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Targeting the APP-Mint2 Protein–Protein Interaction with a Peptide-Based Inhibitor Reduces Amyloid- β Formation

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ASSOCIATED CONTENT

Supporting Information

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Synthesis and additional experimental and analytical details (PDF).

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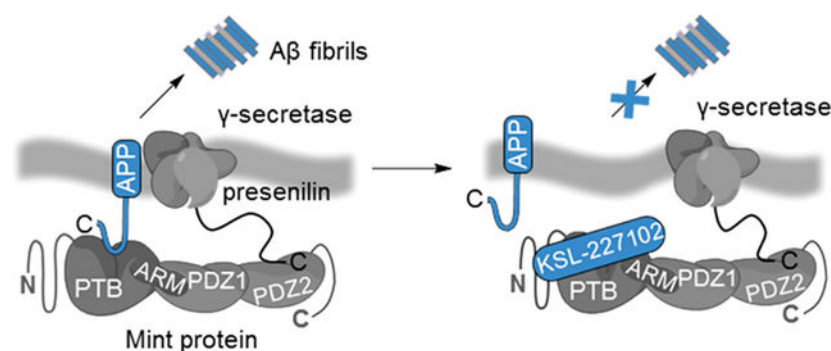
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Abstract

There is an urgent need for novel therapeutic approaches to treat Alzheimer's disease (AD) with the ability to both alleviate the clinical symptoms and halt the progression of the disease. AD is characterized by the accumulation of amyloid- β ($A\beta$) peptides which are generated through the sequential proteolytic cleavage of the amyloid precursor protein (APP). Previous studies reported that Mint2, a neuronal adaptor protein binding both APP and the γ -secretase complex, affects APP processing and formation of pathogenic $A\beta$. However, there have been contradicting results concerning whether Mint2 has a facilitative or suppressive effect on $A\beta$ generation. Herein, we deciphered the APP-Mint2 protein-protein interaction (PPI) via extensive probing of both backbone H-bond and side-chain interactions. We also developed a proteolytically stable, high-affinity peptide targeting the APP-Mint2 interaction. We found that both an APP binding-deficient Mint2 variant and a cell-permeable PPI inhibitor significantly reduced $A\beta_{42}$ levels in a neuronal *in vitro* model of AD. Together, these findings demonstrate a facilitative role of Mint2 in $A\beta$ formation, and the combination of genetic and pharmacological approaches suggests that targeting Mint2 is a promising therapeutic strategy to reduce pathogenic $A\beta$ levels.

Graphical Abstract



INTRODUCTION

Amyloid- β ($A\beta$) peptides, the main component of senile amyloid plaques, play a key role in Alzheimer's disease (AD) pathogenesis.^{1,2} $A\beta$ peptides are generated by the sequential proteolysis of the amyloid precursor protein (APP),³ a type I transmembrane protein of 695–770 amino acids. In the pathogenic amyloid cascade, cleavage by an aspartyl protease (BACE-1 or β -secretase) generates a soluble APP ectodomain (sAPP β) and a membrane-bound C-terminal fragment (C99). Subsequent cleavage of the C99 fragment by γ -secretase generates the APP intracellular domain (AICD) and $A\beta$ fragments of various lengths, including $A\beta_{40}$ and $A\beta_{42}$. The latter exhibits an increased propensity to form insoluble aggregates, which are a hallmark of AD pathology.

The accumulation of $A\beta$ peptides at synaptic sites has detrimental effects on synaptic function, which contributes to the cognitive decline and memory loss associated with AD.⁴ The most advanced anti- $A\beta$ drugs in clinical development are anti- $A\beta$ immunotherapies promoting clearance of the $A\beta$ peptide and its aggregates or β -secretase inhibitors intended to interfere with the initial step of $A\beta$ formation.⁵ However, both strategies have suffered from failure in late-stage clinical trials,^{6,7} and no disease-modifying AD therapy is yet available. Alternative strategies to reduce $A\beta$ formation, such as the modulation of direct binding partners involved in APP trafficking and processing, could therefore increase the chance of developing a successful AD treatment, which is of increasing importance in light of a constantly growing patient population.⁸

Munc18-interacting proteins (Mints, also known as X11s) are a family of adaptor proteins encoded by three distinct genes (*ABPA1*, *ABPA2*, and *ABPA3*) that produce neuron-specific Mint1 (X11 α) and Mint2 (X11 β) and ubiquitously expressed Mint3 (X11 γ).^{9–11} Mint proteins consist of a variable isoform-specific N-terminal region and a conserved C-terminal region that contains a phosphotyrosine binding (PTB) domain and two tandem PSD-95/discs large/zonula occludens (PDZ) domains (Figure 1a).¹² The PTB domain of all three Mint proteins binds the conserved endocytic YENPTY motif of APP, which is essential for regulating APP trafficking and affects $A\beta$ production.^{13–16} A number of studies suggest that the APP-Mint protein-protein interaction (PPI) is of particular biological significance in AD. First, the loss of each individual Mint isoform delays age-dependent $A\beta$ plaque formation in mouse models of AD.¹⁷ Second, Mint proteins interact via their PDZ domains with presenilin-1, the catalytic subunit of the γ -secretase complex, thereby promoting APP/

presenilin-1 colocalization favoring A β formation.^{18–21} Third, Mint proteins are found to be upregulated in AD and are reported to associate with neuritic plaques.^{22,23} Altogether, these findings indicate that the APP-Mint PPI is of therapeutic relevance to A β formation and AD. Efforts have been made to decipher the physiological role of Mint proteins in A β formation, and numerous knockout and knockdown studies, using mainly *in vivo* models, have been reported.^{17,20,24–30} However, these studies revealed both suppressive and facilitative effects of Mint proteins on A β formation. The role of Mint proteins in the formation of pathogenic A β remains controversial since no chemical probe perturbing the APP-Mint PPI has been reported.

Herein, we characterized the APP-Mint2 interaction with molecular resolution through extensive substitutional analyses of both the APP C-terminal peptide ligand (APP_{C-term}) and the Mint2 protein. The introduction of backbone amide-to-ester (A-to-E) modifications uncovered an intricate H-bond network at the APP-Mint2 interface, complementing previously published cocrystal structures. Alanine (Ala) substitutions in the PTB domain of Mint2 identified key residues and led to the generation of an APP binding-deficient Mint2 variant that reduced A β ₄₂ generation *in vitro*. Furthermore, we performed a systematic structure–function analysis of the APP C-terminal peptide which facilitated the development of a proteolytically stable macrocyclic peptide exhibiting nanomolar affinity toward Mint2. In a neuronal *in vitro* model of AD, cell-permeable variants of the high-affinity C-terminal APP mimetic peptide demonstrated that modulation of the APP-Mint2 interaction reduces A β ₄₂ levels.

RESULTS AND DISCUSSION

Minimal APP Mint2-PARM Binding Sequence and Semisynthesis of Mint2-PARM Variants.

Within the Mint protein family, Mint2 exerts the greatest effect on A β formation and was therefore selected as the target protein.¹⁷ The interaction of APP and Mint2 is mediated by the PTB domain of Mint2 and modulated in an autoinhibited open–closed fashion by a C-terminal α -helical linker (termed ARM) adjacent to the PTB domain.^{12,31,32} Consequently, we designed a human Mint2-PTB-ARM (Mint2-PARM) construct (PARMH₆; residues 364–570) as a model protein (Figure 1a). To identify the minimal binding sequence of APP, we systematically tested N- and C-terminal truncated variants of the human 17-mer APP_{C-term} peptide (QNGYENPTYKFFEQMQN; residues 754–770 in APP₇₇₀). Using a fluorescence polarization (FP) assay (SI section Detailed Methods), we identified a 12-residue-long peptide, designated as the APP_{WT} peptide (NGYENPTYKFFE; residues 755–766; $K_d = 4.0 \pm 0.2 \mu\text{M}$), as the minimal Mint2-PARM binding sequence (Figure S1 and Table S1).

