNA to form a silencing complex. Over the past 2 decades, additional noncoding RNA has been shown to negatively regulate BACE1 expression: miR-107, miR-29c, miR-339-5p, miR-186, miR-195, miR-135b, miR-135a, miR-124, and miR-298/328 (see Table 1 for more details) (24–28).

The emerging field of AD transcriptomic signatures, including noncoding RNA that may be involved in negatively regulating BACE1 expression, can offer a promising platform for developing biomarkers. However, diagnostic and therapeutic applications of microRNAs remain challenging owing to multiple reasons, including restricted brain penetration and high-specificity concerns.

Post-translational Regulation of BACE1

BACE1 activity is regulated not only at the expression level but also by posttranslational modification. During the course of AD, BACE1 is subjected to numerous posttranslational modifications and plays a role in regulating signaling associated with $\mbox{A}\beta$ production and AD. BACE1 is a typical aspartyl protease with 2 active aspartate motifs (D₉₃TG and D₂₈₉SG) located in each lobe. BACE1 is first synthesized in the endoplasmic reticulum as a 501-amino-acid immature precursor protein, proBACE1. During maturation, BACE1 is N-glycosylated at 4 Asn sites (Asn153, Asn172, Asn223, and Asn354) in the endoplasmic reticulum lumen (4), and its prodomain (residues 1-21) is removed by furin-like proprotein convertases in the endoplasmic reticulum/early Golgi compartment (3,4). Although the presence of prodomain is not sufficient to suppress BACE1 activity (16), unlike most aspartyl proteases, suppressing glycosylation by site-directed mutagenesis of these aspartic acid residues reduces the protease activity of BACE1 (29). BACE1 is also reported to undergo sugar modifications by bisecting *N*-acetylglucosamine, which is high in brains. In brains of patients with AD, higher N-acetylglucosamine activity may partially cause increased BACE1 activity and A β deposition (30).

The role of sumoylation in regulating BACE1 functionality/ stability and activity is reported in both in vitro and in vivo AD mouse models; it was shown that sumoylation of residue K501 on BACE1 enhances its stability and A β -producing activity (31). By contrast, overexpression of nonsumoylated BACE1 mutant negated memory decline in wild-type mice and did not accelerate senile plaque formation (31).

Posttranslational modifications of BACE1 have been shown to alter its trafficking and/or localization (32). For example, palmitoylation of BACE1 at 4 cysteine residues (Cys474, Cys478, Cys482, and Cys485) in the transmembrane and C-terminal domains targets BACE1 to cholesterol-rich lipid rafts (32).

BACE1 undergoes phosphorylation at both Ser498 and Thr252, but phosphorylation at the Ser498 residue appears to regulate intracellular trafficking by shuttling/recycling BACE1 between endosome and plasma membrane (33). This modification at Ser498 has minimal or no effect on A β levels (34).

Table 1. Regulation of BACE1 Levels by miRNAs

miRNA	Mechanism of Action	Relevance in AD Brain	Reference(s)
miR-107	Downregulates BACE1 mRNA levels by binding to its 3' UTR. Other targets downregulated by miR-107 include granulin, cofilin, CDK5R1, and ADAM10.	Decreased miR-107 levels correlated with increased BACE1 levels in temporal cortex	(135,136)
miR-29c	Targets the 3' UTR of BACE1. Overexpression of miR-29c in cells reduced BACE1 protein expression and A β accumulation.	Decreased miR-29c expression levels correlated with increased BACE1 levels in sporadic AD	(137,138)
miR-186	Suppresses BACE1 expression by targeting the 3' UTR of BACE1 mRNA in primary neuronal cells. Inhibition of miR-186 increased BACE1 protein levels and $A\beta$ levels in neuro-2a cells.	Gradual reduction in miR-186 levels in 13- month-old mouse cortices during aging	(139)
miR-195	Levels inversely correlated with the protein level of BACE1 in SAMP8 mice. miR-195 overexpression in N2a/WT cells decreased the BACE1 protein and A β levels.	Downregulated in human AD CSF samples	(140)
miR-124	Targets BACE1 by binding to 3' UTR. miR-124 mimetic dramatically downregulated BACE1 mRNA and protein, while inhibition of miR- 124 significantly increased the expression in SH-SY5Y cells.	Expression significantly reduced in the hippocampus and anterior temporal cortex in AD brain	(141)
miR-298 and -328	Recognize specific binding sites in the 3' UTR of BACE1 mRNA and regulate BACE1 protein expression in N2a neuronal cells.		(142)

