# Amyloid β oligomer selective antibodies for Alzheimer's therapeutics and diagnostics

- **3 Running title: AβO-antibodies for therapeutics and diagnostics**
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Abbreviations: AD-Alzheimer's disease; AβO-Amyloid β oligomer; CSF-cerebrospinal fluid; GFAP-glial
 fibrillar acidic protein; ICV-intracerebroventricular; MNS-magnetic nanostructures; MRI-magnetic
 resonance imaging; PET-positron emission tomography; NOR-novel object recognition; NLR-novel

23 location recognition; PiB-Pittsburgh Compound B; pTau-phosphorylated tau; ThioS-thioflavin S; Tg-

24 transgenic; WT-wild-type

#### 25 Abstract

26 Improvements have been made in the diagnosis of Alzheimer's disease (AD), manifesting mostly in the 27 development of in vivo imaging methods that allow for the detection of pathological changes in AD by 28 MRI and PET scans. Many of these imaging methods, however, use agents that probe amyloid fibrils and 29 plaques- species that do not correlate well with disease progression and are not present at the earliest 30 stages of the disease. Amyloid  $\beta$  oligomers (A $\beta$ Os), rather, are now widely accepted as the A $\beta$  species 31 most germane to AD onset and progression. Here we report evidence further supporting the role of ABOs 32 as pathological instigators of AD and introduce a promising anti-ABO diagnostic probe capable of 33 distinguishing the 5xFAD mouse model from wild type mice by PET and MRI. In a developmental study, 34 Aβ oligomers in 5xFAD mice were found to appear at 3 months of age, just prior to the onset of memory 35 dysfunction, and spread as memory worsened. The increase is prominent in the subiculum and correlates 36 with concomitant development of reactive astrocytosis. The impact of these ABOs on memory is in 37 harmony with findings that intraventricular injection of synthetic ABOs into wild type mice induced 38 hippocampal dependent memory dysfunction within 24 hours. Compelling support for the conclusion that 39 endogenous AβOs cause memory loss was found in experiments showing that intranasal inoculation of 40 ABO-selective antibodies into 5xFAD mice completely restored memory function, measured 30 days post-41 inoculation. These antibodies, which were modified to give MRI and PET imaging probes, were able to 42 distinguish 5xFAD mice from wild type littermates. These results provide strong support for the role of 43 A $\beta$ Os in instigating memory loss and salient AD neuropathology, and they demonstrate that A $\beta$ O 44 selective antibodies have potential both for therapeutics and for diagnostics.

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47 KEYWORDS: Aβ oligomers; Alzheimer's disease; 5xFAD; MRI; PET; diagnostics; therapeutics

#### 49 Introduction

#### 50 General Alzheimer's disease

51 More than 6 million Americans are currently living with Alzheimer's disease (AD), and Alzheimer's-related 52 deaths have increased 145% from 2000 to 2019 (2021). The financial burden is even more staggering-53 Alzheimer's and other dementias have cost the US more than \$600 billion in medical expenses and 54 unpaid care in 2021 (2021). Despite the great personal and economic burden, progress toward 55 developing effective diagnostics and therapeutics remains slow. Aduhelm® (also known as 56 Aducanumab) was recently approved as a treatment for Alzheimer's disease (Investor Relations, 2021), 57 the first in more than a decade, but it still focuses on A $\beta$  elimination rather than specific A $\beta$ O targets. As 58 AD burden is expected to increase drastically with the aging population, improved diagnostics and 59 therapeutics are more urgent now than ever.

#### 60 **AβOs as a biomarker for early Alzheimer's disease**

The primary pathological hallmarks of Alzheimer's disease are extracellular amyloid plaques and intraneuronal tangles of hyperphosphorylated tau (Masters et al., 1985). It is well known, however, that amyloid plaques do not correlate well with cognitive decline in AD (Terry et al., 1991; Hsia et al., 1999; Lee et al., 2004) and are not present in the earliest stages of the disease (Nyborg et al., 2013). Research from the previous two decades strongly indicates that soluble amyloid beta oligomers (AβOs), not plaques, are the more appropriate amyloid beta species to target in AD (Ashe, 2020; Hampel et al., 2021).

67 ABOs are potent neurotoxins that show AD-dependent accumulation in the brain of AD patients (Gong et 68 al., 2003; Kayed et al., 2003; Lacor et al., 2004) and transgenic (Tg) rodent AD models (Chang et al., 69 2003; Lesne et al., 2006; Ohno et al., 2006). For reviews of other perspectives regarding AD molecular 70 etiology, see (Braak and Del Tredici, 2011; Robakis, 2011; Lasagna-Reeves et al., 2012). AβOs begin to 71 accumulate early in AD, decades prior to symptoms, and are widely held to be the neurotoxic instigators 72 of AD (Rodgers, 2005; Gandy et al., 2010; Schnabel, 2011; Mucke and Selkoe, 2012). AβOs have been 73 shown to exert their toxic effects by instigating failure of synaptic plasticity and memory (Lambert et al., 74 1998; Wang et al., 2002; Lesne et al., 2006; Townsend et al., 2006). Recently, soluble cortical extracts 75 were examined by ELISA and showed that the ratio of ABO levels to plaque density fully distinguished 76 demented from non-demented patients (Esparza et al., 2013); simply put, those with high ABO to plaque 77 ratios were demented and low ABO to plague ratios were not.

#### 78 The 5xFAD mouse model

79 The 5xFAD transgenic mice is an increasingly used AD model that harbors gene mutations in amyloid ß 80 protein precursor (ABPP) (K670N/M671L + I716V + V717I) and presenilins (PS1/2) (M146L + L286V) 81 (Oakley et al., 2006). These mutations are known to increase production of A $\beta$ 42, characteristic of familial 82 AD, and exhibit expedited plaque development compared to other transgenic mice (Oakley et al., 2006). 83 The Mutant Mouse Resource Research Center (MMRRC) found that AB accumulation occurred at 84 different rates, depending on the breeding background, with mice bred on a B6SJL background 85 developing pathology at a significantly more rapid rate (unpublished, available at MMRRC 5xFAD strain 86 data) than those bred on a C57 background. The 5xFAD mouse model is well characterized for memory 87 impairments (Oakley et al., 2006; Kimura and Ohno, 2009; Ohno, 2009; Girard et al., 2013; Girard et al., 88 2014; Zhang et al., 2021a), neuron loss (Jawhar et al., 2012; Oblak et al., 2021), and A $\beta$  plaque 89 accumulation (Devi et al., 2010; Jawhar et al., 2012; Ashe, 2020; Zhang et al., 2021a). Comprehensive 90 studies on the 5xFAD model have also looked at cholesterol and glucose levels (Oblak et al., 2021), 91 activity levels (Oblak et al., 2021), neuroinflammation-related protein levels (Ou-Yang and Van Nostrand, 92 2013; Oblak et al., 2021), tau phosphorylation (Shao et al., 2011; Kanno et al., 2014), and visual acuity 93 (Zhang et al., 2021a).

#### 94 Alzheimer's disease diagnostics

95 Recommended tests (Alzheimer's Disease Diagnostic Guidelines | National Institute on Aging (nih.gov))

for diagnosing Alzheimer's disease include a standard health evaluation and MMSE evaluations. If

97 indicated, these tests are typically followed with cerebrospinal fluid (CSF) assays for tau and A $\beta$  levels,

98 MRI for brain volume and functionality, and positron emission tomography (PET) scans for Aβ plaques, 99 glucose metabolism, and/or tau fibrils in the brain (Albert et al., 2011: Jack et al., 2011: McKhann et al.,

- glucose metabolism, and/or tau fibrils in the brain (Albert et al., 2011; Jack et al., 2011; McKhann et al.,
   2011; Sperling et al., 2011). These analyses may rule out other dementia etiologies and help to determine
- 101 disease severity, but they cannot detect AD at its earliest stages or closely predict disease progression,
- 102 as they do not probe for AD's earliest biomarkers.
- as they do not probe for AD's earliest biomarkers.

### 103 Current diagnostic methods in development

104 Spinal taps are invasive, but cerebrospinal fluid assays show promise (Georganopoulou et al., 2005; 105 Toledo et al., 2013b). Nonetheless, assays using CSF analytes have presented challenges with respect 106 to accuracy and reliable disease-state discrimination (Slemmon et al., 2012). More recently, assays for 107 ABO levels in the blood plasma have been developed with promising results (Meng et al., 2019). These 108 assays show a correlation between A $\beta$ O levels and declining memory scores that appear not to be 109 influenced by age, gender, or ApoE4 status. A promising addition to diagnostic methodology is the 110 detection of AD pathology using targeted in vivo brain imaging. The introduction of PET probes for 111 amyloid plagues has been a great technical advance (Klunk et al., 2004) and has established precedent 112 for the usefulness of brain molecular imaging as a diagnostic tool and for proof of efficacy in drug 113 development (Johnson et al., 2013). Still, these new imaging tools focus on late-stage AD pathology, not 114 on the early biomarkers such as ABOs.

- 115 Prior studies using 5xFAD mice have examined early and late-stage disease development, but none 116 have looked at the progressive development of ABOs in this model. Here, we present an analysis of 117 memory impairment from 2-9 months of age and the progressive accumulation of ABOs across the same 118 age-span. Our studies presented here use an ABO-selective antibody to characterize the spatiotemporal 119 development of ABOs in the 5xFAD mouse model and demonstrate a correlation with memory 120 impairment. Strikingly, intranasal inoculation of the ABO-selective antibody rescued memory performance 121 in 6-month-old 5xFAD mice. We demonstrate the capability of detecting ABO pathology in vivo in the 122 5xFAD mouse by introducing molecular imaging modalities (MRI and PET) with probes for A $\beta$ Os. We 123 additionally present immunofluorescent evidence of a remarkable association between ABOs and GFAP-124 positive reactive astrocytes in the 5xFAD mice. Taken together, we provide further data implicating ABOs 125 as essential diagnostic indicators and therapeutic targets, and show evidence suggesting a mechanism 126 through which A $\beta$ Os instigate pathological abnormalities: by induction of reactive astrogliosis.
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#### 128 Materials and Methods

#### 129 Materials

ACU193 humanized anti-AβO antibody was a generous gift from Acumen Pharmaceuticals, Inc. Aβ<sub>1-42</sub>
 (TFA preparation) was sourced from multiple suppliers. Primary hippocampal cultures were prepared
 from tissue obtained from BrainBits, LLC, using media and reagents also obtained from BrainBits. All

133 chemicals were purchased from Sigma unless otherwise specified.

#### 134 Animals

135 5xFAD Tg mouse model (B6SJL-Tg(APPSwFILon,PSEN1\*M146L\*L286V)6799Vas)(Oakley et al., 2006) 136 (Jackson Laboratories) was bred on a non-transgenic background (B6SJLF1/J mice (Jackson 137 Laboratories, RRID: IMSR JAX:100012)). Aged transgenic and wild-type littermates, 2-20 months old, 138 were used. All mice were kept under a 12/12 h light/dark cycle (7 AM/7 PM) at 22 ± 2 °C. Mice had free 139 access to food and water, including during behavioral experiments, were housed at ≤5/cage (NexGen 140 IVC, Allentown) with enriched environment and daily veterinarian assessment, according to NU's 141 standard procedures. Procedures complied with NIH's Guide for the Care and Use of Laboratory Animals 142 (NIH publication No. 80-23, 1996) and were approved by IACUC (protocol #IS00004010). Behavioral 143 experiments were conducted between 12-6 PM.