The reported X-ray cocrystal structure of Mint2 and the APP peptide (QNGYENPTYKFFEQ; residues 753–767, Figure 1b, blue) suggests that an intricate network of backbone hydrogen bond (H-bond) interactions and side-chain interactions mediates the APP-Mint2 interaction (Figure 1c, dotted lines).^{12,14} Therefore, we probed side-chain interactions by introducing Ala substitutions, whereas backbone H-bond interactions can only be effectively probed by introducing A-to-E substitutions.^{33–37} While we generated Mint2-PARM Ala variants through recombinant expression (Table S2),

introducing A-to-E substitutions into the Mint2-PARM domain required the development of a semisynthesis strategy. Specifically, we first generated a pseudo-wild-type Mint2-PARM variant (pPARM-His6; residues 364–570; R455K, C483A, C501A, C566A) containing three cysteine (Cys) to Ala substitutions enabling the use of native Ala residues in the native chemical ligation (NCL). Next, we synthesized the same version of Mint2-PARM semisynthetically (pPARM_{ss}; residues 364–570; R455K, C483A, C501A, C566A with no His tag) using a three-fragment expressed protein ligation (EPL) approach (Figure 2a). Both protein variants (pPARM-His6 and pPARM_{ss}) displayed similar secondary structure and binding affinity for the 17-mer APP_{C-term} peptide as recombinantly expressed Mint2-PARM (Figures S2 and S3), confirming that neither the His6-tag nor the introduced substitutions affect APP binding. We applied the semisynthetic strategy to introduce eight A-to-E substitutions (R455 κ , T456 τ , I457 ι , S458 σ , Y459 ψ , I460 ι , A461 α , and D462 δ ; α -hydroxy amino acids abbreviated with Greek letters)³⁸ into Mint2-pPARM_{SS} (Tables S3 and S4), which were all obtained with sufficient yields and purities (Figures S4–S7).

Similarly, we introduced A-to-E substitutions in five positions of the APP_{WT} peptide (G756 γ , Y757 ψ , E758 e , N759 ν , and F764 ϕ). The corresponding depsi-peptides were synthesized on the solid phase using α -hydroxy amino acids (SI section Detailed Methods).^{39,40} It is generally accepted that A-to-E substitutions do not perturb the overall protein structure.⁴¹ However, the H-bond donor ability of the respective amide-NH is removed and the acceptor capability of the adjacent carbonyl is reduced, which provides a refined tool for studying H-bond networks. A-to-E mutations have successfully been employed to address the significance of backbone H-bonds for folding β -sheet structures and backbone H-bonds in A β peptides.^{38,42–44}

In total, 21 Mint2-PARM protein variants containing both Ala and A-to-E substitutions and 17 APP_{WT} peptide analogues were generated. We then employed the FP assay to address the consequences of each substitution on the APP-Mint2 interaction (Figure 2b–e).

Backbone H-Bonds and Side-Chain Contacts Are Essential to the APP-Mint2 Interaction.

Single-point Ala substitutions in the APP_{WT} peptide resulted in a >10-fold loss of affinity to Mint2-PARM in 4 out of 12 positions (Figure 2b, Figure S8a,b, and Table S1). The substitution of two residues, Y757A and N759A located in the endocytic YENPTY motif of APP, resulted in 76-fold and >100-fold decreases in affinity, respectively. In addition, Ala substitution of residues F764 and F765 resulted in 14- and 18-fold decreases in affinity, respectively (Figure 2b). The N-terminal region of the APP peptide forms an antiparallel β -sheet with the β 5 strand of Mint2-PARM. On the basis of the X-ray cocrystal structure, H-bond interactions that formed between the APP peptide and Mint2 were probed by A-to-E substitutions (G756 γ , Y757 ψ , E758 e , and N759 ν) (Figure 1c). All four A-to-E substitutions resulted in a reduced binding affinity relative to the APP_{WT} peptide (Figure 2c, Figure S8c,d, and Table S1). Interestingly, the Y757 ψ substitution exhibited a >100-fold loss of affinity to Mint2-PARM, as did the Y757A substitution, rendering Y757 as a key residue in APP.

In Mint2-PARM, three Ala substitutions (L454A, Y459A, and F520A) exhibited a >10-fold reduction in the APP_{C-term} affinity (Figure 2d, Figure S9, and Table S2). The robust effect of

Y459A substitution in Mint2-PARM is likely a result of disrupting the hydrophobic interaction with the C α atom of G756 in APP, an interaction also reported for Mint1 (Y418).¹⁴ Notably, F520A substitution in Mint2-PARM led to a 45-fold reduction in binding affinity, plausibly explained by contacts with key residue N759 in APP (Figure S10a). Corresponding residues in the PTB-domains of Mint1 (F608) and p52 Shc (F198) are also reported to be important for peptide recognition, suggesting that the conserved Phe constitutes an important mediator of ligand recognition for PTB domains.^{13,45} Next, we evaluated the effect of A-to-E substitutions in Mint2-PARM on the binding to the APP_{C-term} peptide. In general, the introduced substitutions reduced the binding affinity of the APP_{C-term} peptide by 2- to 4-fold (Figure 2e and Table S4). However, two A-to-E substitutions, S458 σ and D462 δ , resulted in 14- and 25-fold reduced affinities, respectively. Both residues engage in H-bonds and form the antiparallel β -sheet with the N-terminal region of the APP_{C-term} peptide (Figure 1b,c). Altogether, we identified two key side chains (Y757 and N759) and one backbone amide (Y757 ψ) in APP to be critical to the APP-Mint2 interaction. In the Mint2-PTB domain, we found F520A, Y459A, and one backbone amide substitution (D462 δ) led to a substantial reduction in affinity to the APP_{C-term} peptide (Figure 2d, e).

Compared to the reported cocrystal structure of APP and Mint2, we observed a number of deviations from the suggested interactions. First, the side chains of D462 in Mint2 and Y757 in APP are suggested to interact via an H-bond. In our FP measurements, we detected no effect of the D462A mutation in Mint2-PARM on the binding of the APP_{C-term} peptide (Figure 2d); however, we cannot exclude that the Ala mutation can be compensated for by a water molecule mimicking the H-bond acceptor of D462. Second, Q767 of APP is proposed to H-bond to N387 in Mint2, although the truncated APP_{WT} peptide lacking Q767 retains a wild-type binding affinity (Figure S1). Finally, the side chain of the critical N759 residue in APP is proposed to bind the backbone carbonyls of both L454 and I457 in Mint2.¹⁴ Here, our A-to-E substitutions suggest that only the H-bond protruding from I457 of Mint2 is important, while perturbing the carbonyl of L454 had only minor effects on the affinity to the APP_{C-term} peptide (Figure 1c).

Mutations in the Mint2-PTB Domain Reduce APP Binding and A β Formation.

We took advantage of the Mint2-PARM Ala-scan results and designed a Mint2 variant with reduced APP binding affinity. Introducing two mutations, Y459A and F520A, into human Mint2-PARM (hMint2^{Y459A/F520A}) resulted in a significant 72-fold decrease in the APP_{C-term} peptide binding affinity ($K_d = 895 \pm 43 \mu\text{M}$) in the FP assay (Figure 3a and Figure S10). To examine whether hMint2^{Y459A/F520A} has a cellular effect on APP binding, we produced a full-length GFP-tagged rat Mint2 (GFP-rMint2^{WT}) and corresponding APP binding-deficient variant GFP-rMint2^{Y460A/F521A} (analogous to hMint2^{Y459A/F520A}, Figure S11). We confirmed our FP results by cotransfecting HEK293T with full-length APP and either GFP-rMint2^{WT} or GFP-rMint2^{Y460A/F521A} and performing coimmunoprecipitation assays. APP coimmunoprecipitated with both GFP-rMint2^{WT} (lane 5, Figure 3b) and GFP-rMint2^{Y460A/F521A} (lane 6, Figure 3b); however, GFP-rMint2^{Y460A/F521A} exhibited a significant 85% reduction in APP binding (Figure 3c). Next, we examined the functional effect of an impaired interaction between APP and Mint2 on A β generation. We employed

primary cortical neurons from an established murine AD model with a double transgene for the expression of mutant APP (APP^{swe}) and presenilin-1 proteins (PS1^{E9}), resulting in enhanced A β production.¹⁷ We infected primary neurons with a full-length GFP-rMint2^{Y460A/F521A} lentivirus after 2 days *in vitro* (2 DIV). Expression of the GFP-rMint2^{Y460A/F521A} variant was confirmed by Western blot analysis (Figure 3d). We quantified the effect on A β ₄₂ levels released from primary neurons using an enzyme-like immunosorbent assay (ELISA) on the neuronal conditioned medium. At 15 DIV, we observed a significant 42% decrease in A β ₄₂ levels in neurons infected with GFP-rMint2^{Y460A/F521A} relative to neurons expressing only endogenous Mint2 (Figure 3e). Together, our results confirm that APP interacts with the PTB domain of Mint2 and suggest that Mint2 plays a facilitative role in A β formation in our neuronal AD model.