Aβ, amyloid-β; AD, Alzheimer's disease; BACE1, beta-site amyloid precursor protein cleaving enzyme 1; CSF, cerebrospinal fluid; miRNA, microRNA; mRNA, messenger RNA; SAMP8, senescence-accelerated prone mice; UTR, untranslated region; WT, wild-type.

Others noted that phosphorylation at Thr252 by p25/Cdk5 was associated with increased BACE1 activity and A β production (35). Similarly, acetylation of BACE1 has also been identified at 7 lysine resides (36).

While acetylation imparts BACE1 protein stability (36), ubiquitination of BACE1 promotes its degradation in the lysosome and is impaired in AD (37,38).

Overall, as an aspartyl protease, BACE1 requires low acidic environments to reach optimal proteolytic activity, ideally at \sim pH 4.5 (39,40).

BACE1 PHYSIOLOGICAL FUNCTIONS AND PATHOPHYSIOLOGICAL IMPLICATIONS

Amyloidogenesis

BACE1 is the β -secretase enzyme that cleaves the transmembrane APP and, together with γ -secretase, generates A β species that in AD form increasingly large and conformationally complex soluble regionally deposited brain aggregates (see Figure 1). BACE1 cleavage of APP represents the rate-limiting step for A β production.

For this reason, BACE1 has been extensively studied in the context of brain amyloidogenesis and proven to be directly involved in $A\beta$ production based on data from several knockout (KO) mouse models (41–43). BACE1 has been pharmacologically targeted, with several inhibiting compounds entering clinical development and trials, effectively lowering $A\beta$ concentrations in human individuals.

Cleavage of APP by β -secretase liberates the soluble N-terminus of APP, while the C-terminal fragment (CTF- β or C99) remains bound to the membrane. Two mutations at the β -secretase cleavage site (the Swedish mutation KM/NL and an Italian variant A673V) were reported to be linked to familial AD and consequently raise the sAPP β level owing to strongly increased affinity of BACE1 for the changed recognition motif in APP (44).

The significant protective effect of the A673T variant against AD has provided a robust proof of principle for the

pathophysiological and pharmacological model that reducing the β -cleavage of APP may offer a resilient mechanism against the disease (12). In addition, preliminary evidence suggests that a longtime preventive reduction of BACE1 activity by 20% to 30% may be sufficient to prevent AD. However, to fully exploit the clinical and pharmacological implications of the A673T mutation, it is essential to understand whether A673T mutation A β aggregates display different toxicity rates compared with wild-type carriers and which molecular pathways underlie the finding that the A673T allele also protects against non-AD cognitive decline (12).

High BACE1 enzymatic activity was found in human AD brain extracts, which is consistent with the finding that neurons produce the highest levels of A β (3,45). The highest BACE1 protein level was reported in postnatal brain in mouse. Notably, a relatively large accumulation of BACE1 was described in neuritic dystrophies in the vicinity of A β plaques both in AD amyloidogenic mouse models and in AD brains, most likely by a posttranslational mechanism (2,5,6). Inducing autophagy in mutant human neurons augments retention of BACE1 in distal axons by autophagy, leading to enhanced β -cleavage of APP (46). To produce A β , the fragment CTF- β is cleaved by β -secretase, which finally releases A β into the extracellular space and releases the APP intracellular domain into the cytoplasm (2,5,6).

