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PERSPECTIVE



Open drug discovery in Alzheimer's disease

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Abstract

Alzheimer's disease (AD) drug discovery has focused on a set of highly studied therapeutic hypotheses, with limited success. The heterogeneous nature of AD processes suggests that a more diverse, systems-integrated strategy may identify new therapeutic hypotheses. Although many target hypotheses have arisen from systems-level modeling of human disease, in practice and for many reasons, it has proven challenging to translate them into drug discovery pipelines. First, many hypotheses implicate protein targets and/or biological mechanisms that are under-studied, meaning there is a paucity of evidence to inform experimental strategies as well as high-quality reagents to perform them. Second, systems-level targets are predicted to act in concert, requiring adaptations in how we characterize new drug targets. Here we posit that the development and open distribution of high-quality experimental reagents and informatic outputs—termed target enabling packages (TEPs)—will catalyze rapid evaluation of emerging systems-integrated targets in AD by enabling parallel, independent, and unencumbered research.

KEYWORDS

Alzheimer's disease, drug development, drug targets, genomics, proteomics, systems biology, TEPs

1 | INTRODUCTION

By 2050, Alzheimer's disease (AD) will affect 150 million people worldwide. Caring for these patients will exact a tremendous societal and economic toll,¹ especially because there is a paucity of efficacious disease-modifying therapies approved for use in humans. The past two decades of drug discovery efforts in AD have been focused on a few therapeutic hypotheses. Although these were strongly supported by genetic evidence, none have been validated in patients.² There is, therefore, an urgent need to explore new target

hypotheses, which will necessitate adopting a higher risk tolerance in our target selection and drug discovery strategies. To this end, the National Institute on Aging has launched an initiative, called TREAT-AD (TaRget Enablement to Accelerate Therapy Development for AD). The goals of this program are to provide the community with a set of high-quality core reagents that are needed to explore new targets or disease mechanisms, and to pursue the most promising by generating new high-quality pharmacological tools that are active in animals (drug leads). TREAT-AD comprises two centers. Each is a collective of institutions; one is led from the Indiana University

This is an open access article under the terms of the Creative Commons Attribution-NonCommercial License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited and is not used for commercial purposes. © 2023 The Authors. Alzheimer's & Dementia: Translational Research & Clinical Interventions published by Wiley Periodicals LLC on behalf of Alzheimer's Association. School of Medicine and the other from Emory University (and includes Sage Bionetworks and the Structural Genomics Consortium [SGC]). Here we describe the strategy of the Emory-Sage-SGC TREAT-AD Center to enable rapid diversification of the AD drug discovery pipeline and apply the strategy to targets suggested by systems-level approaches.

Unbiased genome-wide studies from genetics and systems biology have led to an abundance of novel disease-associated proteins and pathways that have not been studied previously in relation to AD and may provide for new therapeutic hypotheses. This is both an advantage and a disadvantage: an advantage in that the approaches offer the potential for truly transformative ideas, but a disadvantage in that the research enterprise is reticent to work on previously understudied proteins.³ In AD, this phenomenon is particularly true, and can be quantified by evaluating the publication intensity over time for the 184 genes reported as associated with AD risk (ie, "AD-related genes") in the National Human Genome Research Institute - European Bioinformatics Institute (NHGRI-EBI) Unbiased genome-wide association studies (GWAS) catalog through 2015⁴ (Figure 1). Specifically, we quantified the publications that mention both Alzheimer's and the gene or protein name in the keywords, abstract, or title in two time periods: before genetic association studies were published (1950-2000) and after these studies were published (2018-2020). As might be expected, prior to 2000 almost all of the AD gene-based publications (93%) were focused on four genes (apolipoprotein E [APOE], amyloid precursor protein [APP], presenilin 1 [PSEN1], and presenilin 2 [PSEN2]) that were discovered through early-onset familial AD studies. But despite strengthening evidence from clinical trials that some anti-amyloid immunotherapies informed by these four genes/proteins have only small although significant benefit in at least a subset of AD cases,⁵ clearly more robust therapeutic strategies are urgently needed. Nonetheless, these four genes remained the most studied genes in