Non-canonical Amino Acids Enhance the Binding of the APP Peptide to Mint2.

Since the APP binding-deficient rMint2^{Y460A/F521A} variant reduced A β production *in vitro* and previous studies revealed that Mint2 knockout in APP^{swe}/PS1^{E9} transgenic mice reduces A β formation *in vivo*, we hypothesized that a peptide-based inhibitor designed to disrupt the APP-Mint2 interaction could reduce A β generation. As such, we used the APP_{WT} peptide as a template for an initial D- and *N*-Me-amino acid (d-AA and *N*Me-AA) scan (Figure S8 and Table S1). D-AAs were tolerated only in the N- and C-termini of the APP_{WT} peptide, whereas the *N*-methylation of N755 resulted in an improved affinity toward Mint2-PARM. Notably, the substitution of Y757 with *N*-MeY757 resulted in a nonbinding APP_{WT} peptide analogue, which is in agreement with the reduced binding of the Y757 ψ analogue (Figure 2c). Next, we prepared a series of APP_{WT} peptide variants containing single substitutions of both canonical (cAAs) and noncanonical amino acids (ncAAs) (Figure 4a). First, we tested position N759, the most critical residue according to the Ala scan, and introduced eight different amino acids. Remarkably, all substitutions, even conservative ones such as N759Q, abrogated the binding to Mint2-PARM ($K_i = 500 \mu\text{M}$) (Figure 4b). We therefore consider N759 to be a key residue in the APP-Mint2 interaction, which is further supported by its conservation as part of the APP endocytic NPXY consensus sequence that interacts with the PTB domain of Mint proteins.⁴⁶ Next, we probed the two aromatic residues, F764 and F765. Individual substitution by Trp had no effect on the affinity. In contrast, the incorporation of L-3-(1-naphthyl)alanine (Nal1) at either position improved the affinity for both variants, and the corresponding double substitution showed a 4-fold improvement in affinity ($K_i = 1.1 \pm 0.3 \mu\text{M}$). We attribute the improved affinity to hydrophobic interactions with Y524 and F527 in Mint2-PTB located in close proximity to F764 and F765. Neither L-3-(2-naphthyl)alanine (Nal2) nor 2-indanyl-L-glycine (Igl) improved the affinity to the same extent (Figure 4b).

Finally, the substitution of Y762 to Phe resulted in a 4-fold increase in affinity ($K_i = 1.0 \pm 0.1 \mu\text{M}$), while an Y762W substitution lowered the affinity to Mint2-PARM (Figure 4b and Table S1).

Taken together, we identified six positions (N755, Y757, Y762, F764, F765, and E766) in the APP_{WT} peptide at which substitutions to cAAs or ncAAs either improved the affinity or were expected to improve the proteolytic stability of the peptide.

Side Chain-to-Side Chain Cyclization in the APP_{WT} Peptide Improves Affinity and Stability.

To improve the binding and metabolic stability of our peptide ligand, we introduced side chain-to-side chain lactam cyclizations employing solid-phase peptide synthesis (SPPS) and on-resin cyclization (Figure S12a). According to the reported X-ray cocrystal structure of APP-Mint2, native residues E758 and K763 are in close proximity when forming an intramolecular noncovalent salt bridge (Figure S12b).¹² We reasoned that introducing a lactam bridge in this position could confer conformational constraint and potentially improve the affinity and stability.⁴⁷ Indeed, a cyclic analogue of the APP_{WT} peptide containing a lactam linkage between E758 and K763 (*i* to *i* + 5, cAPP_{E758-K763}; **1**) exhibited improved affinity compared to the APP_{WT} peptide and was also superior to a variant cyclized between Y762K and E766 (*i* to *i* + 4, cAPP_{Y762-E766}; **2**) (Figure S12b-d). Next, we prepared a series of cyclic variants of **1** in which we systematically modified the lactam bridge by introducing Asp, Orn, or Dap and reversed the orientation of the amide bond in the lactam bridge (Figure 4c,d). The affinity of these 11 peptides (**3-13**) was highly dependent on the nature of the introduced macrocyclization (Figure 4e).

In general, macrocyclic peptides with carboxylic acid side chains in *i* positions (**1** and **3-7**; Figure 4c) were characterized by reduced binding affinities, while peptides with amino side chains in *i* positions (**8-13**, Figure 4d) displayed improved affinities relative to the linear APP_{WT} peptide. Furthermore, smaller ring sizes (**5-7**, **11**, and **13**) resulted in reduced binding affinities with only peptide **12** ($n = 3$, $m = 1$) refuting this trend. Among the 12 macrocycles, peptide **10** ($n = 4$, $m = 1$) exhibited the highest affinity ($K_i = 2.2 \pm 0.3 \mu\text{M}$) (Figure 4e). To test whether macrocyclization improved the proteolytic stability, cyclic peptide **1** was tested in an *in vitro* human plasma stability assay, resulting in a 4-fold-improved half-life time ($T_{1/2}$) over the APP_{WT} peptide, $T_{1/2} = 105$ and 26 min, respectively (Figure S12e).⁴⁸

On one hand, the moderately improved affinity might be a result of the introduced constraint causing a preorganization of the peptide resembling the bound conformation with a well-defined helical turn of residues Y762-E766, which reduces the entropic penalty upon binding. On the other hand, additional favorable interactions by the formed lactam bridge can also account for the improved affinity. We performed molecular dynamics (MD) simulations of cyclized peptides **1** and **3-12** and found a correlation between the degree of helicity in solution and the observed binding affinities. For the two most potent cyclic peptides **10** and **12**, helical propensity predictions of 42 and 47%, respectively, were observed. In contrast, for low-affinity peptides **11** and **13** the molecular dynamics simulations suggested helical propensities of 6% and less (Figure 4f). This suggests that ligand preorganization is the driving force of the improved binding. Altogether, the systematic lactam macrocyclization scan led to the identification of a cyclic APP_{WT} peptide scaffold with improved binding affinity and proteolytic stability.

Identification of a Metabolically Stable, High-Affinity Mint2 Inhibitor Derived from the APP_{WT} Peptide.

To develop a potent and proteolytically stable modulator of the APP-Mint2 PPI, we selected cyclic peptide **10** ($n = 4$, $m = 1$) as a scaffold. Next, we introduced D-*N*-Me-Ala and D-Glu at

positions 755 and 766, respectively. Finally, ncAA L-Nal1 was selected for positions 764 and 765. The corresponding peptide, KSL-221036 (*N*Me-aGY *cyclo*-[KNPTYD]Nal1Nal1e-OH, **14**), was generated through SPPS including on-resin cyclization via orthogonally protected Lys(Aloc) and Asp(All) (Figure 5a). We found that **14** binds Mint2-PARM with nanomolar affinity ($K_d = 53 \pm 2$ nM) as determined by isothermal titration calorimetry (ITC). Compared to the APP_{WT} peptide ($K_d = 2.4$ μ M), a 46-fold improved affinity was achieved (Figure 5b and Figure S13). Furthermore, **14** was found to be proteolytically stable over 24 h (half-life = 1440 min) in human plasma (Figure 5c), while the APP_{WT} peptide exhibited a half-life of <30 min. Finally, we examined the rate of metabolism through cytochrome P450 enzymes using mouse liver microsomes and determined the *in vitro* hepatic clearance of the APP_{WT} peptide and **14**. We found the APP_{WT} peptide was cleared fast [$CL_{(int)} = 15.4 \pm 2.5$ μ L/(mg min)] while **14** was not metabolized over 60 min [$CL_{(int)} = 0.0 \pm 0.8$ μ L/(mg min)] (Figure 5d). This data demonstrates the design of a peptide using the minimal binding sequence of the endogenous APP peptide–ligand as a template. Macrocyclization and the incorporation of selected ncAAs resulted in the design of KSL-221036 (**14**), the first reported modulator of the APP-Mint2 PPI exhibiting nanomolar affinity and suitable stability characteristics for further studies.