In a parallel competing nonamyloidogenic pathway, APP is cleaved by either α -secretase or η -secretase to release 2 additional variants of the APP ectodomain, namely sAPP- α and sAPP- η (47). The η -secretase pathway is used as an alternative when BACE1 is inhibited, with the consequence of increased A η - α activity with an effect on lowering neuronal activity by a so far unknown mechanism (47).

APP is a type I transmembrane protein and is highly expressed in neurons and abundant at the synapse (48–52). Its function remains elusive, although studies implicated it in maintenance of dendritic spines (53), neurotransmission (54), synaptic plasticity (55–58), and maintenance of excitation/in-hibition balance (58). Soluble APP is a GABA_B (gamma-aminobutyric acid type B) ligand that modulates synaptic



Figure 1. Schematic representation of amyloid precursor protein (APP) processing pathways. A β , amyloid- β ; A η - α , amyloid- η - α ; A η - β , amyloid- η - β ; BACE2, beta-site amyloid precursor protein cleaving enzyme 2; CTF, C-terminal fragment; LM, lipid membrane; p3, p3 fragment of the amyloid precursor protein; sAPP, soluble amyloid precursor protein. [Adapted with permission from Barão et al. (143).]

transmission (50). Rescue experiments in APP KO mice show that sAPP α is sufficient to restore defects in spine density (59), long-term potentiation, and spatial learning (60,61). Most of the ectodomain shedding of APP is performed by the α -secretase, which cleaves APP in the A β sequence and therefore is believed to protect against AD (47).

Although some evidence suggests that sAPP β seems much less active in in vitro assays of neural activity and plasticity than sAPP α (62), both sAPP α and sAPP β modulate basal synaptic transmission and short-term synaptic facilitation through binding to GABA_B receptor subunit 1a at the synapse (50). The sushi domains of GABA_B receptor subunit 1a are also able to bind full-length APP intracellularly (63). Interestingly, this interaction is crucial for axonal trafficking of the complex and affects presence of the receptor at the presynaptic terminals. Concomitantly, delivery of the complex to axonal cell surface diminishes the pool of APP available for BACE1 processing in endosomes and lowers A β production (63).

Synaptic Substrates

Initial evaluation of BACE1 KO mice focused on decreased production of A β shortly after BACE1 deficiency was revealed to be associated with subtle neurological deficits (43,64). Many of the BACE1 KO mouse phenotypes, such as the peripheral hypomyelination and synaptic deficits, are due to loss of function of substrates depending on the activation by BACE1. Lack of BACE1 was also reported to cause sensorimotor impairments, seizures, schizophrenia-like phenotypes, and retinal pathology (65–67). In 2012, the neuronal secretome of BACE1 was revealed by 2 independent studies in primary cultures (68,69), followed by a more complete repertoire of BACE1 substrates identified in mouse cerebrospinal fluid (CSF) (see Table 2). Among BACE1 substrates, NRG1 (neuregulin 1), SEZ6 (seizure-related protein 6), and CHL1 (close homologue of neural cell adhesion molecule L1) are known to have important neuronal functions and merit further discussion given the recent reports that BACE1 blockade in patients causes cognitive worsening.

NRG1 interacts with the epidermal growth factor receptor family of receptors to exert signaling cascades crucial for central nervous system development (70) and synaptic plasticity (70). BACE1 cleavage of NRG1 is essential for myelination in the central nervous system and peripheral nerves as well as for muscle spindle formation and maintenance (70). SEZ6 is important for dendritic branching, normal synaptic function, and motor coordination (1,71). In the mouse brain, soluble SEZ6 is almost exclusively produced by BACE1 (1,71). In its absence, achieved by genetic KO of SEZ6 or pharmacological inhibition of BACE1, synaptic plasticity is impaired. In particular, aberrant BACE1 processing of SEZ6 results in lower spine density and attenuated long-term potentiation in the hippocampus (1,71).