For intracerebroventricular (icv) experiments, B6SJLF1/J mice (Jackson Laboratories, RRID: IMSR\_JAX:100012) were utilized at ages ranging from 6 months of age (30-50 g).

#### 146 **A**β Oligomer Preparation

147 Fluorescently-labeled A<sup>β</sup> oligomers were prepared essentially according to the protocol published by 148 Klein and colleagues.(Lambert et al., 2007) Briefly, Aβ<sub>1-42</sub> (American Peptide or Peptides International) 149 or FAM-AB1-42 (Anaspec) was dissolved in hexafluoro-2-propanol (HFIP) and distributed into 150 microcentrifuge tubes. Hexafluoro-2-propanol was removed by evaporation and traces removed under 151 vacuum; the tubes were stored at -80°C. An aliquot of each was dissolved in anhydrous dimethyl sulfoxide 152 (DMSO) to ~5 mM, mixed 5:1 (mol: mol) Aβ: FAM-Aβ, and diluted in ice-cold Ham's F12 medium without 153 phenol red (Caisson Laboratories) to 100 µM. This solution was incubated at 4°C for 24 hr. and 154 centrifuged at 14 000 g for 10 min. The supernatant, defined as the FAM-ABO preparation, was 155 transferred to a clean microfuge tube and stored at 4°C until use. Protein concentration was determined 156 using Coomassie Plus protein assay kit (Pierce).

157 The same protocol was employed to prepare unlabeled A $\beta$  oligomers in the absence of FAM-A $\beta_{1-42}$ . A 158 modification of this protocol was used to produce crosslinked A $\beta$ Os (Cline et al., 2019b).

#### 159 Cell Culture

Hippocampal cells were prepared and maintained for at least 18 days as previously described (Gong et al., 2003) by using (0.002%) poly-L-lysine coated coverslips plated at a density of 1.04 x  $10^4$  cells per cm<sup>2</sup> in Neurobasal media (Brainbits, LLC) with B27 supplements and L-glutamine (2.5  $\mu$ M).

#### 163 **A**β Oligomer Incubation and Immunolabeling of Cells

164 Cells were incubated at 37°C in conditioned media collected from the cell cultures containing crosslinked 165 ABOs or FAM-ABOs or an equivalent dilution of vehicle. Following incubation with ABOs or vehicle for 60 166 min, the cells were rinsed rapidly 3 times with warm media then fixed by adding an equal volume of warm 167 3.7% formaldehyde (in PBS) to the third rinse in each well/dish and allowing it to sit at RT for 5 min. The 168 media/formaldehyde was completely removed and replaced with a volume of 3.7% formaldehyde for 5 169 min at RT. Cells were blocked in 10% normal goat serum (NGS) in PBS or HBSS for 45 min at RT then 170 incubated overnight at 4°C on an orbital shaker with fluorescent-tagged antibody or anti-ABO probe 171 diluted in blocking buffer. The cells were washed 3 times for 5 min each with PBS or HBSS. After

secondary antibody incubation, coverslips were mounted onto glass slides using ProLong Gold Anti-fade reagent with DAPI (Invitrogen) and imaged using an epifluorescence (TE2000, Nikon), a widefield

174 fluorescence microscope (Leica DM6B, Leica Corp.), or confocal microscope (Leica SP2, Leica Corp).

#### 175 **AβO intracerebroventricular (icv) administration in mice**

176 Icv injections and behavior testing were performed in 4 independent experiments of 13-21 mice each.

177 Littermates were arbitrarily assigned to different injection groups, targeting 5-10 mice/group for statistical 178 power (n =  $((Z_{\alpha/2}*\sigma)/E)^2$  at  $\alpha = 0.05$ ;  $\sigma = 10.55$  and E = 6.67 derived from pilot studies).

179 Mice were lightly anesthetized (2% isoflurane) during injection (~1 min). A $\beta$ Os (1, 10 pmol in 3  $\mu$ l) or vehicles were administered icv free-handed (Bicca et al., 2015). Separate needles were used for each 180 181 vehicle, progressing from low-high ABO concentration to minimize carryover. No analgesics or anti-182 inflammatory agents were necessary. Mice were monitored constantly for recovery of consciousness and 183 ambulation, then periodically for food-and-water intake until behavior analysis. Needle placement was 184 confirmed by brain dissection after behavioral experiments (euthanization: CO<sub>2</sub> then decapitation). Mice 185 showing needle misplacement (3 mice) or cerebral hemorrhage (2 mice) were excluded from analysis; 186 final n = 5-7 mice/group.

#### 187 Object Recognition/Location Recognition (NOR/NLR) Tasks

188 Tasks were performed essentially as described (Bicca et al., 2015), to evaluate mouse ability to 189 discriminate between familiar and new, or displaced, objects within an area, measured by object 190 exploration (sniffing, touching). The open-field testing arena was constructed of gray polyvinyl chloride at 191 21x21x12" (WxLxH), with a 5x5 square grid on floor and visual cue on wall. 24 h post-injection, mice 192 underwent 6 min sessions of habituation and training, with 3 min between. All sessions were video 193 recorded and analyzed by two researchers blind to experimental groups. During habituation and training, 194 mice were screened for ability to move about the arena and explore the objects, two activities required 195 for accurate memory assessment in subsequent testing sessions. Locomotive inclusion criteria (>100 196 grid crossings and >15 rearings; evaluated in habituation) were based on extensive previous experiments 197 with the same mouse strain and arena; 3/65 mice did not meet this criterion. During training, mice were 198 placed at the arena center with two objects, which were plastic and varied in shape, color, size and 199 texture. Exploration inclusion criteria were low exploration (<3 sec total) or object preference (>50% of 200 total time for either object); 7 of remaining 62 mice did not meet this criterion.

Hippocampal-related memory function was assessed 24 h post-training by displacing one of the two training objects. Cortical-related memory function was assessed 24 h later by replacing the displaced object with a novel object. Hippocampal-related memory function was re-tested 31-38 days post-injection by displacing the novel object. Memory dysfunction was defined as an exploration of the familiar object for >40% total time. Mice were arbitrarily assessed by cage. The arena and objects were cleaned thoroughly between sessions with 20% (v/v) alcohol to minimize olfactory cues.

#### 207 Immunolabeling of slices

208 Free floating 45 µm thick sagittal sections were cut using a Leica SM2010 R sliding microtome and 209 transferred to sterile TBS for storage. Sections were gathered and placed sequentially into wells (~4 per 210 well). Sections were then randomly selected from each well to perform antibody staining using the primary 211 antibodies ACU193 (0.2 µg/ml), Alexa Fluor® 555-conjugated NU4 (0.92 µg/ml), Cy3-conjugated anti-212 GFAP (1:800, Sigma) and the secondary antibody Alexa Fluor® 633 goat anti-human IgG (1:2000, 213 Invitrogen). Floating slices were rinsed 3x10 min with TBS and blocked with blocking buffer (10% NGS 214 with 0.3% Triton X-100 in TBS) for 60 min at room temperature. Slices were then incubated with the 215 respective antibodies in blocking buffer overnight at 4°C with gentle rotation. Sections were washed 3 x 216 10 mins in TBS and incubated with secondary antibody for 3 hours at room temperature (RT) with orbital 217 agitation in the dark. Secondary was prepared in blocking buffer diluted 10-fold with TBS. Sections were 218 then washed 3 x 10 mins in TBS, mounted using ProLong Diamond® antifade mounting media with DAPI

(Invitrogen) and 24x60mm No.1.5 glass coverslips (Thermo Scientific). Z-stacks of the brain sections
 were collected at 10x or 100x on a Leica SP5 confocal microscope and analyzed with ImageJ.

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#### 222 **Thioflavin S counterstain.**

223 Thioflavin-S counterstaining to NU4 immunofluorescence labeling was performed as previously 224 described (Guntern et al., 1992) with a few modifications (Viola et al., 2015). 5xFAD and WT brains were 225 sliced at a thickness of 50µm and immunolabeled following the same protocol described above 226 (immunolabeling of slices). Slices were incubated with antibody as described above. The slices were 227 then washed with PBS for 5 times 5 min each and incubated with 0.002% of Thioflavin-S solution in TBS-228 T (diluted from a stock solution 0.02% of Thioflavin-S in distillated water) for 10min. Slices were then 229 washed 3 times for 1 min in 50% ethanol and 2 times in TBS-T for 5 min. The slices were mounted with 230 ProLong Gold Antifade reagent for examination by fluorescence microscopy. Images were acquired at 231 40x magnification and analyzed by ImageJ software.

#### 232 Radiolabeling and Quality Control

Antibodies, NU4 and non-specific mouse IgG or ACU193 and non-specific human IgG were radiolabeled with positron emitter <sup>64</sup>Cu (<sup>64</sup>CuCl2 in 0.1 M HCl; radionuclide purity >99%, Washington University). For radiolabeling, Wipke and Wang's method was applied (Wipke et al., 2002). Basically, antibodies mentioned above were conjugated with DOTA-NHS-ester (Macrocyclics, Dallas, TX) and then radiolabeled with <sup>64</sup>Cu.

#### 238 Conjugation.

Antibody solutions were buffer exchanged with PBS using YM-30 Centricon® centrifugal filters (Millipore, Billerica, MA). For conjugation, antibodies were reacted with DOTA-NHS-ester in 0.1 M Na<sub>2</sub>HPO<sub>4</sub> buffer of pH 7.5 at 4°C for 12 - 16 h in a molar ratio of DOTA-NHS-ester:antibody = 100: 1. After conjugation, the reaction mixture was centrifuged repeatedly (5 times) through a YM-30 Centricon® centrifugal filter with 0.1M pH 6.5 ammonium citrate buffer to remove unconjugated small molecules. The concentrations of purified antibody-conjugate was determined by measuring the absorbance at 280 nm in a UV spectrophotometer.

#### 246 Labeling.

When labeling with <sup>64</sup>Cu, 1 mg DOTA-conjugated NU4 and 5 mCi (185 MBq) of <sup>64</sup>Cu as incubated in 0.1
 M ammonium citrate buffer, pH 6.5, at 43°C for 1 hour. Labeled antibody was separated by a size exclusion column (Bio-Spin6, BIO-RAD Laboratories).

#### 250 **Quality Control.**

Radiochemical purity of antibody was determined by integrating areas on the Fast Protein Liquid Chromatography (FPLC) equipped with a flow scintillation analyzer. This analysis was conducted on a superpose 12 size-exclusion column and characterized by the percentage of radioactivity associated with the 150 kDa protein peak. The stability of the <sup>64</sup>Cu radiolabeled mAbs was determined by bovine serum challenge at 44 hours.