KSL-221036 (**14**) Interacts with Mint2 and Reduces A β Levels.

To determine whether KSL-221036 (**14**) binds full-length Mint2 from primary neuronal lysate, we loaded the APP_{WT} peptide and **14** onto epoxy-coated magnetic beads via an N-terminal Cys-PEG linker (Table S1). Pull-down from the lysate of primary mouse neurons confirmed that **14** efficiently interacts with endogenous Mint2 at higher affinity (lane 2, Figure 5e) than for the APP_{WT} peptide under the same conditions (lane 3, Figure 5e; see also Figure S14).

To examine the effect of **14** on A β_{42} production in neurons, we attached cell-penetrating peptide (CPP) TAT^{47–57} (YGR-KKRRQRRR)⁴⁹ to the N-terminus of **14** (TAT-**14**) and the APP_{WT} peptide (**15**, Table S1). TAT is a widely used CPP and was recently demonstrated to be compatible with late-stage clinical development in two phase III clinical trials (ESCAPE⁵⁰ and FRONTIER), rendering TAT the most advanced CPP tag in the clinic. We first treated neurons with a concentration curve of peptide TAT-**14** to determine the highest concentration at which no adverse effect on cell viability was observed. Using primary neurons and a colorimetric readout of metabolic activity, peptides **14** (no CPP) and **15** (TAT-APP_{WT}) exhibited no adverse effect while peptide TAT-**14** affected cell viability at 20 μ M (75%) but had no measurable effect at 10 μ M (Figure S15). To determine whether modulation of the APP-Mint2 interaction affects A β production, primary cortical neurons (14 DIV) carrying a double transgene for the expression of mutant APP (APP^{swe}) and presenilin-1 proteins (PS1^{E9}) were incubated for 24 h with peptides **14**, TAT-**14**, and **15**. ELISA quantification of A β_{42} from the conditioned medium revealed that TAT-**14** (10 μ M) led to a 44% reduction of A β_{42} levels relative to vehicle control. In contrast, TAT-APP_{WT} (**15**, 50 μ M) and **14** (10 μ M, no CPP) did not affect A β_{42} levels. High-affinity γ -secretase inhibitor DAPT (*N*-[*N*-(3,5-difluorophenacetyl)-l-alanyl]-s-phenyl-glycine *tert*-butyl ester or LY-374973) served as a positive control, reducing A β_{42} levels by 47% (Figure 5f and Figure S16).⁵¹ In addition, we assessed the effect of TAT-**14** on cellular integrity by quantifying

lactate dehydrogenase (LDH) release (a proxy for cytotoxicity) in the same conditioned medium used for the aforementioned ELISA. We found low LDH levels for both DAPT- and TAT-**14**-treated neurons relative to vehicle control, indicating that the effect of peptide TAT-**14** on A β production is not due to cellular toxicity (Figure 5g; see also Figure S16). Furthermore, p-value analysis found no significant difference between the vehicle and TAT-**14** ($p = 0.2448$) or DAPT and TAT-**14** ($p = 0.6525$) LDH release. To further support the TAT-mediated intracellular delivery of **14**, we detected TAT immunostaining in primary neurons incubated with TAT-**14** (Figure S17), indicating that TAT-**14** is cell-permeable.

Following the promising A β_{42} results of TAT-**14**, we investigated the dose–response relationship and found a weak correlation between the concentration of TAT-**14** and reduced A β_{42} levels (Figure S16e). We reasoned that the delivery efficiency of the cargo potentially depends on the employed CPP tag; therefore, we synthesized four additional variants of **14** containing different CPPs attached to the N-terminus: mixTAT-**14** (rRrGrKkrK-**14**), polyArg-**14** (RRRRRRRRR-**14**),⁵² D-SynB3-**14** (frrsyslrr-**14**),⁵³ and MiniAp4-**14** (cyclo(DLATEPAK(Dap)-**14**)).⁵⁴ Next, we determined the highest concentration at which no adverse effect on cell viability was observed. Peptide MiniAp4-**14** exhibited no toxicity, while peptides mixTAT-**14** and polyArg-**14** were tolerated at concentrations of 5 and 7.5 μM , respectively (Figure S15). Peptide D-SynB3-**14** was poorly soluble and not further tested. The peptides' effect on A β formation was evaluated analogously to that of peptide TAT-**14**. The quantification of A β_{42} from the conditioned medium of primary neurons revealed that incubation with mixTAT-**14** (5 μM) and polyArg-**14** (5 μM) reduced A β_{42} levels by 80 and 84%, respectively. In contrast, incubation with MiniAp4-**14** (10 μM) did not affect A β_{42} production (Figure S16c). Of note, we detected a small increase in LDH activity following incubation with mixTAT-**14** (5 μM) compared to the vehicle control (Figure S16d). Consequently, polyArg-**14**, which reduced A β_{42} production at a lower concentration (5 μM) with minimal cellular toxicity, is a promising CPP alternative to TAT-**14**.

Our results indicate that peptide **14** successfully binds Mint2 *in vitro* with greater affinity than the APP_{WT} peptide. Further studies in primary neurons cultured from our AD mouse model revealed that TAT-**14** significantly decreases A β_{42} production with minimal toxicity. Further investigation into alternative CPPs showed that polyArg also provides attractive properties that facilitate the cellular uptake of peptide ligand **14**. The observation that TAT-APP_{WT} (15) has no effect on A β_{42} formation is in excellent agreement with the pull-down results and highlights the importance of ligand optimization. According to the reported effects of Mint knockout on APP trafficking and A β_{42} formation,²¹ inhibition of the Mint2-APP PPI by compounds TAT-**14** and polyArg-**14** could either result in reduced APP endocytosis or prevent the formation of the tertiary protein complex comprising APP (C99), Mint2, and the γ -secretase. Both avenues ultimately result in reduced A β formation. As such, the lowered A β_{42} levels seen in the *in vitro* AD model following treatment with compounds TAT-**14** and polyArg-**14** provide evidence that modulation of the APP-Mint2 PPI might serve as a pharmacological strategy to reduce pathologic A β levels in connection with AD.

CONCLUSIONS

In the pursuit of novel strategies to treat AD, pharmacological approaches modulating direct binding partners involved in APP trafficking and processing are promising alternatives to currently pursued therapies. Here, the APP-Mint PPI is an attractive target as the effects of Mint deletions are relatively mild, and our strategy, which selectively inhibits APP binding to the PTB domain of Mint proteins, would likely not interfere with the critical function of Mint proteins in synaptic vesicle exocytosis,¹⁷ which is mediated through Mint's other PPI domains.⁹ Although numerous studies have examined the biological importance of Mint proteins, particularly with regard to A β formation in the context of AD,³⁰ these studies have been informative but contradictory.⁵⁵

Here, we extensively characterized the APP-Mint2 interaction at the molecular level by performing comprehensive mutational scans of the APP-Mint2 interface, covering both side-chain and backbone interactions. Compared to the reported structures, we observed distinct differences and showed that two backbone alterations, Y757 ψ in the APP_{WT} peptide and D462 δ in Mint2, had major impacts on the APP-Mint2 interaction. The introduction of Ala substitutions demonstrated that the side chains of two residues in APP, N759, and Y757 and one residue in Mint2, F520, are pivotal to the APP-Mint2 interaction. These observations led to the generation of a Mint2 variant containing two distinct substitutions, Y459A/F520A, which resulted in impaired APP binding. We showed that reduced APP binding results in significantly reduced A β production, which supports a facilitative role of Mint2 in A β formation and suggests that targeting the APP-Mint2 interaction with a PPI inhibitor is of potential therapeutic relevance.