RESEARCH IN CONTEXT

- SYSTEMATIC REVIEW: The authors reviewed the scientific literature using standard approaches (GoogleScholar, PubMed) to develop our perspective on open drug discovery in Alzheimer's disease (AD).
- INTERPRETATION: We present an overview of a novel approach to target validation in AD, which utilizes open science practices to accelerate drugs through the drug discovery pipeline.
- 3. FUTURE DIRECTIONS: Our center is generating openly available resources to enable target validation and drug discovery efforts in AD. Future directions based on our work include the adoption of these specific resources in experimental programs. For example, with an in vivo chemical or molecular probe inhibitor to a target of interest, a researcher could design experiments to examine a role for that target in relevant AD-related phenotypes in human induced pluripotent stem cells (iPSCs) and relevant AD mouse models.

the 2018–2020 period, garnering fully 73% of the AD-related gene publications. In contrast, the number of genes associated with AD risk continues to explode with larger well-powered studies, with at least 75 gene associations confirmed—most with links to additional, understudied biology.⁶

In our center, we hypothesize that researchers will be more willing to devote attention to understudied genes and proteins if provided with open access to high-quality molecular and cellular research tools,³





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TABLE 1 Quality acceptance criteria for target enabling package components.

TEP component	Quality criteria
Genetic tools	
CRISPR-deleted in selected cell lines	 Parental cell line or iPSC expressing "native" protein levels Sequencing of CRISPR target site confirms knockout of all alleles Target RNA undetectable using RT-PCR Target protein undetectable by immunoblot Growth rate and viability of KO lines are communicated
CRISPRa, CRISPRi, or induced-degradation in relevant cell lines (or iPSCs) or RNAi	 gRNA/degrader/RNAi-induced decrease/increase in protein levels by > 10 fold (using validated antibodies or quantitative mass spectrometry) Maintaining cell phenotype/pluripotency Genomic stability (karyotyping and analysis of genome stability by qPCR) Report growth parameters of engineered cell vs WT cell line
CRISPR-edited cell lines containing disease-linked mutations	 Parent iPSC-derived cell type expresses "native" protein levels Sequencing of CRISPR target site confirms targeted genetic mutation in both alleles Genomic stability (karyotyping and analysis of genome stability by qPCR) Maintaining cell phenotype/pluripotency Report growth rate and viability of KO lines
Validated antibodies	
General	Preferably monoclonal or recombinant and with registered Research Resource Identifiers (RRIDs)
WB	 Immunoreactivity with band(s) at the predicted size(s) in lysates from native expressing cells; no reactivity with an isogenic KO cell under the same conditions. Oddly sized bands or multiple bands may be acceptable if they do not occur in the KO cells Ideally the rest of the lane is blank, but some nonspecific bands might be acceptable If protein is essential, then the above with multiple RNAis
IF	 Immunoreactivity in cells, no signal in isogenic KO or nonexpressing knockdown cells under the same conditions Native and KO cells are stained contemporaneously on the same coverslip (each cell line is marked with a different fluorescent protein)
IP	 Desirable: full depletion of the target protein from lysates (by WB if there is a WB-validated antibody) IP-MS: the target protein is recovered among the top hits in MS/MS analysis of immunoprecipitates from WT cells or tissues
IHC	• Immunoreactivity in tissue or cell blocks, no signal in isogenic KO or nonexpressing knockdown tissue or cell blocks under the same conditions
Protein expression	
Purified protein/functional domain	 Expression construct fully sequenced Reproducible expression/purification protocol Identity of purified protein verified by intact mass spectrometry Thermostability profile demonstrates a folded state Data provided on protein stability: uniform pattern on size exclusion chromatography, optimal buffer, freeze-thaw behavior
Modified versions of the target protein if required for assays	 The above, plus evidence of the correct tags/modification Intact biochemical activity
If relevant, assembly into relevant protein complex	• The above, plus evidence of copurification of the purified proteins (eg, size exclusion chromatography)
Fundamental assays	
 Biophysical assays: Assay demonstrating binding of any ligand to the target protein (eg, SPR, DSF, Prometheus, BLI, FP, ITC) Optional: 1-D, HSQC NMR profiles 	 Protein stable and giving interpretable data under assay conditions without aggregation or other detection artifacts Measurement of binding of a test compound, ideally with correlation between changes in the chemical structure and binding affinity (ie, a structure–activity relationship)
 In vitro biochemical assays: Ligand displacement assay Quantitative assay for SAR (additional criterion) Quantitative assay for HTS (additional criterion) 	 Protein stable and giving interpretable data under assay conditions without aggregation or other detection artifacts Control ligand binds with saturable, specific binding isotherm Assay suitable for measuring IC50 values in the range of 0.01-10.00 μM HTS compatibility: available (affordable) substrate, suitable for automation, robust assay performance (S/B >4, Z' >0.5) in a HTS format (384- or 1536-well plates)