#### 256 **Conjugation efficiency.**

Based on our preliminary data, > 90% of conjugation rate, >70% of labeling rate is achieved by following
 prescribed protocol.

#### 259 **Overall details of micro PET and micro CT acquisition**

260 Mice were placed in a 37.5 °C heated cage 20-30 minutes prior to radiotracer injection and moved to a

261 37.5 °C heated induction chamber 10 minutes prior to injection where they were anesthetized with 2-3%

262 isoflurane in 1000 cc/min O<sub>2</sub>. A dose of 40  $\mu$ g/200  $\mu$ Ci in 100  $\mu$ L of proposed PET tracers was

administered intravenously through the tail vein. Each animal was administered a dose ranging from 30-

264 40 µg NU4PET, ACU193PET, or non-immune IgGPET. Probes were administered in a single dose. 265 PET/CT imaging was conducted at 0, 4, 24, and 48 h to measure for changes in distribution and time 266 required for probe clearance or decay.

267 NU4PET scans were acquired using a Genisys<sup>4</sup> PET (Sofie Biosciences, Culver City, CA) system and 268 CT scans were acquired using a Bioscan NanoSPECT/CT (Washington, D.C.). When scanning, all mice 269 were placed prone on the bed of the scanner. A 10 minute static acquisition was used for PET imaging 270 followed immediately by a 6.5 minute CT acquisition both utilizing the mouse imaging chamber from the 271 Genisys<sup>4</sup>. PET reconstruction was performed without attenuation correction using 3D Maximum 272 Likelihood Expectation Maximization (MLEM) with 60 iterations and CT reconstruction used Filtered Back 273 Projection with a Shepp-Logan Filter. PET and CT reconstructions were exported in dicom image format 274 and fused using custom software developed by the Small Animal Imaging Facility at Van Andel Institute. 275 Fused PET/CT images were analyzed using VivoQuant Image Analysis Suite (inviCRO, LLC, Boston, 276 MA). Standardized Uptake Values (SUV) were calculated using the mouse body weight and corrected 277 for residual dose in the injection syringe and the injection site, as applicable. The formula used to 278 calculate SUV was

 $SUV = \frac{\text{Activity}_{\text{tissue}}/\text{Volume}_{\text{tissue}}}{\text{Injected Activity}/\text{BodyWeight}}.$ 279

#### Evaluation NU4PET (<sup>64</sup>Cu-NU4) in AβOs detection 280

281 2 groups (n = 3/ group) of 6 months old 5xFAD Tg AD mouse model and 2 groups (n = 3/ group) WT 282 mouse model were used for evaluating the capability of AβOs detection. NU4PET (<sup>64</sup>Cu-NU4) or non-283 specific IgGPET (<sup>64</sup>Cu-IgG) was injected into each 5xFAD Tg AD mouse model and WT mouse model 284 groups, respectively.

285 Target (AβOs)–Background (normal tissue) contrasts in PET images were used to distinguish the 286 difference of the capability of ABOs detection between NU4PET and IgGPET in different mouse models. 287 Tracer uptake of high intensity (hot) areas and background tissues in the brain were chosen by drawing 288 regions-of-interest (ROI) along the edges of the areas from the PET images. Average pixel values of 289 each ROIs were acquired and use in Target ( $A\beta Os$ )–Background (normal tissue) contrasts calculation. 290 The formula used to calculate Target-Background contrast was

 $T - B \ Contrast = \frac{1}{Background_{Average Pixel Value}}$ 291

#### 292 **Tissue Biodistribution Assessment**

299

293 Animals were sacrificed immediately after the 44 hour post injection image was acquired. Blood was 294 collected, while brains and 13 other organs and tissues were harvested and weighed. After the blood 295 sample was taken from the heart (~500-1000µl), 10 ml of saline was injected into left ventricle while the 296 heart was still beating to flush out the residual blood in the organs. Radioactivity in each tissue (cpm) wa 297 measured using the y-scintillation counter. Percentages of the injected dose/gram (%ID/g) were 298 calculated for each tissue/ organ by the following formula.

$$\% {}^{ID}/g = \frac{(Sample \ Activity - Background)}{(Injected \ Activity - Background)(Sample \ weight(g))} \times 100\%$$

300 Student's t-test was conducted to the results between different groups. P<0.05 is considered statistically 301 significant.

#### 302 Synthesis of Magnetic Nanostructures (MNS)

303 16 nm magnetite nanoparticles were synthesized by decomposition of iron-oleate at 320°C as described 304 in an earlier report.(Park et al., 2004)

305 Synthesis of Iron-oleate complexes: 10.8 g of iron (III) chloride hexahydrate and 36.5 g sodium oleate 306 were dissolved in a mixture of 60 ml distilled water, 80 ml ethanol and 140 ml hexane and heated at 60°C

- for 4 hr. The organic layer of the biphasic mixture becomes dark, indicating phase transfer of iron (III) ions and formation of iron oleate complex. The resulting dark solution is separated and washed with water three times
- 309 three times.
- 310 *Synthesis of 16 nm magnetite nanoparticles:* 18 g of iron oleate complex and 2.58 g of oleic acid were
- dissolved in 100 g of octadecene at room temperature and heated to 320°C at a rate of 3.3°C per minute.
- 312 The reaction mixture is kept at 320°C for 40 min., then cooled down to room temperature. Resulting
- 313 nanoparticles are separated from the solution by addition of ethanol and ethyl acetate followed by
- 314 centrifugation.

#### 315 Preparation of Dopamine-TEG-COOH and Phase Transfer

- To make the organic phase synthesized MNS suitable for biological application, we functionalized the
- 317 MNS using an in-house synthesized ligand with carboxylate as terminal group (for antibody conjugation),
- tetraehylene glycol(TEG) as a stabilizer, and nitrodopamine (nDOPA) as an anchor due to its high affinity for Ee (Nandwana et al. 2016)
- 319 for Fe (Nandwana et al., 2016).
- Synthesis of carboxylate terminated nDOPA ligand and functionalization of the MNS was carried out according to the following protocol. Tetraethylene diacide, N-hydroxysuccinimide (NHS), N,N'-Dicyclohexylcarbodiimide (DCC), nDOPA hydrochloride and anhydrous sodium bicarbonate was dissolved in chloroform under argon atmosphere and stirred for 4 hr. Hexane stabilized MNS were added and stirred for another 24 hr. The precipitate formed was separated by magnet, dispersed in water and purified by dialysis.

#### 326 Conjugation of antibody to MNS

The conjugation of buffer stabilized MNS with antibody was done using a conventional carboxyl-amine crosslinking method. We first activated the carboxyl terminated MNS by sulfo-N-hydroxy succinimide (SNHS) and 1-Ethyl-3-(3-dimethylaminopropyl)carbodiimide (EDC) followed by incubation with corresponding antibody (NU4 or IgG<sub>1</sub>, with or without fluorescent label) overnight. Conjugated MNS were separated by magnet to remove excess reagent and antibody then re-dispersed in working media. Conjugation efficiency was estimated using UV spectroscopy (absorbance at 280nm) of the magnetically separated supernatant.

Ab conc. = (total mg added Ab) - (mg Ab in supernatant)

#### 335 Intranasal immunization.

Mice were anesthetized with isoflurane and then placed on their backs with their heads positioned to maximize the residency time for the delivered material to remain on the olfactory surface. Each naris was administered with ACUMNS or non-immune IgGMNS (10 µl/naris), using a sterile micropipette, slowly over a period of 1 min, keeping the opposite naris and mouth closed to allow complete aspiration of delivered material. Steps were repeated up to 5 times, maintaining anesthetization in between inoculations, for maximum doses of up to 50µl/naris

#### 342 Magnetic Resonance Imaging of Tg and WT mice in vivo

343 Following intranasal inoculation, the probe was allowed to distribute for 4 hours before MR imaging was 344 performed according to imaging methodology described in Mundt et al. (Mundt et al., 2009) T1, T2, and 345 T2\* weighted MR images were acquired on a Bruker BioSpec 9.4T magnet, using a 25 mm RF guadrature 346 coil. The in-plane resolution was 75 µm with slice thickness 0.4 mm. T1- and T2-weighted images provide 347 anatomical guidance as well as some localization of the ACUMNS and were acquired with a fat 348 suppressed spin echo sequence (Rapid Acquisition with Relaxation Enhancement, RARE) with the 349 following parameters for T1-weighted (TR=1000 ms, TEeff=13.2 ms, rare factor 2, number of excitations, 350 NEX=4) and for T2-weighted (TR=3500 ms, TEeff=58.5 ms, rare factor 4, NEX=4). T2\*-weighted imaging 351 provides more of the localization of the NU4MNS as the iron causes local changes in magnetic 352 susceptibility which T2\* weighted images can be sensitive to. A gradient echo sequence was used with

the following parameters (gradient echo fast imaging, GEFI; TR=1200 ms, TE=5.6 ms, flip angle 35° and

354 NEX=4).

#### 356 **Results**

# Memory dysfunction in 5xFAD mice begins shortly after AβO emergence and progressively worsens with concomitant AβO accumulation in the hippocampus

#### 359 Tg 5xFAD NOR/NLR

360 Amyloid plague development and intraneuronal Aβ42 accumulation are well-established in the 5xFAD 361 transgenic (Tg) mouse model of Alzheimer's disease. There is robust plaque buildup around 5-6 months 362 of age (Ohno et al., 2006) and intraneuronal Aβ42 accumulation begins as early as 2 months (Oakley et 363 al., 2006). The majority of neuropathological studies in 5xFAD mice have used probes that show amyloid 364 plaque development; how 5xFAD memory impairment coincides with AβOs pathology and development 365 is much less well-characterized. In order to characterize how memory loss correlates with ABOs in the 366 5xFAD mice, we used the well-established novel object recognition task (NOR) for non-spatial (cortical) 367 memory (Cohen and Stackman, 2015; Denninger et al., 2018) and the novel location recognition task 368 (NLR) for spatial (hippocampal) memory (Antunes and Biala, 2012; Bengoetxea et al., 2015; Grayson et 369 al., 2015; Denninger et al., 2018). We assessed memory in mice aged 2-18 months. 5xFAD mice showed 370 no evident memory impairment at 2 to 3 months old (Figure 1a). By 4 to 5 months old, most transgenic 371 mice showed memory impairment, and by 6 to 7 months of age memory impairment was apparent in all 372 5xFAD mice. Importantly, at 4 months old, the majority of 5xFAD mice were impaired in both the 373 hippocampal-dependent and cortical-dependent tasks; there were, however, some mice that showed 374 only cortical-impairment. Though less obvious than their Tg littermates, memory loss was detected at 9 375 months of age in wild-type mice. In summary, we showed that 5xFAD mice first present memory 376 impairment between 3 and 4 months of age. This memory dysfunction afflicts more mice as their age 377 increases until, at 6 to 7 months, all of the Tg mice are impaired in both hippocampal-dependent and 378 cortical-dependent tasks. These data indicate that memory impairment begins before observed amyloid 379 plaque build-up in the 5xFAD mice.