Using the APP_{WT} peptide as a template, we developed a high-affinity, proteolytically stable cyclic peptide **14**. We demonstrated that introducing noncanonical amino acids, N-methylation, and D-amino acids into a macrocyclic peptide scaffold provided a vast improvement in affinity for Mint2 and stability relative to the APP_{WT} peptide. Notably, compared to APP_{WT}, peptide **14** exhibited an enhanced affinity to robustly pull down full-length Mint2 from neuronal lysate. We confirmed that targeting the Mint2-APP interaction with cell-permeable variants of **14** (TAT-**14** and polyArg-**14**) significantly reduced A β ₄₂ formation, whereas parent APP_{WT} peptide **15** had no effect on A β ₄₂ formation. Peptides TAT-**14**, mixTAT-**14**, and polyArg-**14** therefore provide first proof-of-concept evidence that targeting a direct interaction partner of APP affects its metabolism and A β formation. Furthermore, the comparison of different CPP tags indicates that the selection is critical and affects not only efficacy but also neuronal viability. Here, the combined analysis of LDH levels and A β ₄₂ reduction suggests that polyArg-**14** might be preferred over TAT-**14** (Figure S16).

Previous work has shown that APP and BACE-1 converge in acidic microdomains.^{56,57} Because the BACE-1 cleavage of APP is the rate-limiting step in A β production,⁵⁸ this convergence in endosomes is a critical initiator of the APP amyloidogenic cascade. Sullivan et al. reported that the knockout of Mint proteins resulted in reduced APP endocytosis and A β formation.²¹ We suspect that the APP binding-deficient rMint2^{Y460A/F521A} variant and TAT-, mixTAT-, and polyArg-**14** could have a similar effect. The same study reported that

the knockout of Mint proteins also affects the internalization of presenilin-1 of the γ -secretase complex and reduced colocalization with APP.

The impaired binding of APP (C99 fragment) to Mint2 and presenilin-1 would reduce the probability of γ -site cleavage in APP and hence $A\beta$ formation. Our results encompass both observations.

In summary, we demonstrated that targeting the APP-Mint PPI may present an alternative strategy in the pursuit of new therapeutic approaches in AD treatment. We believe that the compounds reported herein will be of help in identifying mechanisms of Mint2-mediated $A\beta$ formation and enable future *in vivo* studies.

METHODS

Detailed experimental procedures are provided in the Supporting Information.

Peptide Synthesis.

Peptides and depsipeptides were assembled in a manual or automated manner (LibertyBlue, CEM, Matthews, NC, USA) on a solid support using Boc- or Fmoc-based chemistry, purified by RP-HPLC, and characterized by LCMS and UPLC. When relevant, side-chain cyclization was performed on resin by the deprotection of Alloc and Allyl protecting groups, followed by PyBOP-induced cyclization.

Protein Expression.

Human Mint2 constructs were cloned in pRSET, pET, or pTXB1 vectors, expressed in *E. coli* B121(DE3)pLysS (Invitrogen), and purified using HisTrap columns, followed by size-exclusion chromatography on a HiLoad 16/600 Superdex 75 pg column or purification by reverse-phase chromatography (C4 column, YMC-Pack-C4, 250 × 20 mm²).

Expressed Protein Ligation.

The three-step ligation was initiated by ligating PARM_N to the thioester peptide (PARM_{PeP}) fragment. Following buffer exchange, the second ligation was performed using the PARM_C fragment. Obtained proteins were purified using a reverse-phase C4 column (Jupitor, Phenomenex, 250 × 10 mm²). After desulfurization, the semisynthetic protein was refolded into storage buffer and characterized by LCMS and UPLC.

Circular Dichroism (CD).

Experiments were performed on an Olis DSM 100 (Olis Inc.) at 15 μ M protein concentration, and the obtained millidegrees of ellipticity (mo) was converted to the mean residue ellipticity (θ_{MRE}).

ThermoFluor.

ThermoFluor experiments were performed in a 96-well plate format on a Stratagene Mx3005p qPCR cycler (Agilent) using SYPRO orange as the dye at 15 μ M protein concentration.

Fluorescent Polarization (FP).

Saturation binding (for semisynthetic and mutated proteins) using 50 nM TAMRA-APP_{C-term} [(TAMRA)-NNG-QNGYENPTYKFFEQMQN] as a probe was performed using a Safire2 plate reader (Tecan) and increasing concentrations of Mint2-PARM. Inhibition experiments were performed using 50 nM TAMRA-APP_{C-term} and 15 μ M Mint2-PARM at increasing concentrations of peptide ligands.

Rat Mint2^{Y460A/F521A} Cloning.

Using a QuikChange II Site-Directed Mutagenesis Kit (Agilent Technologies), pEGFP-rMint2^{Y460A/F521A} was created from rat pEGFP-Mint2 and subcloned into the lentiviral pFUW vector for neuronal expression.

HEK293T Coimmunoprecipitation.

Transfected HEK293T cells were lysed, and protein extracts were incubated with precipitating antibody and protein A Ultralink resin (Thermo Scientific). Precipitated proteins were resolved by SDS-PAGE and immunoblotted for APP, GFP, and α -tubulin.

Primary Neuron Infection with rMint2^{Y460A/F521A} and A β ELISA.

Primary neuronal cultures were prepared from newborn mice and infected with lentivirus for rMint2^{Y460A/F521A}. At 15 DIV, neuronal lysate was collected, resolved by SDS-PAGE, and immunoblotted for APP, Mint2, and α -tubulin. Conditioned media were collected and handled according to a Human A β ₄₂ Ultrasensitive ELISA Kit (Invitrogen).

Molecular Dynamics Simulations.

The Desmond package in Schrodinger's Maestro suite was used to perform molecular dynamics simulations. Each simulation was run in duplicate (200 ns) under standard temperature and pressure. The simulation output was parsed frame by frame using Python, and helical propensity values were determined.

In vitro Plasma Stability.

Peptides were incubated in human plasma at 37 °C for up to 24 h. The peptides were extracted from the plasma at various time points and analyzed using UPLC.

In vitro Hepatic Clearance.

Peptides were incubated in mouse hepatic microsomes supplemented with reduced β -nicotinamide adenine dinucleotide 2'-phosphate (NADPH) and MgCl₂ at 37 °C for up to 60 min. The peptides were extracted from the microsomes at various time points and analyzed using UPLC.

Isothermal Titration Calorimetry (ITC).

Experiments were performed using an ITC200 (Malvern). Ligand-to-buffer titrations were performed to subtract the heat produced by injection, mixing, and dilution. The binding enthalpy was directly measured, and the dissociation constant (K_d) and stoichiometry (N) were obtained by data analysis using Origin software (OriginLab).

Neuronal Cell Viability.

Primary neuron cultures were treated with peptides or DMSO as indicated. The cell viability and cytotoxicity were determined using a CellTiter 96 AQueousOne Solution Cell Proliferation Assay (MTS) Kit (Promega) and a CyQUANT LDH Cytotoxicity Assay (Thermo Fisher).

Immobilization of Peptides.

Peptide KSL-221036 and the APP_{WT} peptide comprising a C-terminal Cys and a PEG2 linker were loaded onto Dynabeads M-270 Epoxy Beads (Thermo Fischer Scientific) according to the protocol of the manufacturer.

Protein Isolation from Neuronal Cell Lysate.

Primary neuronal cultures were prepared as described above. Neurons expressing Mint2 were infected with lentiviral inactive cre recombinase, while Mint2 knockout neurons were infected with active cre recombinase. At DIV15, neuronal lysate was collected and incubated with peptide-loaded Dynabeads. Isolated proteins were resolved by SDS-PAGE and immunoblotted for Mint2 and GAPDH.

ELISA A β Assay Following Peptide Treatment.