One of the most interesting substrates of BACE1 is CHL1. This cell adhesion molecule mediates axonal guidance in response to Sema3A (semaphorin 3A) (72,73). BACE1 cleavage yields an intracellular membrane-bound C-terminal fragment of CHL1 that is able to influence cytoskeleton dynamics, leading to growth cone collapse on presentation of the Sema3A cue. This particular substrate and/or its homologue L1CAM (L1 cell adhesion molecule), both cleaved by BACE1 (69,72–74), might be responsible for axonal organization defects in BACE1 KO mice (69,72–74). Intriguingly, axon guidance abnormalities in the hippocampus persist in the adult

Table 2. BACE1 Substrates and Their Physiological Roles

BACE1 Substrate	Physiological Role	
APP	Regulates neurite outgrowth, synapse formation, and synaptic plasticity; also regulates metal homeostasis	
APLP1	Regulates neurotransmission and plasticity in CNS synapses	
APLP2	Regulates synaptic function and plasticity in CNS	
Contactin 2	Regulates axon guidance, cell adhesion, and neurite outgrowth	
Jagged 1	Balances astrogenesis and neurogenesis; notch signaling influences neural plasticity, long-term memory, and synapse remodeling transmitter release through astrocytes	
CHL1	Regulates axon guidance, cell adhesion, neuronal migration, and neurite outgrowth	
Neurexin 1α and 3β	Regulates synapse assembly and maintenance	
NRG1	Regulates myelination, neuronal migration, and oligodendrocyte differentiation; also regulates synaptic transmission and plasticity via neurotransmitter receptors	
SEZ6	Regulates dendritic arborization and affects excitatory synapse development and maintenance and formation of neuronal circuits	
SEZ6L	Regulates synapse maturation, tumor suppressor function, and free cholesterol levels	
β (β 1–4) Auxiliary Subunits of the VGSC Subtype Nav1	Modulates cell surface expression of Nav1 sodium channels and thus controls excitability and propagation of action potentials in the neuronal membrane	
VGSC Accessory Subunits KCNE1 and KCNE2	Regulates cardiac and brain potassium channel subunit trafficking and maintenance of membrane excitability	

APLP1/2, amyloid-like protein 1/2; APP, amyloid precursor protein; CHL1, neural cell adhesion molecule L1; CNS, central nervous system; NRG1, neuregulin 1; SEZ6, seizure-related protein 6; SEZ6L, seizure-related protein 6 precursor protein; VEGFR1, vascular endothelial growth factor receptor 1; VGSC, voltage-gated sodium channels. [Adapted with permission from Das and Yan (144).]

conditional KO of BACE1, confirming an important role of the secretase in adult circuitry architecture as well as its established developmental functions (69,72–74).

BACE2 Physiological Functions: A Brief Update

Much less is known about the brain-relevant substrates and functions of the sister secretase BACE2 that is more prominently expressed in the colon, kidney, and pancreas (7). In pancreatic β cells, the proproliferative plasma membrane protein Tmem27 and islet amyloid polypeptide are proposed BACE2 substrates (75,76). BACE2 also processes the PMEL (pigment cell–specific melanocyte protein) in pigment cells (9). Pharmacological inhibition of BACE2 results in depigmentation, the most consistent side effect seen in preclinical studies of BACE1/BACE2 inhibition (9). Thus, BACE1 and BACE2 shedding events seem to be tissue, cell type, and context dependent, revealing the intricacy of their functions (7). Inhibition of BACE2 brain substrates might contribute to some of the side effects seen with BACE1 inhibitors.

Human postmortem studies showed high expression levels of BACE2 and strong correlation with BACE1 expression in neurons and astrocytes of patients with AD but not of control subjects (77). Huentelman *et al.* reported that different single nucleotide polymorphism variations at the BACE2 locus are associated with AD risk and altered A β processing (78). This was the first genetic evidence of a role for BACE2 in AD pathophysiology. Larger genome-wide association studies are needed to confirm such important findings that may have significant pharmacological implications.