380 Immunohistofluorescence validation of AβO development

381 The development of amyloid plaque pathology is well- established in the 5xFAD mouse model (Oakley 382 et al., 2006; Ohno et al., 2006). Amyloid plagues, however, are no longer considered the most germane 383 Aβ species to AD pathology (Overk and Masliah, 2014; Viola and Klein, 2015; Selkoe and Hardy, 2016; 384 Cline et al., 2018; Li and Selkoe, 2020). Characterizing the development of the most relevant species, 385 putatively A $\beta$ Os, and their association with other pathological changes in AD, such as glial activation or 386 pTau accumulation, is necessary to better understand disease progression in this model. Sagittal 387 sections of brain tissue, collected and fixed from WT and 5xFAD mice at ages 2, 3, 4, 6, and 8 months 388 of age, were immunolabeled with ACU193 and imaged using confocal microscopy. ACU193, a 389 humanized monoclonal antibody that targets ABOs, has been shown to selectively bind oligomers in vitro 390 (Krafft et al., 2013; Goure et al., 2014; Savage et al., 2014) and in the TG2576 mouse model. Here, using 391 ACU193 to probe for ABOs, we show the progressive, spatio-temporal accumulation of ABOs in the 392 hippocampus of 5xFAD mice (Figure 1b). AβOs first appear in the subiculum as early as 2 months of age 393 (not shown), followed by continued accumulation in the subiculum and a spreading of pathology to CA1, 394 CA2 and the dentate gyrus. This timing suggests that A $\beta$ Os are associated with the observed memory 395 loss.

#### 396 <u>ACU193 detects AβOs bound to primary neurons with high specificity</u>

To validate the specificity of ACU193 for A $\beta$ Os, the antibody was used *in vitro* to detect synthetic preparations of oligomers introduced into primary hippocampal neurons in culture (Supplemental Figure 1). Primary hippocampal neurons were treated with cross-linked A $\beta$ Os, which have been shown to preserve A $\beta$ O structure *in vitro* (Cline et al., 2019b), or vehicle control. The cells were subsequently fixed and labeled with ACU193 at increasing dosages. Confocal imaging of the cells showed somatic staining of A $\beta$ Os in addition to small, nanoscale puncta along dendritic processes (labeled with MAP2). These ACU193-positive puncta are likely A $\beta$ Os binding to dendritic spines, as seen in previously published work 404 (Lacor et al., 2007; De Felice et al., 2009; Pitt et al., 2017). Minimal ACU193 labeling was observed on
 405 vehicle-treated neurons, indicating its specificity for AβOs.

#### 406 ACU193 and NU4 detect AβOs

407 Additional support for the specificity of ACU193 can be seen in comparing the distribution of ACU193 in

- 408 brain sections with the distribution of NU4, a well-established AβO monoclonal antibody (Lambert et al.,
  409 2007; Xiao et al., 2013; Viola et al., 2015). Using ACU193 and NU4 conjugated to Alex Fluor 555 we
- 410 found that both antibodies similarly detected ABOs in the subiculum and other areas of the hippocampus
- 411 (Figure 2) including CA1, CA2 and the dentate gyrus. ACU193- (cyan) and NU4-positive (magenta) cells
- 412 were observed accumulating in a nearly identical pattern, from 3 months to nine months of age. ACU193
- 413 and NU4 selectively detect ÅβOs in the 5xFAD mice with virtually no signal in WT mice.

# 414 Other Alzheimer's-associated pathologies also show developmental regulation in the 415 5xFAD mouse model

416 To determine whether other Alzheimer's related pathologies show developmental regulation or 417 accumulation in the 5xFAD mouse model for AD in association with ABOs, we examined 418 immunohistochemical patterns of glial fibrillary acidic protein (GFAP), activated microglia (Iba1), and 419 phosphorylated tau (pTau). Immunolabeling for pTau yielded difficult to interpret results which varied 420 amongst the different antibodies for the same epitope and often did not match the literature. Instead, we 421 focused on the inflammatory pathways, stimulated by the strong interest in the involvement of 422 inflammatory responses in AD, in particular a new and growing interest in astrocytes (Wang et al., 2021). 423 Sagittal sections from 5xFAD mice, aged 3-9 months, or their WT littermates were immunolabeled with 424 antibodies against GFAP and co-labeled with ACU193, then imaged by confocal microscopy. We found 425 a marked spatiotemporal association of GFAP pathology with ACU193-positive ABOs in the 5xFAD mice. 426 GFAP (Figure 3, magenta) pathology first appeared in the subiculum at 3 months of age concurrent with 427 the first appearance of ABOs (cyan) in the subiculum and in close proximity to one another. As the mice 428 aged, GFAP and ACU193-positive pathology concomitantly spread throughout the subiculum and 429 hippocampus (Figure 3, B & E). At 9 months, WT mice have minimal GFAP expression (Figure 3C) and 430 no AβOs (Figure 3F). These patterns are consistent with possible induction of reactive astrogliosis by 431 ABOs. At higher magnification, we observed GFAP-positive reactive astrocytes surrounding an ACU193-432 positive neuron and projecting their processes onto the cell soma (Figure 3I). In addition, we observed 433 micron-wide ACU193-positive puncta adjacent to astrocytic processes distant from the cell soma. 434 Immunolabeling for activated microglia (Iba1) (data not shown) indicated that the WT mice have more 435 ramified microglial cells (resting) while 5xFAD littermates have more amoeboid and activated-shaped 436 microglial cells. Microglial activation was evident at 2 months, with some increase in abundance seen in 437 older animals.

#### 438 AβOs given to WT littermates induces memory impairment within 24 hours

#### 439 <u>ICV AβOs induce impairment in NLR/NOR</u>

440 While the previous data indicate a relationship between ABO accumulation and memory dysfunction in 441 the 5xFAD mice, the question remained whether AβOs cause the observed memory loss. We therefore 442 asked whether injection of ABOs into WT littermate mice would induce similar behavioral dysfunction. 443 Wild-type littermates from the 5xFAD colony were injected with 10pmol synthetic ABOs, or vehicle control 444 into the right lateral ventricle, following our previously established protocol (Cline et al., 2019b). After 24 445 hours, the mice were assessed by the NLR task, and later, the NOR assay at 48 hours post-injection. 446 We found that ICV injection of ABOs induce memory dysfunction within 24 hours and impacts both cortical 447 (NOR) and hippocampal (NLR) memory (Figure 4). As in the 5xFAD mice, A $\beta$ O injected mice showed no 448 preference to either new or old objects and explored both equally. Vehicle-injected mice scored no 449 different from wild-type in these tasks. These data show that ABOs are sufficient to induce memory 450 impairment within 24 hours post-injection in wild-type mice. We next sought to establish the functional
 451 effect of neutralizing these AβOs in the 5xFAD mice.

# 452 Oligomer-selective antibodies engage and neutralize AβOs responsible for memory 453 dysfunction in 5xFAD mice

#### 454 <u>ACU193-based probes ameliorate memory dysfunction</u>

455 We have previously observed no short-term detrimental impact after inoculation of our ABO antibodies 456 into 5xFAD mice, but no studies have been done to determine the long-term positive or negative effects 457 in these mice. To determine the impact of AβO-neutralization in 5xFAD mice, 6- and 7-month-old mice 458 were first assessed for memory impairment using the NLR/NOR assay. Mice were then inoculated with 459 ACU193-based probes and imaged 24 hours later in vivo to ensure target engagement (see next section). 460 The mice were then housed for 30-40 days to monitor any adverse effects or changes in behavior before 461 being reassessed for memory impairment in the NLR/NOR tasks. Strikingly, we found that 6-month-old 462 5xFAD mice inoculated with the ACU193-based MRI probe had reversal of memory dysfunction, with 463 performance the same as WT controls in the NOR task 30 days post-inoculation (Figure 5). The NLR 464 assay showed the same restoration of memory (not shown). The ACUPET probe similarly ameliorated 465 memory impairment, measured 40 days post-injection. As a control, 5xFAD mice injected with a human 466 IgGPET probe showed no memory improvement (data not shown). Results show that the ABO-selective 467 ACU193 antibody engages ABOs in vivo, completely reversing memory dysfunction in the 5xFAD mice 468 with no evidence of pathological side effects. The data establish  $A\beta Os$  as the primary instigators of 469 cognitive dysfunction in 5xFAD mice and support the therapeutic relevance of A $\beta$ O-selective probes.

### 470 AβOs imaged *in vivo* using ACU193-based probes distinguish 5xFAD from wild-type 471 mice

#### 472 MRI signal from ACUMNS distinguishes 5xFAD from wild-type mice.

473 Our previous work showed that ABOs can be detected *in vivo* in the 5xFAD mouse model using antibody-474 based MRI probes which were conjugated to magnetic nanostructures (MNS) (Viola et al., 2015). These 475 prior studies used NU4 as the AβO-targeting antibody, which as shown above, binds similarly to ACU193. 476 Here we show that ACU193 can also be developed into a molecular probe for ABO detection in vivo. 477 After baseline imaging by MRI, 12-month-old mice were intranasally inoculated with MNS-conjugated 478 ACU193 and allowed to recover overnight (about 16 hours) before imaging again (Figure 6). MRI data 479 shows an accumulation of the ACUMNS probe in the hippocampus and cortex of the 5xFAD mice that is 480 absent in WT controls. ImageJ quantification of signal intensity in the hippocampi of inoculated mice 481 shows a ~ 30-fold increase in 5xFAD mice over their WT littermates. Using the ACUMNS probe in 18-482 month-old mice showed similarly robust AD-dependent MRI signal in the hippocampus of the 5xFAD 483 animals, but signals obtained in younger animals (6-months old) were less consistent. These data add to 484 previous studies with the NU4 probe and show that non-invasive in vivo imaging of ABOs is possible 485 using the ACUMNS probe, suggesting its potential diagnostic value and ability to confirm target 486 engagement

#### 487 <u>Development of an ACU193-based PET imaging probe for early AβO detection.</u>

488 While the spatial resolution of MRI is excellent, its sensitivity is lower than other imaging modalities such 489 as positron emission tomography (PET). Given PET sensitivity is at least 100 times greater than MRI, we 490 thought it might detect very low levels of ABOs during early stages of AD development. ACU193 was 491 conjugated to DOTA, a chelator, as the initial step in the PET probe development. To ensure that this 492 conjugation did not interfere with the antibody's ability to target ABOs, sagittal brain slices from 5xFAD 493 mice were probed with the ACU193-DOTA probe and counterstained with Thioflavin S (ThioS) for amyloid 494 plaques (Supplemental Figure 2). Results show that ACU193-DOTA detected ABOs in the 5xFAD brain 495 and did not co-localize with ThioS, consistent with previously obtained results showing that ACU193 does 496 not bind amyloid plaques cores (Cline et al., 2019a).

#### 497 ACUPET detects pathology in the brains of 4-month and older 5xFAD mice.

- The next step was to determine if radiolabeled ACU193-DOTA (ACUPET) detects AD-related AβOs in the 5xFAD mouse brain at an early age. ACU193-DOTA was incubated with <sup>64</sup>Cu and free isotopes were removed prior to tail vein injection into mice of either 4 or 18 months old, Mice were then imaged at 1, 4, and 24 hours post-injection for ACUPET distribution. At 4 hours post-injection, ACUPET accumulation in the brain was detectable (not shown), but not robust. By 24 hours, accumulation of the ACUPET probe in the brains of the 5xFAD animals was evident in both the 4-month-old animals (Supplemental Figure 3A) and the 18 month old animals (Supplemental Figure 3B-D). Animals at 6, 7, 8 and 12 months were
- also examined and similarly were able to distinguish 5xFAD from WT mice (data not shown).