TABLE 1 (Continued)

TEP component	Quality criteria
 Enzyme assay Quantitative assay for SAR (additional criterion) Quantitative assay for HTS (additional criterion) 	 Protein stable and giving interpretable data under assay conditions without aggregation or other detection artifacts. Characterized mechanism and kinetic parameters (Km, kcat) including demonstration of reversibility/irreversibility (kinact/Ki) Assay suitable for measuring IC₅₀ values in the range of 0.01-10 μM HTS compatibility: available (affordable) substrate, suitable for automation, robust assay performance (S/B >4, Z' >0.5) in a HTS format (384- or 1536-well plates)
 Intracellular target engagement: Thermal denaturation profile of the protein in cells, for cellular thermal shift assay (CETSA) or Assay with unique readout of target binding to a competitive tracer (i.e., BRET) or Assay of an enzymatic activity or signaling event uniquely attributed to the target 	 Thermal denaturation profile of the target in cells established, Tm <60°C. Preferably a shift seen with a control agent known to bind Target-dependent readout established with sensitivity for measuring 1 μM binding. Test compound shows tracer displacement. S/B > 1.5 Target-dependent readout established with sensitivity for measuring 1 μM binding. Test compound shows inhibition. S/B > 1.5
 Cellular phenotypes Assay measuring cellular function uniquely attributed to the target, such as migration or immune cell activation Control assay measuring cell viability or toxicity 	 Positive and negative experiments with mechanistically similar reagents are distinguishable in blind experiments with machine readout Little cell toxicity seen using mechanistically related reagents under the conditions and duration of the assay. Correlation between changes in the chemical structure, activity in primary assay and phenotypic readout (i.e., a structure–activity relationship)
Low-throughput assays (additional criterion)	 Replication in at least 3 biologically independent samples with p value < .05 compared with control in nonplate reader-based assays
High-throughput assays (additional criterion)	• Robust assay performance (S/B > 4, Z' >0.5) in a HTS format (384- or 1536-well plates)
Structures	
Crystal structures	Details at https://www.thesgc.org/scientists/strubio
Chemical probes	
	 Potency refers to EC₅₀/IC₅₀ In vitro potency: <100 nM potency in activity/binding and biophysical interaction assay Selectivity: >30-fold selectivity over homologous (target-class members) proteins and confounding targets in pathway. Data available to assess activity in general pharmacology profiling assays (eg, CEREP); selectivity sufficient to use the probe in an in vivo study is required Cellular potency and target engagement: <1 μM with quantitative evidence of target engagement in cells via direct binding or activity on a proximal readout of target function Pharmacokinetic (PK) properties: sufficient half-life and CNS penetration to dose orally once daily (QD) or twice daily (BID) and achieve free brain fraction > cellular EC₅₀ Functional activity in cell and in vivo AD models consistent with AD target hypothesis at a tolerated dose Cellular probes will meet all criteria except PK and in vivo activity in an AD model
Biological probes	
	 In vitro affinity of at least 20 nM Can immunoprecipitate its endogenous full-length protein target from a complex mixture with high selectivity Significant efficacy in one or more cell-based assays

Abbreviations: BID, twice daily; BRET, Bioluminescence Resonance Energy Transfer; CETSA, cellular thermal shift assay; CNS, central nervous system; CRISPR (Clustered Regularly Interspaced Short Palindromic Repeats) - deleted; CRISPRa (CRISPRactivation); CRISPRi (CRISPRinterference); HTS, high-throughput screening; IF, immunofluorescence; IHC, immunistochemistry; IP, immunoprecipitation; IP-MS, IP-mass spectrometry; KO, knock-out; PK, pharmacokinetic; QD, once daily; SAR, structure activity relationship; Tm, melting point, WB, western blot.

and our aim is to generate these tools for under-studied proteins linked to AD.