BACE1 BIOMARKERS: STATE OF THE ART ON THE VALIDATION AND QUALIFICATION FOR DRUG-BIOMARKER CODEVELOPMENT PIPELINES

During the past 15 years, a few human in vivo studies reported good diagnostic performance of CSF BACE1 concentration (supposed to reflect gene expression levels) and activity in discriminating among patients with AD dementia, patients with MCI, and cognitively healthy individuals (79–85). Some studies also showed an association between BACE1 biomarkers and other core CSF and neuroimaging biomarkers of AD as well as the presence of apolipoprotein E (*APOE*) ε 4 allele (79–85). Significant predictive power regarding conversion from MCI to AD dementia has also been reported.

Regarding the blood matrix, BACE1 biomarkers have been investigated in both plasma (82,86,87) and platelets (88,89), displaying good correspondence with CSF and association with brain AD alterations.

A multicenter study reported good correspondence between CSF and plasma BACE1 concentrations (87). In particular, plasma BACE1 activity demonstrated good diagnostic performance in discriminating patients with AD dementia from patients with MCI and cognitively normal individuals (87). A recent study showed an association between plasma BACE1 concentrations and amyloid-positron emission tomography quantitative measures in a cohort of cognitively healthy individuals at risk for AD (86).

By contrast, some studies showed no acceptable diagnostic performance of BACE1 biomarkers (90–92) or no association between them and AD established biomarkers.

Moreover, lower levels of BACE1 were found in advanced dementia stages of AD, potentially owing to advanced neuronal and synaptic loss (93–95).

For more details about study populations and outcomes, see Table S1.

Interstudy results variability may be partially explained by several differences in the study design, confounding factors (e.g., disease stage, sex, APOE genotype, comorbidities), and the methodology. Regarding the latter, preanalytical factors such as the sample collection, processing, and storage protocols, as well as analytical factors such as assays used, are likely the most relevant determinants. The conflicting data reported above call for harmonization and standardization of research protocols.

In summary, there are enough promising data to boost development of BACE1 biomarkers and investigate whether they may enrich the current AD biomarkers panel and potentially support different contexts of use in BACE1 clinical trials, including target engagement and proof of mechanism, dose finding, efficacy, and safety monitoring.

HUMAN CLINICAL TRIALS WITH BACE INHIBITORS: A SCHEMATIC OVERVIEW

All BACE inhibitors investigated in randomized clinical trials were discontinued for either futility or safety reasons.

A Phase 3 trial of verubecestat conducted in mild to moderate patients with AD (EPOCH) was terminated owing to futility (96).

A Phase 2/3 trial of atabecestat investigated in preclinical individuals with AD (EARLY) was discontinued owing to liver toxicity (97).

The Phase 3 trials of lanabecestat investigated in patients with prodromal AD and mild AD (AMARANTH and DAYBREAK-ALZ, respectively) were stopped owing to futility (98).

A Phase 2 trial of LY3202626 involving patients with mild AD (NAVIGATE-AD) was discontinued owing to interim futility analysis (99).

The phase 2/3 trials of umibecestat (CNP520) investigating asymptomatic individuals at risk for AD (i.e., *APOE* ε 4 allele carriers) (GENERATION) were discontinued owing to cognitive worsening in the active treatment group (100).

The Phase 2 trial of elenbecestat (E2609), conducted in participants with MCI to moderate AD, was discontinued after recommendation by the Data Safety Monitoring Board owing to an unfavorable risk/benefit ratio (https://www.alzforum.org/therapeutics/elenbecestat).

See the Supplement for more details.