#### 506 **AβOs are specifically detected in vivo by NU4PET**

#### 507 <u>NU4-based PET probe development</u>

- 508 Given the success of the NU4-based MRI probe (Viola et al., 2015), an NU4-based probe was 509 synthesized for PET imaging. NU4 was conjugated to DOTA and tested to ensure that this conjugation 510 did not interfere with the antibody's ability to target A $\beta$ Os. Primary hippocampal neurons, pre-treated with 511 fluorescently conjugated AβOs (FAM-AβOs) and were probed with NU4-DOTA (Supplemental Figure 4). 512 Data show that nearly all FAM-ABOs (magenta) were also labeled with the NU4-DOTA probe 513 (colocalization seen as dark blue) and no free NU4-DOTA (cyan) was detected. Vehicle treated cells 514 showed no NU4-DOTA binding. Data confirm the specificity of the NU4-DOTA probe for A $\beta$ Os, necessary 515 for its use for in vivo imaging.
- 516 NU4PET detects AD-related pathology *in vivo* in 5xFAD mice, distinguishing them from WT
- 517 Validation of the ABO-PET probes as effective for early AD diagnostics requires verification that they 518 produce an *in vivo* signal that depends on the presence of ABOs. To validate our new probe, NU4 519 (Lambert et al., 2007; Acton et al., 2010) and non-specific IgG antibodies were conjugated to DOTA and 520 then radiolabeled with positron emitter <sup>64</sup>Cu using Wipke and Wang's method (Wipke et al., 2002). Our 521 next step was to image for AβOs by PET following probe delivery. Animals (12 total), 7 months of age, 522 were injected via tail vein with either NU4PET or IgGPET and then imaged at T=1, 2, 4, 8, 20, 30, 40, 523 and 44 hours after injection. After 44 hours, the animals were euthanized and their brains removed for a 524 final ex vivo image of all 12 brains simultaneously (3 animals per group). Results showed the NU4PET 525 specifically identified 5xFAD animals (Figure 7). No signal was detected in all three control groups (5xFAD 526 with IgGPET;WT with NU4PET; WT with IgGPET).
- 527 The fraction of NU4PET probe retained (Supplemental Figure 5) showed good uptake into the brains of 528 the 5xFAD mice but not the WT littermates (quantification of uptake; see Methods). For all mice, the 529 IgGPET probe showed negligible signal. Quantification showed uptake into the brain was comparable to 530 levels of uptake seen with the commercially available Pittsburgh Compound B (PiB) tracer (Mathis et al., 531 2003; Klunk et al., 2004). To corroborate the presence of A $\beta$ Os in the animals used for these studies, we 532 analyzed the brain tissue with immunofluorescence. After final PET imaging, the brains were fixed and 533 stored in 10% sucrose until no longer radioactive. Brains were then sliced sagittally at 50 µm and probed 534 with ACU193. Images were collected and analyzed for ACU193 signal intensity (Supplemental Figure 6). 535 Data showed that only 5xFAD mice, and not WT littermates, had A $\beta$ O pathology. Results confirm the 536 NU4 PET probe gives a signal selective for AβO-positive mice.
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538 <b>DISCUSSION</b>
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539 Alzheimer's disease is costly and marked by accumulation of pathological hallmarks such as amyloid 540 plaques and neuronal tangles of hyperphosphorylated tau. Because AB plaques have shown poor 541 correlation with AD progression, there has been a rise in the exploration and development of therapeutics 542 that are not based on amyloid (Cummings et al., 2021). This shift in focus has resulted in numerous 543 potential therapies that have made it into clinical trials, but so far there have been limitations on the impact 544 of these potential therapies. As an alternative, focusing on ABOs as the target for diagnostics and 545 therapeutics appears to be a promising strategy for developing disease modifying treatments and early 546 diagnosis. Here, we confirm that ABOs can induce memory dysfunction in wild type mice and that ABOs 547 build up in 5xFAD mice in a manner concomitant with astrocyte pathology and with memory dysfunction. 548 Importantly, targeting this buildup with  $A\beta O$ -selective antibodies rescues memory performance. 549 Furthermore, we demonstrate that antibody-based brain imaging probes that target A $\beta$ Os can be used 550 to identify animals that present with AD pathology, indicating the value of A $\beta$ O-selective antibodies both 551 for diagnostics and therapeutics.

552 Recent interest in inflammatory processes and their involvement in AD has grown. Our data showed a 553 striking association between GFAP-positive astrocytes and ACU193-positive ABOs. This association and 554 concomitant increase indicates a potential mechanism for ABO-induced behavioral abnormalities. These 555 findings are particularly intriguing given recent studies indicating AD's dependence on astrocytes. One 556 especially interesting study showed that when apolipoprotein E (ApoE), a protein expressed in astrocytes 557 which AβOs associate with at synapses, was knocked out in astrocyte-only populations of P301S mice, 558 AD pathology markedly improved (Wang et al., 2021). As ApoE4 is the greatest genetic risk factor of late 559 onset AD, we propose that it may mediate ABO-induced reactive astrogliosis and the subsequent 560 neuropathology instigated by reactive astrocytes. Another study showed that astrocytes were activated 561 into their reactive state via the JAK/STAT3 pathway in 6 month-old 5xFAD mice (Choi et al., 2020). 562 Consistent with the idea that reactive astrogliosis is necessary for behavioral dysfunction in 5xFAD mice, 563 STAT3 phosphorylation inhibition restored cognitive function in the 5xFAD mice. Taken together with our 564 data, we propose that ABOs may induce JAK/STAT3 pathway-dependent reactive astrogliosis in 565 astrocytes which is necessary for observed cognitive dysfunction in 5xFAD mice. In addition to 566 astrocytes, microglia play a major role in AD pathology. The Triggering Receptor Expressed on Myeloid 567 cells 2(TREM2)- expressed in microglia- has already been shown to be involved in AD, with mutations 568 being neuroprotective and TREM2 accumulation being detected in AD patients (Jiang et al., 2013; 569 Benitez et al., 2014; Guven et al., 2020). Previous studies have shown that AβOs associate with TREM2 570 (Zhao et al., 2018; Zhong et al., 2019; Price et al., 2020), but TREM2 has no impact on established 571 pathology (Yuan et al., 2021).

572 While interest increases in alternatives to the Amyloid Hypothesis, we are still left with no effective 573 diagnostic tools for identifying AD at its earliest stages when therapeutics have the greatest impact. 574 Currently recommended tests may rule out other dementia etiologies and help to determine disease 575 severity, but they cannot detect AD at its earliest stages or closely predict disease progression as they 576 do not probe for AD's earliest biomarkers. While AD diagnosis has significantly improved with the 577 incorporation of a multiple assay evaluation currently being recommended, the tests still cannot predict 578 disease progression or diagnose AD at its earliest stages because they are not quantifying the earliest 579 biomarkers of the disease. However, alternative detection assays are being developed. Pre-tangle Tau, 580 thought to be the toxic form of tau, has now been detected in MCI and AD and has been found to be one 581 of the earliest tau lesions that correlates with cognitive status (Mufson et al., 2014). Synapse loss (Bastin 582 et al., 2020; Buchanan et al., 2020; Camporesi et al., 2020; Mecca et al., 2020; Pereira et al., 2021), 583 changes in hormone levels (Cheng et al., 2021), changes in blood biomarker levels (Guzman-Martinez 584 et al., 2019; Montoliu-Gaya et al., 2021), electroencephalogram (EEG) readings (Hulbert and Adeli, 2013; 585 Siwek et al., 2015; Lin et al., 2021), retinal assays (Ashok et al., 2020; Mirzaei et al., 2020), and changes 586 in specific protein levels (Buchanan et al., 2020; Colom-Cadena et al., 2020) are some of the myriad

assays being developed to try to detect AD earlier and predict when and if the change from mild cognitive
 impairment (MCI) to AD will occur (Zhang et al., 2021b). All of these new developments are focused
 towards enabling earlier therapeutic intervention when chances for success would be greatest.

590 AβOs as a diagnostic resource are currently unavailable. Cerebrospinal fluid assays show promise 591 (Georganopoulou et al., 2005; Toledo et al., 2013a; Savage et al., 2014; Yang et al., 2015; Yang et al., 592 2019), but spinal taps are invasive and assays using CSF analytes have presented challenges with 593 respect to accuracy and reliable disease-state discrimination (Slemmon et al., 2012). Other assays for 594 AβO levels are under development and show promise as well (Meng et al., 2019). For example, AβO 595 guantification in blood plasma shows a correlation between ABO levels and declining memory scores that 596 appear to not be influenced by age, gender, or ApoE4 status. Recently, the examination of soluble 597 cortical extracts by ELISA found a link between the ratio of ABOs and fibrils with disease. "The ratio of 598 ABO levels to plaque density fully distinguished demented from non-demented patients, with no overlap 599 between groups in this derived variable." (Esparza et al., 2013)

- 600 Because ABOs are regarded as the first toxin to appear in disease progression, they should provide an 601 excellent target for diagnostic imaging (Hefti et al., 2013; Goure et al., 2014). The usefulness of targeting 602 ABOs is indicated by human neuropathology studies in which ABOs initially appear bound to discrete 603 neurons, localizing to synapses in dendritic arbours (Lacor et al., 2004) through putative association with 604 clustered cell surface receptors (Ferreira and Klein, 2011). FAM-ABOs bind at discrete sites on dendrites, 605 showing saturable, concentration-dependent synaptic binding (Viola et al., 2015), further suggesting their potential as a suitable target for an antibody-based diagnostic probe. Pronucleon<sup>™</sup> imaging used 606 607 engineered peptides that deliver a readout when associated with beta-rich Aß fibers and oligomeric Aß 608 (Nyborg et al., 2013). Several PET probes have also been developed including a probe from curcumin<sup>18</sup>F 609 (Rokka et al., 2014), a probe created by modifying 6E10 antibody with PEG and <sup>64</sup>Cu that distinguished Tg from control mice (McLean et al., 2012), and a probe developed from an <sup>124</sup>I-labeled mAb158 against 610 611 A $\beta$  protofibrils (Magnusson et al., 2013). Still, none of these probes specifically target A $\beta$ Os.
- 612 Previously, we described a molecular MRI probe that is targeted against AβOs (Viola et al., 2015). Based 613 on the success of our initial MRI probe and the antibody-based probes being explored by others, it is 614 reasonable to predict that AβO-specific antibodies can be used to target probes and provide better signal-615 to-noise ratios. Here we showed that anti-AβO antibodies can be used to develop molecular MRI ad PET 616 probes that distinguish WT mice from their 5xFAD littermates at ages as early as 4 months old. These 617 probes have proven to be non-toxic over the periods examined and, in fact, showed *in vivo* efficacy.
- 618 Early diagnostics are critical to combating this devastating disease, but without effective therapeutics, 619 they have limited value. The first FDA-approved drug to treat Alzheimer's disease (AD) in nearly two 620 decades, Aduhelm<sup>®</sup>, shows a preferential affinity for all aggregated forms of amyloid beta (A $\beta$ ), rather 621 than targeting only the toxic A $\beta$ Os. Currently, there are more than 126 agents in clinical trials, with most 622 aimed at disease modification (Cummings, 2021; Cummings et al., 2021). While less than 10% of these 623 target A $\beta$ , there remains evidence that A $\beta$  is a significant target for the rapeutic development. Lowering 624 AβO levels by enhancing fibril formation has been shown to be protective (Mucke et al., 2000). This is 625 supported by previous antibody-based studies (Lambert et al., 2007; Xiao et al., 2013). The data 626 presented here importantly show that A $\beta$ O-selective antibodies rescue memory performance in a widely 627 used AD model. These antibodies, which have been modified for use in brain imaging of A $\beta$ O, show great 628 promise as potential agents for AD therapeutics and diagnostics; the potential of one AβO-selective 629 antibody is now being assessed in a recently begun clinical trial.
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### 650 **Contribution to the field statement**