1.1 | The target enabling packages (TEPs): Open reagents for target evaluation

Our center is committed to providing a core set of data and reagents (target-enabling packages [TEPs]) for a range of protein targets nominated through systems approaches and otherwise by the community.⁷ As we define it, the minimal TEP comprises the purified protein, validated renewable or recombinant antibodies for a range of applications, and a verified gene knockout cell line. These minimal TEP components are the basic and validated experimental tools needed to start probing target biology. Each TEP component must meet our quality criteria prior to its release and open distribution. The quality criteria are listed in Table 1. For prioritized targets, we are generating additional TEP components that might include crystal structures, biochemical and biophysical assays, and cell-based assays that would facilitate the identification of drug candidates. For each protein target, we pre-define the composition of the TEPs, based on its function and localization, and with input from the wider community. All TEP components are being made available through trusted commercial and/or non-profit distributors.

In practice, the first step in assembling a TEP is to secure and assay existing commercial reagents (such as antibodies and cell lines) to determine if they meet our quality criteria. If any do, we will make this information freely available. If we identify reagents that are missing, our center will produce and characterize them. To date our center has generated TEPs for seven AD-relevant targets. These TEPs are distributed through the AD Knowledge Portal.⁸

The generation of reagents for the chromosome 9 open reading frame 72 (C9ORF72) protein provides an excellent example of the potential impact of TEPs, and of our approach. The C9orf72 gene is one of the most commonly mutated genes in individuals with frontotemporal dementia or amyotrophic lateral sclerosis (ALS). This link was discovered almost 10 years ago, and in the immediate years after there was significant interest in its molecular and cellular biology. Many of the studies on C9ORF72 conflicted; however, different studies localized the protein to different subcellular components. The situation was clarified by focusing on the quality of the reagents. In 2018, McPherson et al. launched a TEP project to create or identify existing tools for the C9ORF72 proteins.⁹ Focusing on antibodies to start, they used knockout cells as controls to screen commercially available antibodies for those that recognized the protein specifically and selectively. They found a high-performing antibody, made the results publicly available, and used the antibody to localize C9ORF72 to the lysosomal membrane. Their work also revealed that many of the antibodies used previously to localize the protein to other compartments did not in fact even recognize the protein. The trajectory of C9ORF72 cell biology research has now changed in response to the availability of high-quality, specific, and selective reagents.

TABLE 2 TREAT-AD emerging gene targets in Alzheimer's disease for TEP development.

Target genes	
APOE	NDUFS2
APP	NTN1
ARHGEF2	NTN3
BDH2	NXPH1
C4A	OLFML3
CAPN2	PAK1
CD44	PLEC
CNN3	PRDX1
COL11A1	PRDX6
COL25A1	PTN
CTHRC1	QPRT
СТЅН	RABEP1
DAG1	RENBP
DDX1	SDC4
DHX58	SFRP1
ECE1	SLIT1
EPHX2	SLIT2
FCER1G	SMOC1
FLT1	SNX32
FRZB	SPOCK1
GPC5	SPOCK2
GPNMB	SPOCK3
HTRA1	SPON1
IFIH1	STX4
LRP1	SYK
MDK	TICAM1
MSN	TMEFF2

1.2 | **TEP** targets from AD systems biology

The current target portfolio in our center (Table 2) was assembled by evaluating prioritized proteins that were nominated as promising, yet understudied, targets for AD through the Accelerating Medicines Partnership in AD (AMP-AD) program.¹⁰⁻¹⁵ We select targets using an iterative process in which we evaluate lists of targets from AMP-AD studies by calculating an unbiased target risk score, summarizing existing literature, and assessing the tractability and therapeutic intervention potential of each target. Our full target scoring pipeline is detailed in Cary et al.,¹⁶ but briefly, all targets are scored across a hierarchy of criteria that is broadly organized into genetic, multiomic, neuropathological, and literature-based metrics (Figure 2). Following this initial scoring, targets are evaluated for therapeutic potential, with an emphasis on tractability. If targets are judged to not be tractable, the starting gene lists are expanded to incorporate other genes that