Potential Explanation Coming From Translational Data: Selectivity and Toxicity of BACE1 Inhibitors

To optimize next-generation BACE1 inhibitor clinical trials, it is essential to understand all major biological and pharmacological factors that might account for the high attrition rates of the previous trials. For this purpose, 3 points should be considered: 1) the biological rationale for BACE1 as a pharmacological target; 2) BACE 1/2 selectivity, druggability, and inhibition strength by dose adjustment as well as the overall benefit/risk ratio; and 3) timing of intervention with BACE1 inhibitors over the course of AD.

Regarding point 1, in consideration of all evidence reported in the sections above, we argue that the target has a good scientific rationale and was properly and extensively validated in experimental models of AD ahead of clinical studies. Regarding point 2, there are more than 40 known BACE1 substrates (see Table 2), and BACE inhibitors may block one or more of them, causing functional consequences. Current compounds tested in the clinic had variable amounts of selectivity for BACE1, but all exerted inhibition of BACE2 activity as well. Inhibiting BACE1 lowers A β production, but in combination with BACE2 inhibition the processing of a number of other substrates are blocked with potential negative impact, changing the benefit/risk ratio (see also Table S2).

Provided these data have not been systematically disclosed during early development of these compounds, it is arguable that all compounds in clinical development have been tested in standard good laboratory practice/good clinical practice toxicity studies as requested and reviewed by regulatory authorities. Therefore, it is conceivable that these preclinical tests did not demonstrate systematic biological signatures indicating the observed effects. Dose levels selected for all prior and current ongoing studies may be too high, targeting more than 50% inhibition of BACE1, leading to unwanted side effects, while potentially lower levels of inhibition could have been therapeutically active.

In this regard, it is not possible to rule out that an excessive suppression of BACE1 activity has determined cognitive dysfunction in patients with AD by depleting A β monomers that have physiological functions and display poor toxicity. A β monomers can trigger or sustain intracellular signaling essential for synaptic plasticity and homeostasis (101–103).

Concerning point 3 above, timing of intervention with BACE1 inhibitors over the course of AD, robust evidence indicates that cerebral A β accumulation is one of the earliest mechanistic alterations of the whole pathophysiological dynamic of AD (104–107).

The stronger correlation found between tau biomarkers with neurodegeneration outcome measures and long-term cognitive scores than between A^β markers and long-term cognitive scores has raised the question of whether A β pathophysiology triggers downstream pathways, including tau-mediated toxicity, and facilitates tau spreading (104-107). However, CSF and positron emission tomography longitudinal studies support the hypothetical pathophysiological model of AD for which amyloidosis proceeds, either promoting or being permissive to the spreading of tau pathology that is likely to drive disease clinical evolution (104-107). Such spatial and temporal dynamics of AD brain proteinopathies imply that Aβdirected treatments should be initiated at the earliest preclinical stages of the disease and not in the dementia stages. If so, BACE1 inhibitors started prior to the spreading of tau pathology may represent the most suitable path to pursue (108).

Some detrimental effects were induced during the initial phase of treatment and were irrespective of disease stage. It is conceivable that these negative effects may be caused by acute synaptic impairment via BACE1 inhibition for some substrates other than APP and should be assessed for reversibility after off-treatment.

New Potential BACE Inhibition Strategies: Drug Repositioning Programs and Modulation of Posttranslational Modification

A pursuable path for BACE1 inhibition may be represented by drug repurposing (also called drug repositioning or reprofiling) pipelines that aim at identifying new therapeutic avenues for already approved or investigational drugs irrespective of their original medical indication. Two recent animal trials used chronic exposure to lithium chloride—a therapeutic agent approved for major psychiatric disorders—and showed slowdown of cognitive decline and histopathological alterations associated with reduced BACE1 activity (109,110).

In particular, Wilson *et al.* reported that an innovative experimental formulation of lithium microdose release is associated with the lowering of BACE1 gene expression and overall cerebral A β accumulation (110), thereby confirming previous translational studies pointing at a potential neuroprotective effect of lithium (111).