651 Alzheimer's disease is costly and marked by pathological damage and progressive memory loss. While 652 there has been progress made towards developing better therapeutics and diagnostics, it has been 653 limited. Diagnostic improvements have primarily been in the development of better imaging methods. 654 mostly using agents that probe amyloid fibrils and plaques- species that do not correlate well with disease 655 progression and are not present at the earliest stages of the disease. Amyloid β oligomers (AβOs) are 656 now widely accepted as the Aβ species most germane to AD onset and progression. Here we report 657 evidence further supporting the role of ABOs in Alzheimer's disease and introduce a promising anti-ABO 658 diagnostic probe capable of distinguishing the 5xFAD mouse model from wild type mice by PET and MRI. 659 Our studies also showed a concomitant development of memory impairment with the accumulation of 660 ABOs and reactive astrocytes. Compelling support for the conclusion that ABOs cause memory loss was 661 found in experiments showing that AβO-selective antibodies into 5xFAD mice completely restored 662 memory function. These antibodies, modified to give imaging probes, were able to distinguish 5xFAD 663 mice from wild type littermates. These results demonstrate that ABO selective antibodies have potential 664 both for therapeutics and for diagnostics.

665	References
666	(2021). 2021 Alzheimer's disease facts and figures. <i>Alzheimers Dement</i> 17(3), 327-406. doi:
667	10.1002/alz.12328.
668	Acton, P.Q., PA, US), An, Z.A., PA, US), Bett, A.J.L., PA, US), Breese, R.Q., PA, US), Chang,
669	L.W., IL, US), Dodson, E.C.S., PA, US), et al. (2010). Anti-ADDL antibodies and uses
670	<i>thereof.</i> United States patent application 11/256332. 08/24/2010.
671	Albert, M.S., DeKosky, S.T., Dickson, D., Dubois, B., Feldman, H.H., Fox, N.C., et al. (2011).
672	The diagnosis of mild cognitive impairment due to Alzheimer's disease: recommendations
673	from the National Institute on Aging-Alzheimer's Association workgroups on diagnostic
674	guidelines for Alzheimer's disease. Alzheimers Dement 7(3), 270-279. doi:
675	10.1016/j.jalz.2011.03.008.
676	Antunes, M., and Biala, G. (2012). The novel object recognition memory: neurobiology, test
677	procedure, and its modifications. Cogn Process 13(2), 93-110. doi: 10.1007/s10339-011-
678	0430-z.
679	Ashe, K.H. (2020). The biogenesis and biology of amyloid beta oligomers in the brain.
680	Alzheimers Dement 16(11), 1561-1567. doi: 10.1002/alz.12084.
681	Ashok, A., Singh, N., Chaudhary, S., Bellamkonda, V., Kritikos, A.E., Wise, A.S., et al. (2020).
682	Retinal Degeneration and Alzheimer's Disease: An Evolving Link. Int J Mol Sci 21(19).
683 684	doi: 10.3390/ijms21197290. Bastin, C., Bahri, M.A., Meyer, F., Manard, M., Delhaye, E., Plenevaux, A., et al. (2020). In vivo
685	imaging of synaptic loss in Alzheimer's disease with [18F]UCB-H positron emission
686	tomography. Eur J Nucl Med Mol Imaging 47(2), 390-402. doi: 10.1007/s00259-019-
687	04461-x.
688	Bengoetxea, X., Rodriguez-Perdigon, M., and Ramirez, M.J. (2015). Object recognition test for
689	studying cognitive impairments in animal models of Alzheimer's disease. Front Biosci
690	(Schol Ed) 7, 10-29.
691	Benitez, B.A., Jin, S.C., Guerreiro, R., Graham, R., Lord, J., Harold, D., et al. (2014). Missense
692	variant in TREML2 protects against Alzheimer's disease. Neurobiol Aging 35(6),
693	1510.e1519-1526. doi: 10.1016/j.neurobiolaging.2013.12.010.
694	Bicca, M.A., Costa, R., Loch-Neckel, G., Figueiredo, C.P., Medeiros, R., and Calixto, J.B. (2015).
695	B(2) receptor blockage prevents Abeta-induced cognitive impairment by
696	neuroinflammation inhibition. <i>Behav Brain Res</i> 278, 482-491. doi:
697 698	10.1016/j.bbr.2014.10.040.
699 699	Braak, H., and Del Tredici, K. (2011). Alzheimer's pathogenesis: is there neuron-to-neuron propagation? <i>Acta Neuropathol</i> 121(5), 589-595. doi: 10.1007/s00401-011-0825-z.
700	Buchanan, H., Mackay, M., Palmer, K., Tothová, K., Katsur, M., Platt, B., et al. (2020). Synaptic
701	Loss, ER Stress and Neuro-Inflammation Emerge Late in the Lateral Temporal Cortex
702	and Associate with Progressive Tau Pathology in Alzheimer's Disease. <i>Mol Neurobiol</i>
703	57(8), 3258-3272. doi: 10.1007/s12035-020-01950-1.
704	Camporesi, E., Nilsson, J., Brinkmalm, A., Becker, B., Ashton, N.J., Blennow, K., et al. (2020).
705	Fluid Biomarkers for Synaptic Dysfunction and Loss. Biomark Insights 15,
706	1177271920950319. doi: 10.1177/1177271920950319.
707	Chang, L., Bakhos, L., Wang, Z., Venton, D.L., and Klein, W.L. (2003). Femtomole
708	immunodetection of synthetic and endogenous amyloid-beta oligomers and its application
709	to Alzheimer's disease drug candidate screening. <i>J Mol Neurosci</i> 20(3), 305-313. doi:
710	10.1385/JMN:20:3:305.

- Cheng, Y.J., Lin, C.H., and Lane, H.Y. (2021). From Menopause to Neurodegeneration Molecular Basis and Potential Therapy. *Int J Mol Sci* 22(16). doi: 10.3390/ijms22168654.
- Choi, M., Kim, H., Yang, E.J., and Kim, H.S. (2020). Inhibition of STAT3 phosphorylation
  attenuates impairments in learning and memory in 5XFAD mice, an animal model of
  Alzheimer's disease. *J Pharmacol Sci* 143(4), 290-299. doi: 10.1016/j.jphs.2020.05.009.
- Cline, E., Viola, K., Klein, W., Wang, X., Bacskai, B., Rammes, G., et al. (2019a). "Synaptic intervention in Alzheimer's disease: soluble Aβ oligomer directed ACU193 monoclonal antibody therapeutic for treatment of early Alzheimer's disease", in: *Clinical Trials on Alzheimer's disease*. (San Diego, CA, USA).
- Cline, E.N., Bicca, M.A., Viola, K.L., and Klein, W.L. (2018). The Amyloid-beta Oligomer
   Hypothesis: Beginning of the Third Decade. *J Alzheimers Dis* 64(s1), S567-S610. doi:
   10.3233/JAD-179941.
- Cline, E.N., Das, A., Bicca, M.A., Mohammad, S.N., Schachner, L.F., Kamel, J.M., et al. (2019b).
  A novel crosslinking protocol stabilizes amyloid beta oligomers capable of inducing
  Alzheimer's-associated pathologies. *J Neurochem* 148(6), 822-836. doi:
  10.1111/jnc.14647.
- Cohen, S.J., and Stackman, R.W., Jr. (2015). Assessing rodent hippocampal involvement in the
   novel object recognition task. A review. *Behav Brain Res* 285, 105-117. doi:
   10.1016/j.bbr.2014.08.002.
- Colom-Cadena, M., Spires-Jones, T., Zetterberg, H., Blennow, K., Caggiano, A., DeKosky, S.T.,
  et al. (2020). The clinical promise of biomarkers of synapse damage or loss in Alzheimer's
  disease. Alzheimers Res Ther 12(1), 21. doi: 10.1186/s13195-020-00588-4.
- Cummings, J. (2021). Drug Development for Psychotropic, Cognitive-Enhancing, and Disease Modifying Treatments for Alzheimer's Disease. *J Neuropsychiatry Clin Neurosci* 33(1), 3 13. doi: 10.1176/appi.neuropsych.20060152.
- Cummings, J., Lee, G., Zhong, K., Fonseca, J., and Taghva, K. (2021). Alzheimer's disease drug
  development pipeline: 2021. *Alzheimers Dement (N Y)* 7(1), e12179. doi:
  10.1002/trc2.12179.
- De Felice, F.G., Vieira, M.N., Bomfim, T.R., Decker, H., Velasco, P.T., Lambert, M.P., et al.
  (2009). Protection of synapses against Alzheimer's-linked toxins: insulin signaling
  prevents the pathogenic binding of Abeta oligomers. *Proc Natl Acad Sci U S A* 106(6),
  1971-1976. doi: 10.1073/pnas.0809158106.
- Denninger, J.K., Smith, B.M., and Kirby, E.D. (2018). Novel Object Recognition and Object
   Location Behavioral Testing in Mice on a Budget. *J Vis Exp* (141). doi: 10.3791/58593.
- Devi, L., Alldred, M.J., Ginsberg, S.D., and Ohno, M. (2010). Sex- and brain region-specific
  acceleration of beta-amyloidogenesis following behavioral stress in a mouse model of
  Alzheimer's disease. *Mol Brain* 3, 34. doi: 10.1186/1756-6606-3-34.
- Esparza, T.J., Zhao, H., Cirrito, J.R., Cairns, N.J., Bateman, R.J., Holtzman, D.M., et al. (2013).
  Amyloid-beta oligomerization in Alzheimer dementia versus high-pathology controls. *Ann Neurol* 73(1), 104-119. doi: 10.1002/ana.23748.
- Ferreira, S.T., and Klein, W.L. (2011). The Abeta oligomer hypothesis for synapse failure and
  memory loss in Alzheimer's disease. *Neurobiol Learn Mem* 96(4), 529-543. doi:
  10.1016/j.nlm.2011.08.003.
- Gandy, S., Simon, A.J., Steele, J.W., Lublin, A.L., Lah, J.J., Walker, L.C., et al. (2010). Days to
  criterion as an indicator of toxicity associated with human Alzheimer amyloid-beta
  oligomers. *Ann Neurol* 68(2), 220-230. doi: 10.1002/ana.22052.