Primary neuronal cultures were prepared from newborn transgenic mice carrying the APP^{swe}/PS1 E9 transgene. At 15 DIV, neurons were incubated with peptides, DAPT (γ -secretase inhibitor), and vehicle control 0.25% DMSO. The collected conditioned media were handled according to the protocol of the Human A β ₄₂ Ultrasensitive ELISA Kit (Invitrogen).

Data Analysis.

All data analysis was performed in GraphPad Prism 7.0 (GraphPad Inc.), Excel 2010 (Microsoft), and Origin 9.0 (OriginLab). Significance was evaluated using the Student's *t* test in the case of two groups. The one-way ANOVA test was used in the case of three or more groups, combined with either Dunnett's or Sidak's multiple comparison test.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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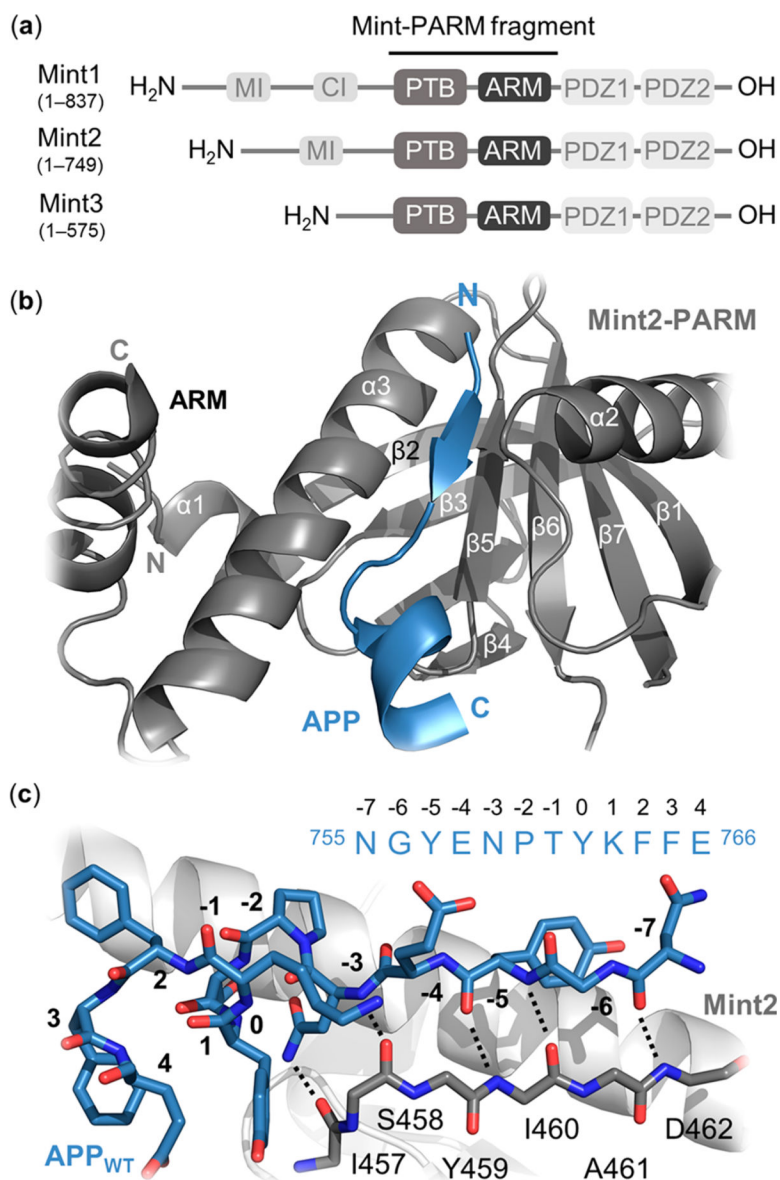


Figure 1. Structural overview of the Mint protein family and the Mint2-APP interaction. (a) Schematic architecture of Mint proteins illustrating their conserved C-terminal region consisting of a phosphotyrosine binding (PTB) domain (gray), an α -helical ARM linker (dark gray), two PSD-95/drosophila discs large/zonula occludens (ZO-1) (PDZ) domains (light gray), and a variable N-terminal region. Numbering corresponds to human residues. MI = Munc-18 interaction domain, CI = CASK interaction domain. (b) Cartoon representation of the interaction between rat Mint2-PARM (gray) and APP (residues 754–767; blue). The APP_{C-term} is binding at the interface between the α 3-helix and β 5-strand of Mint2, forming an antiparallel β -sheet (N-terminal) followed by a β -turn and a C-terminal α -helix. The ARM linker (dark gray) is found in an open conformation enabling APP binding. (c) Structure of the APP peptide (blue sticks) bound to the rat Mint2-PTB domain (gray). The β 5-strand of the PTB domain (gray stick, side chains not depicted) is shown to highlight the

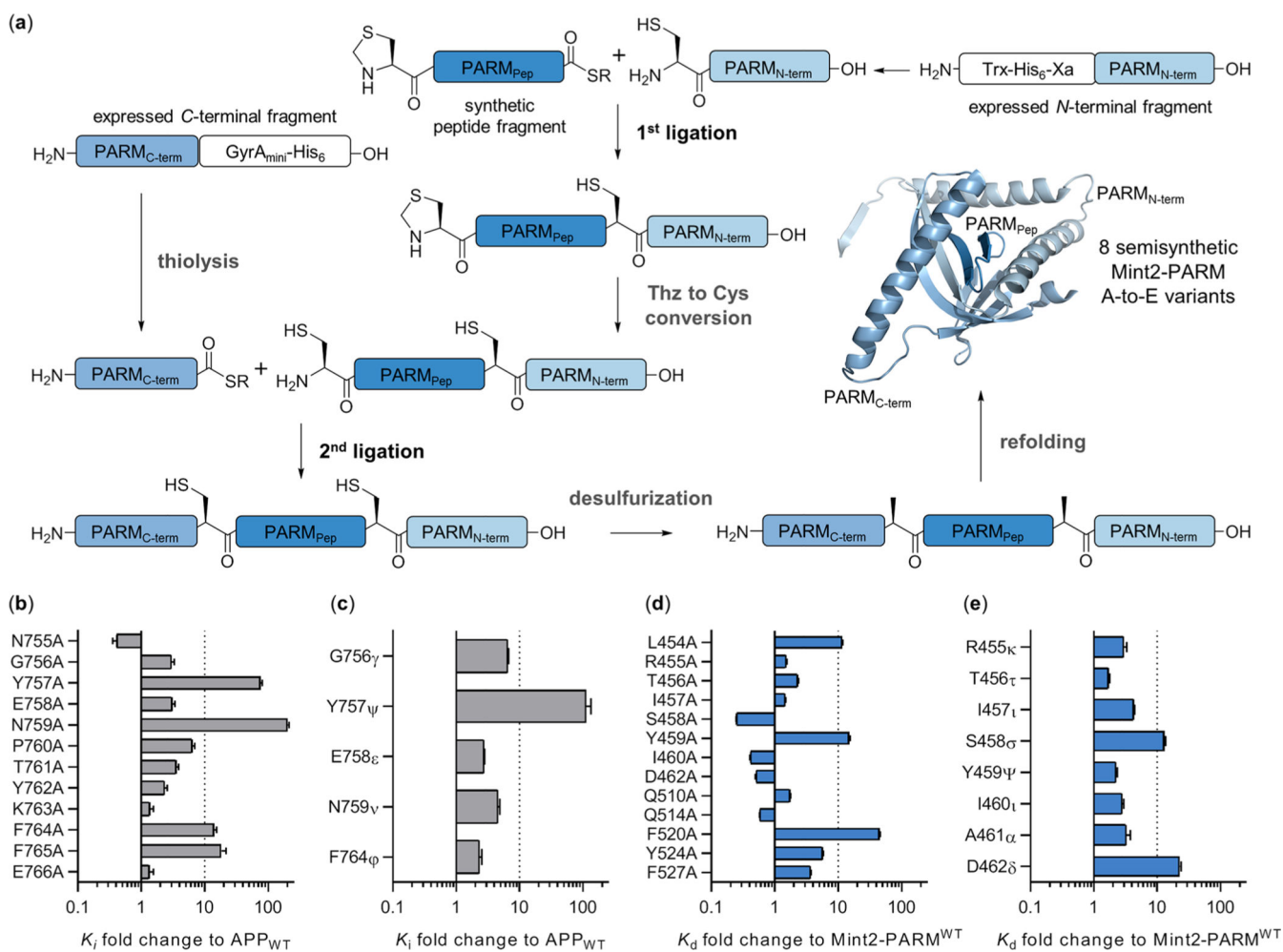
backbone–backbone H-bond network (black dotted lines) between the Mint2-PTB domain and APP (PDB ID: 3SV1).¹²