Modulation of posttranslational modification of BACE1 may represent a viable therapeutic avenue. For instance, it was shown that AD mouse models expressing S-palmitoylationdeficient BACE1 had a significant decrease in A β burden and improved memory function, indicating that posttranslational Spalmitoylation of BACE1 influences A β pathogenesis (32). In line with this, HEK293 cells treated with KMI-574 specifically caused dissociation of BACE1 from lipid raft to nonraft membranes, and BACE1 processing activity was reduced (112). Hence, blocking BACE1 activity in the raft membrane is another venue for reducing A β deposition.

OPEN ISSUES

Several genetic data studies (306 autosomal dominant mutations plus the APP gene duplication and trisomy 21) and multimodal biomarker studies indicate that an imbalance between A β production and clearance plays a critical and early role in AD pathogenesis (104–106,113,114). A large amount of evidence also supports the hypothesis that cerebral A β deposition begins decades before AD clinical onset and prior to cortical spreading of tau pathology. However, the full understanding of the molecular dynamics of A β species, from loss of proteostasis to synaptic toxicity (either tau mediated or not), has not yet been achieved. In this context, BACE1 is established to play a key role in A β homeostasis and may have an important function in synaptic plasticity.

Incomplete knowledge of the physiological functions of BACE and its downstream pathways may have contributed to the failures of BACE1 inhibitor clinical trials.

Soluble A β peptides, oligomers, protofibrils, fibrils, and plaques still remain attractive targets (see Figure 2).

The recently discovered human APP Arctic (115) and E693 Δ (Osaka) (116) mutations show a type of AD with low cerebral deposition of plaques, as indicated by modest A β -positron emission tomography radiotracer binding (117) and higher production of oligomers and protofibrils that are likely to be the initiators of A β toxicity (115,116,118,119). These findings indicated that other forms of A β , besides fibrils and plaques, may trigger brain toxicity and contribute to AD-synaptic failure (115,116,118,119). Such evidence has fostered the development of novel biomarkers for tracking all A β aggregation states that may be used for novel surrogate end points.

Time for Biomarkers of Synaptic Dysfunctions?

From a functional standpoint, synaptogenic mechanisms of AD cognitive decline-that is, network activation and



Figure 2. BACE1 (beta-site amyloid precursor protein cleaving enzyme 1) and the amyloid- β (A β) cycle. Despite the fact that several clinical trials investigating anti-A β compounds did not reach primary end points, A β peptides, oligomers, protofibrils, and plaques still remain attractive targets. Of note, the nature of the toxic A β species remains unclear. Evidence suggests that, besides fibrils, dimeric or oligomeric A β species, but not monomeric A β peptides, cause neuronal hyperactivity and downstream toxicity in the vicinity of A β plaques.

deactivation deficits, abnormal oscillatory rhythmic activity, and network hypersynchrony—account for AD-related synaptic failure (120–122).

Clinical trials can benefit from resting-state and task-related functional magnetic resonance studies to detect aberrant patterns at the large-scale brain network level, including the default mode network (123–126). Brain networks' functional shifts, as well as their association with molecular dynamics, have already been described in aging and AD (127,128). Very recently, it was shown that genetic risk factors with a pleiotropic biological effect, such as the *APOE* ε 4 allele, affect the trajectories of the default mode network of cognitively healthy individuals at risk for AD (129,130).

Fluid biomarkers of synaptic dysfunction are currently under development (131–133). Neurogranin (a key regulator of the calcium-binding protein calmodulin), synaptogamin (a calcium sensor protein), and SNAP-25 (a component of the SNARE [soluble *N*-ethylmaleimide sensitive factor attachment protein receptor] complex) are the strongest candidates for in vivo tracking synaptic homeostasis (131–133).

CONCLUSIONS

Despite a number of robust discovery stage studies, a number of phase 3 small-molecule BACE1 inhibitor clinical trials did not reach primary end points, showed cognitive worsening, or were discontinued owing to safety reasons.