- Georganopoulou, D.G., Chang, L., Nam, J.M., Thaxton, C.S., Mufson, E.J., Klein, W.L., et al.
  (2005). Nanoparticle-based detection in cerebral spinal fluid of a soluble pathogenic
  biomarker for Alzheimer's disease. *Proc Natl Acad Sci U S A* 102(7), 2273-2276. doi:
  10.1073/pnas.0409336102.
- Girard, S.D., Baranger, K., Gauthier, C., Jacquet, M., Bernard, A., Escoffier, G., et al. (2013).
  Evidence for early cognitive impairment related to frontal cortex in the 5XFAD mouse
  model of Alzheimer's disease. *J Alzheimers Dis* 33(3), 781-796. doi: 10.3233/jad-2012120982.
- Girard, S.D., Jacquet, M., Baranger, K., Migliorati, M., Escoffier, G., Bernard, A., et al. (2014).
  Onset of hippocampus-dependent memory impairments in 5XFAD transgenic mouse
  model of Alzheimer's disease. *Hippocampus* 24(7), 762-772. doi: 10.1002/hipo.22267.
- Gong, Y., Chang, L., Viola, K.L., Lacor, P.N., Lambert, M.P., Finch, C.E., et al. (2003).
  Alzheimer's disease-affected brain: presence of oligomeric A beta ligands (ADDLs)
  suggests a molecular basis for reversible memory loss. *Proc Natl Acad Sci U S A* 100(18),
  10417-10422. doi: 10.1073/pnas.1834302100.
- Goure, W.F., Krafft, G.A., Jerecic, J., and Hefti, F. (2014). Targeting the proper amyloid-beta
   neuronal toxins: a path forward for Alzheimer's disease immunotherapeutics. *Alzheimers Res Ther* 6(4), 42. doi: 10.1186/alzrt272.
- Grayson, B., Leger, M., Piercy, C., Adamson, L., Harte, M., and Neill, J.C. (2015). Assessment
  of disease-related cognitive impairments using the novel object recognition (NOR) task in
  rodents. *Behav Brain Res* 285, 176-193. doi: 10.1016/j.bbr.2014.10.025.
- Guntern, R., Bouras, C., Hof, P.R., and Vallet, P.G. (1992). An improved thioflavine S method
   for staining neurofibrillary tangles and senile plaques in Alzheimer's disease. *Experientia* 48(1), 8-10. doi: 10.1007/BF01923594.
- Guven, G., Bilgic, B., Samanci, B., Gurvit, H., Hanagasi, H., Donmez, C., et al. (2020). Peripheral
   TREM2 mRNA levels in early and late-onset Alzheimer disease's patients. *Mol Biol Rep* 47(8), 5903-5909. doi: 10.1007/s11033-020-05661-7.
- Guzman-Martinez, L., Maccioni, R.B., Farías, G.A., Fuentes, P., and Navarrete, L.P. (2019).
   Biomarkers for Alzheimer's Disease. *Curr Alzheimer Res* 16(6), 518-528. doi: 10.2174/1567205016666190517121140.
- Hampel, H., Hardy, J., Blennow, K., Chen, C., Perry, G., Kim, S.H., et al. (2021). The Amyloidbeta Pathway in Alzheimer's Disease. *Mol Psychiatry*. doi: 10.1038/s41380-021-012490.
- Hefti, F., Goure, W.F., Jerecic, J., Iverson, K.S., Walicke, P.A., and Krafft, G.A. (2013). The case
  for soluble Abeta oligomers as a drug target in Alzheimer's disease. *Trends Pharmacol Sci* 34(5), 261-266. doi: 10.1016/j.tips.2013.03.002.
- Hsia, A.Y., Masliah, E., McConlogue, L., Yu, G.Q., Tatsuno, G., Hu, K., et al. (1999). Plaqueindependent disruption of neural circuits in Alzheimer's disease mouse models. *Proc Natl Acad Sci U S A* 96(6), 3228-3233.
- Hulbert, S., and Adeli, H. (2013). EEG/MEG- and imaging-based diagnosis of Alzheimer's disease. *Rev Neurosci* 24(6), 563-576. doi: 10.1515/revneuro-2013-0042.
- Investor Relations, B. (2021). "FDA grants accelerated approval for ADUHELM<sup>™</sup> as the first and
   only Alzheimer's disease treatment to address a defining pathology of the disease".
   (www.biogen.com: Biogen).
- Jack, C.R., Jr., Albert, M.S., Knopman, D.S., McKhann, G.M., Sperling, R.A., Carrillo, M.C., et
   al. (2011). Introduction to the recommendations from the National Institute on Aging-

Alzheimer's Association workgroups on diagnostic guidelines for Alzheimer's disease.
 *Alzheimers Dement* 7(3), 257-262. doi: 10.1016/j.jalz.2011.03.004.

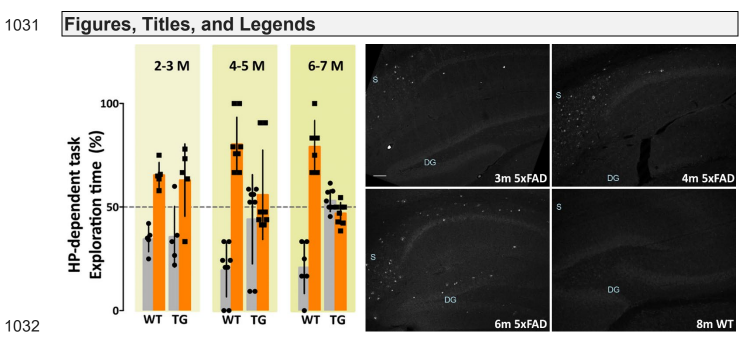
- Jawhar, S., Trawicka, A., Jenneckens, C., Bayer, T.A., and Wirths, O. (2012). Motor deficits, neuron loss, and reduced anxiety coinciding with axonal degeneration and intraneuronal Abeta aggregation in the 5XFAD mouse model of Alzheimer's disease. *Neurobiol Aging* 33(1), 196 e129-140. doi: 10.1016/j.neurobiolaging.2010.05.027.
- 809 Jiang, T., Yu, J.T., Zhu, X.C., and Tan, L. (2013). TREM2 in Alzheimer's disease. *Mol Neurobiol* 810 48(1), 180-185. doi: 10.1007/s12035-013-8424-8.
- Johnson, K.A., Minoshima, S., Bohnen, N.I., Donohoe, K.J., Foster, N.L., Herscovitch, P., et al.
  (2013). Appropriate use criteria for amyloid PET: a report of the Amyloid Imaging Task
  Force, the Society of Nuclear Medicine and Molecular Imaging, and the Alzheimer's
  Association. *Alzheimers Dement* 9(1), e-1-16. doi: 10.1016/j.jalz.2013.01.002.
- Kanno, T., Tsuchiya, A., and Nishizaki, T. (2014). Hyperphosphorylation of Tau at Ser396 occurs
  in the much earlier stage than appearance of learning and memory disorders in 5XFAD
  mice. *Behav Brain Res* 274, 302-306. doi: 10.1016/j.bbr.2014.08.034.
- Kayed, R., Head, E., Thompson, J.L., McIntire, T.M., Milton, S.C., Cotman, C.W., et al. (2003).
  Common structure of soluble amyloid oligomers implies common mechanism of pathogenesis. *Science* 300(5618), 486-489. doi: 10.1126/science.1079469.
- Kimura, R., and Ohno, M. (2009). Impairments in remote memory stabilization precede
   hippocampal synaptic and cognitive failures in 5XFAD Alzheimer mouse model. *Neurobiol Dis* 33(2), 229-235. doi: 10.1016/j.nbd.2008.10.006.
- Klunk, W.E., Engler, H., Nordberg, A., Wang, Y., Blomqvist, G., Holt, D.P., et al. (2004). Imaging
  brain amyloid in Alzheimer's disease with Pittsburgh Compound-B. *Ann Neurol* 55(3),
  306-319. doi: 10.1002/ana.20009.
- Krafft, G., Hefti, F., Goure, W., Jerecic, J., Iverson, K., and Walicke, P. (2013). ACU-193: A
  candidate therapeutic antibody that selectively targets soluble beta-amyloid oligomers. *Alzheimer's & Dementia* 9(4, Supplement), P326. doi:
  <u>http://dx.doi.org/10.1016/j.jalz.2013.04.166</u>.
- Lacor, P.N., Buniel, M.C., Chang, L., Fernandez, S.J., Gong, Y., Viola, K.L., et al. (2004).
  Synaptic targeting by Alzheimer's-related amyloid beta oligomers. *J Neurosci* 24(45), 10191-10200. doi: 10.1523/JNEUROSCI.3432-04.2004.
- Lacor, P.N., Buniel, M.C., Furlow, P.W., Clemente, A.S., Velasco, P.T., Wood, M., et al. (2007).
  Abeta oligomer-induced aberrations in synapse composition, shape, and density provide a molecular basis for loss of connectivity in Alzheimer's disease. *J Neurosci* 27(4), 796-807. doi: 10.1523/JNEUROSCI.3501-06.2007.
- Lambert, M.P., Barlow, A.K., Chromy, B.A., Edwards, C., Freed, R., Liosatos, M., et al. (1998).
   Diffusible, nonfibrillar ligands derived from Abeta1-42 are potent central nervous system
   neurotoxins. *Proc Natl Acad Sci U S A* 95(11), 6448-6453.
- Lambert, M.P., Velasco, P.T., Chang, L., Viola, K.L., Fernandez, S., Lacor, P.N., et al. (2007).
   Monoclonal antibodies that target pathological assemblies of Abeta. *J Neurochem* 100(1),
   23-35. doi: 10.1111/j.1471-4159.2006.04157.x.
- Lasagna-Reeves, C.A., Castillo-Carranza, D.L., Sengupta, U., Guerrero-Munoz, M.J., Kiritoshi,
   T., Neugebauer, V., et al. (2012). Alzheimer brain-derived tau oligomers propagate
   pathology from endogenous tau. *Sci Rep* 2, 700. doi: 10.1038/srep00700.
- Lee, S.P., Falangola, M.F., Nixon, R.A., Duff, K., and Helpern, J.A. (2004). Visualization of betaamyloid plaques in a transgenic mouse model of Alzheimer's disease using MR