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**Figure 2.**

Semisynthesis of Mint2-PARM and effects of A-to-E and Ala mutations in APP and Mint2-PARM. (a) Semisynthetic approach to introducing A-to-E substitutions in Mint2-PARM. N-terminal fragment PARM_N (A479-Q570) is fused to a Xa site to generate an N-terminal Cys by factor Xa cleavage. C-terminal fragment PARM_C (E364-H452) is expressed with a C-terminal intein to generate the required C-terminal thioester. Semisynthesis is initiated by ligating synthetic peptide fragment PARM_{pep} to PARM_N. Next, the Thz group is converted to a free Cys and PARM_{pep+N} is ligated to PARM_C. Semisynthetic Mint2-pPARM_{SS} is obtained after desulfurization and refolding (Figures S2–S7). (b) Affinity fold-change of the APP_{WT} peptide's Ala-scan and (c) A-to-E substitutions toward Mint2-PARM obtained in an inhibition FP assay (K_i) reported relative to the APP_{WT} peptide [$K_i(\text{mutant})/K_i(\text{APP}_{\text{WT}}) + \text{SEM}$]. (d) Affinity fold change of the APP_{C-term} peptide binding to Mint2-PARM Ala and (e) A-to-E variants obtained by the saturation FP assay (K_d) relative to Mint2-PARM^{WT} [$K_d(\text{variant})/K_d(\text{wild type}) + \text{SEM}$]. See Figures S8 and S9 for FP data.

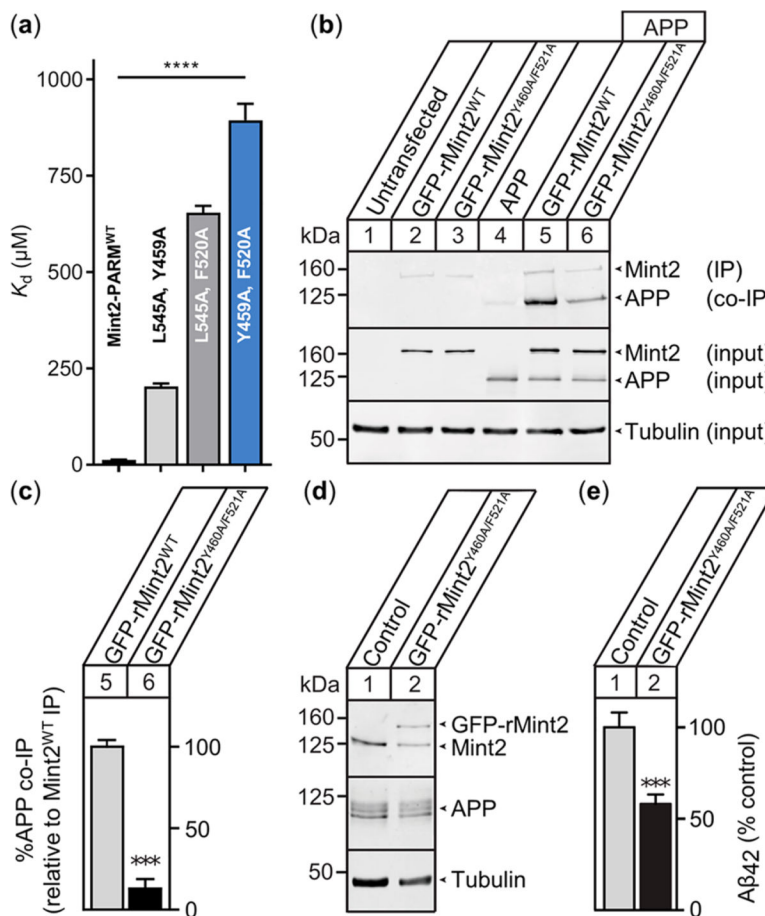
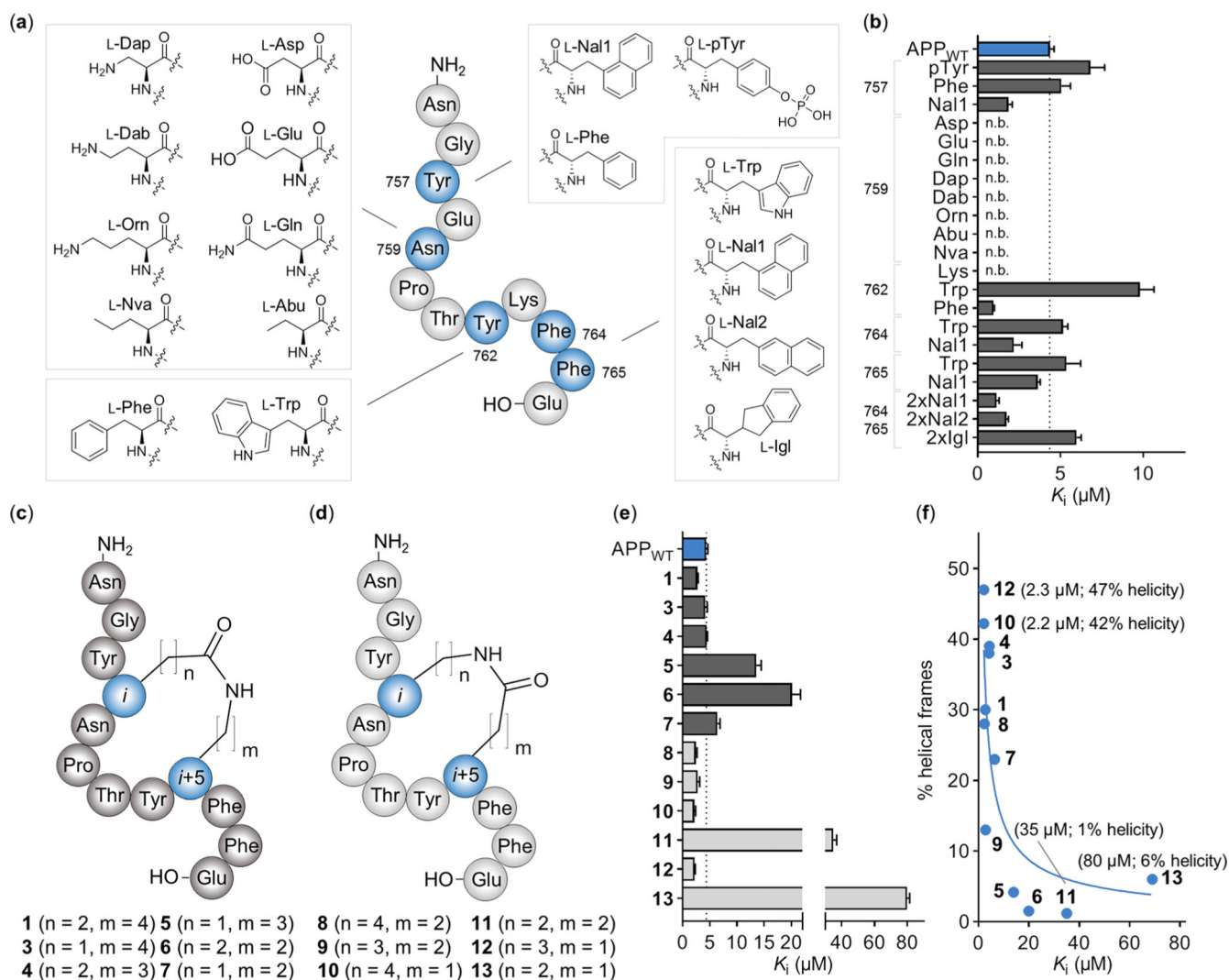
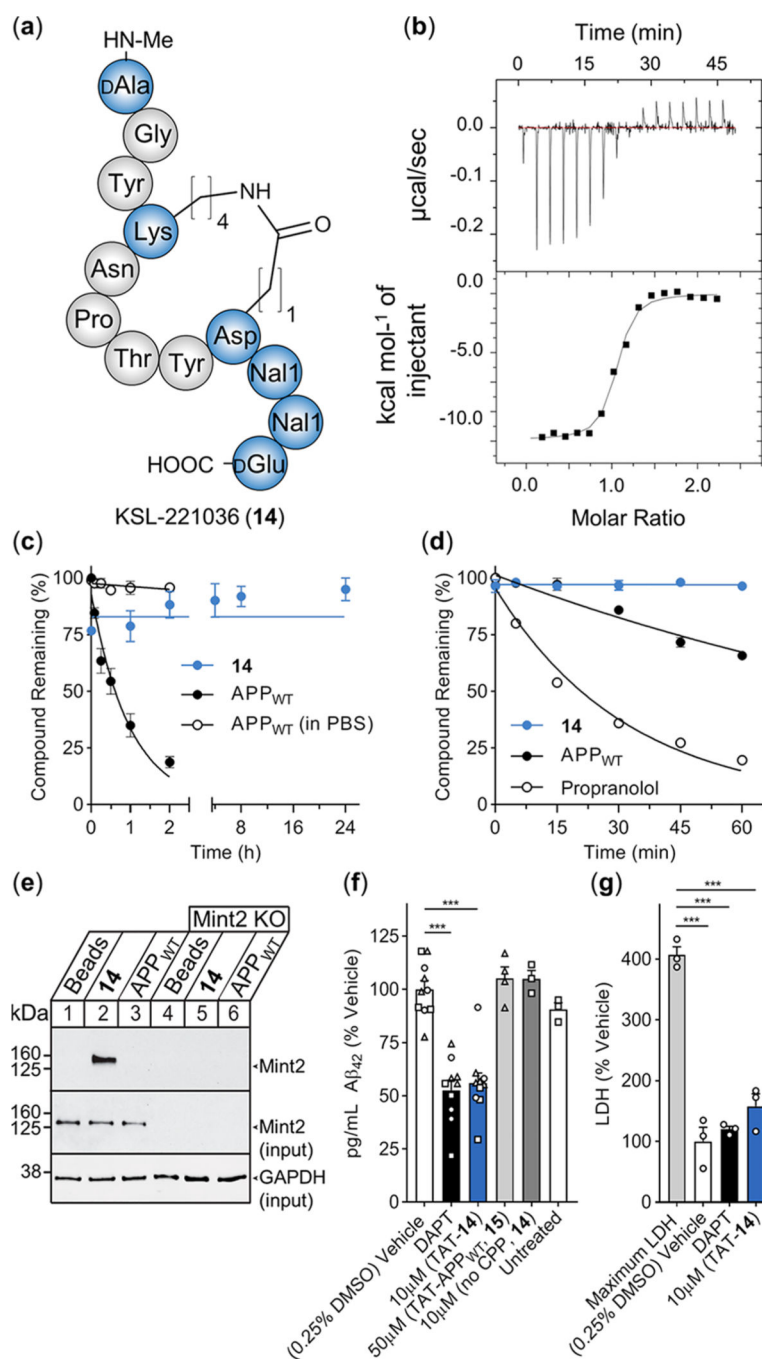