PSEN1





SYK



FIGURE 2 Representative histograms of TaRget Enablement to Accelerate Therapy Development for AD (TREAT-AD) consortium target-ranking scores encompassing genomics, genetics, and literature evidence for a classical Alzheimer's disease (AD) example target, presenilin1 (PSEN1), as well as a target prioritized by our TREAT-AD center, spleen-associated tyrosine kinase (SYK).

are in the same network (using co-expression or pathway annotation information). $^{16} \,$

In addition to selection of individual understudied targets, we are developing resources to interrogate a set of functionally connected targets, by developing TEPS for multiple targets within one specific protein module. Deep proteomic profiling study of AD and control brains from AMP-AD revealed new AD-related protein co-expression modules that were highly preserved across cohorts and brain regions.¹⁵ One of these modules, Module 42, ontologically linked to the matrisome, was highly associated with the global burden of pathology in the brain, with a correlation coefficient of 0.75. Strikingly, the small group of 32 proteins co-expressed in this module was enriched in AD risk genes and included APP and apoE. Evaluation of the

potential therapeutic relevance of matrisome module proteins is a considerable challenge given the paucity of understanding of most of the proteins in the module, and the many possible mechanisms by which they influence disease pathology individually or collectively. By developing TEPs for multiple matrisome module targets, we aim to provide resources that can be used to dissect the function of this module and understand the relevance of individual module members to AD.

1.3 | TEPs and open science

One of the anticipated major public benefits of this program is our commitment to share all reagents, including chemical inhibitors,

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FIGURE 3 Screenshot depicting the Target Enabling Resources page within the AD Knowledge Portal, found at https://adknowledgeportal.synapse.org/Explore/Target%20Enabling%20Resources.

without restriction or patent protection. This policy is designed to promote rapid and broad scientific discovery so that target validation can be robustly and rapidly performed across multiple independent research groups. Emerging evidence indicates that scientific discovery is slowed by the cautionary actions required to protect potential future intellectual property, such as imposing restrictive legal agreements on potential collaborators.¹⁷ In further support of this, scientific discovery-as measured by citations-has been shown to be greater for openly available small molecule inhibitors than for analogous inhibitors that are encumbered by restrictions on use.¹⁸ Several avenues exist to advance an unpatented TREAT-AD drug candidate toward a therapy, including through the National Institute on Aging (NIA) drug discovery pipeline. Companies such as M4ND Pharma¹⁹ do so by practicing open science business models. Alternatively, TREAT-AD assets can be used to enable others to invent new patentable chemistry for a target. Our model offers the promise of stimulating new approaches to drug discovery by removing intellectual property restrictions as a barrier to research and openly sharing all tools that we generate.

2 CONCLUSION

The primary aim of this work is to increase the numbers of new targets considered in AD drug discovery pipelines by providing the community with the TEPs and drug-like tools necessary to carry out robust target characterization. The TREAT-AD program is the early translational partner of the NIA Translational Research Program and, so, operates on target priorities identified within this ecosystem. Reagents developed through TREAT-AD are designed to uncover new biology, invalidate some target hypotheses, and increase the scientific attention on lesser-studied human proteins and other targets. This alone will be a tremendous contribution to AD research. If one of the targets proves promising and advances into drug discovery and development, then its impact would extend from basic understanding to translational medicine.

Our center is designed as a community resource. As such, we actively solicit input on experimental prioritization from domain experts and promote the use of center outputs, in particular to test emergent therapeutic hypotheses. To encourage use by others, all available TREAT-AD resources, including TEP reports, are cataloged in the AD Knowledge Portal⁸ (Figure 3). An overview of the full TREAT-AD portfolio, including resources currently in development, can be found on the TREAT-AD website.²⁰

We are committed to generating high-quality probes that can be applied alongside an appropriate suite of reagents to enable discoveries about fundamental biology and validate the roles of these and other potential emerging AD targets. We will continue to make resources available and to collaborate widely with the research community—and we call on the community to use these reagents and engage with us to guide future reagent development.

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CONFLICT OF INTEREST STATEMENT

The authors declare no conflicts of interest. Author disclosures are available in the Supporting Information.

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SUPPORTING INFORMATION

Additional supporting information can be found online in the Supporting Information section at the end of this article.

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