The failure of several BACE1 inhibitor clinical trials appears to involve insufficient understanding of BACE1 biology and physiology, limited knowledge of the natural history of AD and the optimal stage of disease at which to treat, and lack of biomarker-based outcomes and end points. In this regard, BACE1 trial failures may benefit from the previous pitfalls of γ -secretase pharmacological investigation (134).

The field needs to fully uncover the physiological functions of BACE1 substrates, including those involved in including synaptic homeostasis, and needs to better understand the physiological role(s) of BACE2.

During the past 20 years, several key genetic, epigenetic, and posttranslational factors have been established to influence BACE1 gene expression levels and enzymatic activity that may explain interindividual heterogeneity in BACE1related pathophysiological processes and drug response. While the research community continues to debate the most plausible biological and pharmacological explanations for BACE1 clinical trial failures, there is emerging evidence encouraging a new generation of compounds with an ultra APP selectivity BACE inhibitory effect.

Robust evidence indicates that BACE1 concentrations and the rate of activity assessed in body fluids, including plasma, may serve as multiple contexts of use (including trial enrollment, proof of mechanism, response and toxicity dose estimation, and drug resistance prediction) in drug biomarker codevelopment programs (see Figure S1).

Within this conceptual framework, BACE1-oriented therapies continue to represent a rational and central development area for time-sensitive and effective pathway (mechanism)– based preventive strategies for AD.

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ARTICLE INFORMATION

From the Neurology Business Group (HH, AATM, MI, BA, AK, LY, LK, AV), Eisai Inc., Woodcliff Lake, New Jersey; Department of Neurology (RV), Mesulam Center for Cognitive Neurology and Alzheimer's Disease, Feinberg School of Medicine, Northwestern University, Chicago, Illinois; Department of Neuroscience (NS, JZ, RY), University of Connecticut Health, Farmington, Connecticut; Sorbonne University (HH, SL, AV), GRC No. 21, Alzheimer Precision Medicine, and Institute of Memory and Alzheimer's Disease (SL. AV), Department of Neurology, Pitié-Salpêtrière Hospital, Paris; and Brain & Spine Institute (SL, AV), INSERM U 1127, CNRS UMR 7225, Paris, France; Department of Neurosciences (BDS, IV), KU Leuven; Centre for Brain and Disease Research (BDS, IV), VIB (Flanders Institute for Biotechnology), Leuven; ADx NeuroSciences NV (ED, ADV), Ghent; Reference Center for Biological Markers of Dementia (JS, MT), Institute Born-Bunge, University of Antwerp, Antwerp; UCB Biopharma SPRL (JS), Braine-I'Alleud; and Janssen Research and Development (MT), a division of Janssen Pharmaceutica, Beerse, Belgium: Dementia Research Institute (BDS) and Department of Molecular Neuroscience and Reta Lilla Weston Laboratories (JH), Queen Square Institute of Neurology, University College London, London, and ALBORADA Drug Discovery Institute (IV), University of Cambridge, Cambridge, United Kingdom; Chair of Metabolic Biochemistry (MW), Biomedical Center, Faculty of Medicine, Ludwig Maximilians University Munich, Munich, Germany; Laboratory of Neuropharmacology (RN), EBRI Rita Levi-Montalcini Foundation, and School of Pharmacy (RN), Department of Biology, University of Rome "Tor Vergata," Rome; Department of Neurorehabilitation

Sciences (MC), Casa Cura Policlinico, Milan; and Department of Research and Development (BPI), Chiesi Farmaceutici, Parma, Italy; Department of Epidemiology (AATM), Biostatistics and Occupational Health, McGill University, Montreal, Quebec, Canada; and Eisai Company Ltd. (NW, TK), Tokyo, Japan.

Address correspondence to Andrea Vergallo, M.D., at andrea_vergallo@ eisai.com, or Harald Hampel, M.D., Ph.D., at harald_hampel@eisai.com.

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