Figure 3. APP binding-deficient rMint2^{Y460A/F521A} variant reduces A β ₄₂ levels in primary mouse neurons. (a) APP_{C-term} peptide affinity for selected Mint2-PARM variants. Data are expressed as the mean + SEM ($n = 3$). The statistical significance was evaluated using one-way ANOVA with Dunnett's multiple comparison test, **** $P < 0.0001$ (Figure S10). (b) Coimmunoprecipitation of APP with the Mint2 antibody from HEK293T cells cotransfected with APP and GFP-rMint2^{WT} or GFP-rMint2^{Y460A/F521A}. Western blotting indicates less APP binding to rMint2^{Y460A/F521A} (lane 6) than to GFP-Mint2^{WT} (lane 5). α -Tubulin served as a loading control. (c) Quantification of APP coimmunoprecipitation using GFP-rMint2^{WT} and GFP-rMint2^{Y460A/F521A}. Data were normalized to GFP-rMint2^{WT} and expressed as the mean + SEM. The statistical significance was evaluated with the Student's t -test, *** $P < 0.001$. ($n = 4$ independent experiments). (d) Western blot for Mint2, APP, and α -tubulin from neuronal lysates carrying the APP^{swe}/PS1 E9 mutation and infected with the lentiviral GFP-rMint2^{Y460A/F521A} mutant. (e) A β ₄₂ ELISA quantification of conditioned media from neurons overproducing A β shows reduced A β ₄₂ levels when neurons were infected with the GFP-rMint2^{Y460A/F521A} mutant. Data were normalized to the endogenous Mint2^{WT} control and expressed as the mean + SEM ($n = 7$ biological replicates from two independent experiments). The statistical significance was evaluated using the Student's t -test, *** $P < 0.001$.

**Figure 4.**

Incorporation of ncAAs and evaluation of side chain-to-side chain macrocyclization in the APP_{WT} peptide. (a) Overview of synthesized APP_{WT} peptide variants, including the structure of the introduced amino acids for each position (blue). (b) K_i values of each APP_{WT} peptide variant measured by FP. n.b. indicates nonbinding peptide (i.e., $K_i > 500 \mu\text{M}$). Data are expressed as the mean + SEM ($n = 3$). (c) Structure of side chain-to-side chain cyclized APP_{WT} peptide analogues with native residue order (Glu in the i position, dark gray) and (d) inverse residue order (Lys in the i position, light gray). (e) K_i values of cyclic APP_{WT} peptide variants measured by FP. Data are expressed as the mean + SEM ($n = 3$). (f) Helical propensity of residues Y762 to E766 from the molecular dynamics simulation of cyclic APP_{WT} peptide variants **1** and **3–13** (% helical frames) plotted against the binding affinity (the mean K_i value) including the trend line (blue).

**Figure 5.**

PPI inhibitor of the APP-Mint2 interaction reduces Aβ₄₂ production in the neuronal *in vitro* model of AD. (a) Schematic structure of KSL-221036 (**14**). (b) Representative ITC of titrating Mint2-PARM with **14**; raw heat signature (top) and integrated molar heat release (bottom). (c) *In vitro* plasma stability of **14** (blue) and the APP_{WT} peptide (black) including buffer control for the APP_{WT} peptide (circles); data are expressed as the mean ± SEM (*n* = 3). (d) *In vitro* hepatic clearance of **14** (blue) and the APP_{WT} peptide (black) including propranolol control (circles, *n* = 1); data are expressed as the mean ± SEM (*n* = 3). (e)

Representative pull down of Mint2 from neuronal cell lysate (15 DIV). Lanes 1–3: expression of Mint2. Lanes 4–6: knockout of Mint2. The eluent was resolved by SDS-PAGE and immunoblotted for Mint2 and GAPDH ($n = 3$). (f) $A\beta_{42}$ ELISA quantification from conditioned media of neurons overproducing $A\beta$; data are normalized to the vehicle control and expressed as the mean + SEM ($n = 3$ independent experiments, $n = 10$ biological replicates for the vehicle, DAPT, and TAT-14 ($10 \mu\text{M}$)). Independent experiments are represented by different shapes. Statistical significance evaluated using one-way ANOVA with Dunnett's multiple comparison test, *** $P < 0.001$. (g) LDH levels from conditioned medium of cultured neurons overproducing $A\beta$ collected after 24 h of treatment with DAPT and TAT-14 ($10 \mu\text{M}$). Maximum LDH represents the maximum amount of LDH released from neurons lysed using detergent. Individual data points in (g) correspond to data in panel (f) (circles = same assay). Data are expressed as %LDH normalized to vehicle control and shown as the mean + SEM. Statistical significance was evaluated using one-way ANOVA with Sidak's multiple comparison test, *** $P < 0.001$.