- 849 microscopy without contrast reagents. *Magn Reson Med* 52(3), 538-544. doi: 10.1002/mrm.20196.
- Lesne, S., Koh, M.T., Kotilinek, L., Kayed, R., Glabe, C.G., Yang, A., et al. (2006). A specific amyloid-beta protein assembly in the brain impairs memory. *Nature* 440(7082), 352-357.
  doi: 10.1038/nature04533.
- Li, S., and Selkoe, D.J. (2020). A mechanistic hypothesis for the impairment of synaptic plasticity
   by soluble Abeta oligomers from Alzheimer's brain. *J Neurochem* 154(6), 583-597. doi:
   10.1111/jnc.15007.
- Lin, N., Gao, J., Mao, C., Sun, H., Lu, Q., and Cui, L. (2021). Differences in Multimodal Electroencephalogram and Clinical Correlations Between Early-Onset Alzheimer's Disease and Frontotemporal Dementia. *Front Neurosci* 15, 687053. doi: 10.3389/fnins.2021.687053.
- Magnusson, K., Sehlin, D., Syvanen, S., Svedberg, M.M., Philipson, O., Soderberg, L., et al.
  (2013). Specific uptake of an amyloid-beta protofibril-binding antibody-tracer in AbetaPP
  transgenic mouse brain. *J Alzheimers Dis* 37(1), 29-40. doi: 10.3233/jad-130029.
- Masters, C.L., Simms, G., Weinman, N.A., Multhaup, G., McDonald, B.L., and Beyreuther, K.
   (1985). Amyloid plaque core protein in Alzheimer disease and Down syndrome. *Proc Natl Acad Sci U S A* 82(12), 4245-4249.
- Mathis, C.A., Wang, Y., Holt, D.P., Huang, G.F., Debnath, M.L., and Klunk, W.E. (2003).
  Synthesis and evaluation of 11C-labeled 6-substituted 2-arylbenzothiazoles as amyloid imaging agents. *J Med Chem* 46(13), 2740-2754. doi: 10.1021/jm030026b.
- McKhann, G.M., Knopman, D.S., Chertkow, H., Hyman, B.T., Jack, C.R., Jr., Kawas, C.H., et al.
  (2011). The diagnosis of dementia due to Alzheimer's disease: recommendations from
  the National Institute on Aging-Alzheimer's Association workgroups on diagnostic
  guidelines for Alzheimer's disease. *Alzheimers Dement* 7(3), 263-269. doi:
  10.1016/j.jalz.2011.03.005.
- McLean, D., Cooke, M.J., Wang, Y., Green, D., Fraser, P.E., George-Hyslop, P.S., et al. (2012).
   Anti-amyloid-beta-mediated positron emission tomography imaging in Alzheimer's disease mouse brains. *PLoS One* 7(12), e51958. doi: 10.1371/journal.pone.0051958.
- Mecca, A.P., Chen, M.K., O'Dell, R.S., Naganawa, M., Toyonaga, T., Godek, T.A., et al. (2020).
   In vivo measurement of widespread synaptic loss in Alzheimer's disease with SV2A PET.
   *Alzheimers Dement* 16(7), 974-982. doi: 10.1002/alz.12097.
- Meng, X., Li, T., Wang, X., Lv, X., Sun, Z., Zhang, J., et al. (2019). Association between
   increased levels of amyloid-β oligomers in plasma and episodic memory loss in
   Alzheimer's disease. *Alzheimer's Research & Therapy* 11(1), 89. doi: 10.1186/s13195 019-0535-7.
- Mirzaei, N., Shi, H., Oviatt, M., Doustar, J., Rentsendorj, A., Fuchs, D.T., et al. (2020).
  Alzheimer's Retinopathy: Seeing Disease in the Eyes. *Front Neurosci* 14, 921. doi: 10.3389/fnins.2020.00921.
- Montoliu-Gaya, L., Strydom, A., Blennow, K., Zetterberg, H., and Ashton, N.J. (2021). Blood
   Biomarkers for Alzheimer's Disease in Down Syndrome. *J Clin Med* 10(16). doi:
   10.3390/jcm10163639.
- Mucke, L., Masliah, E., Yu, G.Q., Mallory, M., Rockenstein, E.M., Tatsuno, G., et al. (2000).
   High-level neuronal expression of abeta 1-42 in wild-type human amyloid protein precursor transgenic mice: synaptotoxicity without plaque formation. *Journal of Neuroscience* 20(11), 4050-4058.

- Mucke, L., and Selkoe, D.J. (2012). Neurotoxicity of amyloid beta-protein: synaptic and network
   dysfunction. *Cold Spring Harb Perspect Med* 2(7), a006338. doi:
   10.1101/cshperspect.a006338.
- Mufson, E.J., Ward, S., and Binder, L. (2014). Prefibrillar tau oligomers in mild cognitive
  impairment and Alzheimer's disease. *Neurodegener Dis* 13(2-3), 151-153. doi:
  10.1159/000353687.
- Mundt, A.P., Winter, C., Mueller, S., Wuerfel, J., Tysiak, E., Schnorr, J., et al. (2009). Targeting
   activated microglia in Alzheimer's pathology by intraventricular delivery of a
   phagocytosable MRI contrast agent in APP23 transgenic mice. *Neuroimage* 46(2), 367 372. doi: 10.1016/j.neuroimage.2009.01.067.
- Nandwana, V., Ryoo, S.-R., Kanthala, S., De, M., Chou, S.S., Prasad, P.V., et al. (2016).
   Engineered Theranostic Magnetic Nanostructures: Role of Composition and Surface
   Coating on Magnetic Resonance Imaging Contrast and Thermal Activation. ACS Applied
   Materials & Interfaces 8(11), 6953-6961. doi: 10.1021/acsami.6b01377.
- Nyborg, A.C., Moll, J.R., Wegrzyn, R.D., Havas, D., Hutter-Paier, B., Feuerstein, G.G., et al.
  (2013). In vivo and ex vivo imaging of amyloid-beta cascade aggregates with a Pronucleon peptide. *J Alzheimers Dis* 34(4), 957-967. doi: 10.3233/jad-122107.
- Oakley, H., Cole, S.L., Logan, S., Maus, E., Shao, P., Craft, J., et al. (2006). Intraneuronal betaamyloid aggregates, neurodegeneration, and neuron loss in transgenic mice with five familial Alzheimer's disease mutations: potential factors in amyloid plaque formation. *J Neurosci* 26(40), 10129-10140. doi: 10.1523/JNEUROSCI.1202-06.2006.
- Oblak, A.L., Lin, P.B., Kotredes, K.P., Pandey, R.S., Garceau, D., Williams, H.M., et al. (2021).
   Comprehensive Evaluation of the 5XFAD Mouse Model for Preclinical Testing
   Applications: A MODEL-AD Study. *Front Aging Neurosci* 13, 713726. doi:
   10.3389/fnagi.2021.713726.
- Ohno, M. (2009). Failures to reconsolidate memory in a mouse model of Alzheimer's disease.
   *Neurobiol Learn Mem* 92(3), 455-459. doi: 10.1016/j.nlm.2009.05.001.
- Ohno, M., Chang, L., Tseng, W., Oakley, H., Citron, M., Klein, W.L., et al. (2006). Temporal memory deficits in Alzheimer's mouse models: rescue by genetic deletion of BACE1. *Eur J Neurosci* 23(1), 251-260. doi: 10.1111/j.1460-9568.2005.04551.x.
- Ou-Yang, M.H., and Van Nostrand, W.E. (2013). The absence of myelin basic protein promotes
   neuroinflammation and reduces amyloid beta-protein accumulation in Tg-5xFAD mice. J
   *Neuroinflammation* 10, 134. doi: 10.1186/1742-2094-10-134.
- Overk, C.R., and Masliah, E. (2014). Toward a unified therapeutics approach targeting putative
   amyloid-beta oligomer receptors. *Proc Natl Acad Sci U S A* 111(38), 13680-13681. doi:
   10.1073/pnas.1414554111.
- Park, S.Y., Avraham, H.K., and Avraham, S. (2004). RAFTK/Pyk2 activation is mediated by
   trans-acting autophosphorylation in a Src-independent manner. *J Biol Chem* 279(32),
   33315-33322. doi: 10.1074/jbc.M313527200.
- 934 Pereira, J.B., Janelidze, S., Ossenkoppele, R., Kvartsberg, H., Brinkmalm, A., Mattsson-935 Carlgren, N., et al. (2021). Untangling the association of amyloid- $\beta$  and tau with synaptic 936 and axonal loss in Alzheimer's disease. Brain 144(1), 310-324. doi: 937 10.1093/brain/awaa395.
- Pitt, J., Wilcox, K.C., Tortelli, V., Diniz, L.P., Oliveira, M.S., Dobbins, C., et al. (2017).
   Neuroprotective astrocyte-derived insulin/insulin-like growth factor 1 stimulates endocytic

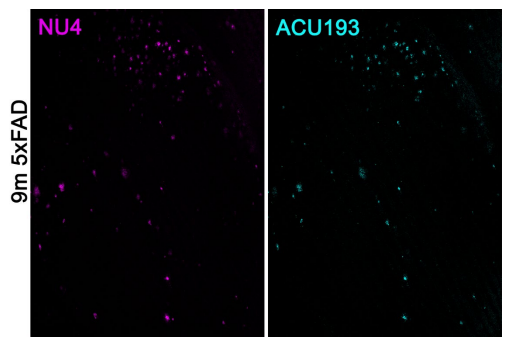
- processing and extracellular release of neuron-bound Abeta oligomers. *Mol Biol Cell*28(20), 2623-2636. doi: 10.1091/mbc.E17-06-0416.
- Price, B.R., Sudduth, T.L., Weekman, E.M., Johnson, S., Hawthorne, D., Woolums, A., et al.
  (2020). Therapeutic Trem2 activation ameliorates amyloid-beta deposition and improves
  cognition in the 5XFAD model of amyloid deposition. *J Neuroinflammation* 17(1), 238. doi:
  10.1186/s12974-020-01915-0.
- Robakis, N.K. (2011). Mechanisms of AD neurodegeneration may be independent of Abeta and
  its derivatives. *Neurobiol Aging* 32(3), 372-379. doi:
  10.1016/j.neurobiolaging.2010.05.022.
- Rodgers, A.B. (2005). "Progress report on Alzheimer's disease 2004-2005". U.S.Department of
   Health and Human Services; National Institutes on Aging; National Institutes of Health).
- Rokka, J., Snellman, A., Zona, C., La Ferla, B., Nicotra, F., Salmona, M., et al. (2014). Synthesis
   and evaluation of a (18)F-curcumin derivate for beta-amyloid plaque imaging. *Bioorg Med Chem* 22(9), 2753-2762. doi: 10.1016/j.bmc.2014.03.010.
- Savage, M.J., Kalinina, J., Wolfe, A., Tugusheva, K., Korn, R., Cash-Mason, T., et al. (2014). A
  sensitive abeta oligomer assay discriminates Alzheimer's and aged control cerebrospinal
  fluid. *J Neurosci* 34(8), 2884-2897. doi: 10.1523/jneurosci.1675-13.2014.
- 957 Schnabel, J. (2011). Amyloid: little proteins, big clues. *Nature* 475(7355), S12-14. doi: 10.1038/475S12a.
- Selkoe, D.J., and Hardy, J. (2016). The amyloid hypothesis of Alzheimer's disease at 25 years.
   *EMBO Mol Med* 8(6), 595-608. doi: 10.15252/emmm.201606210.
- Shao, C.Y., Mirra, S.S., Sait, H.B., Sacktor, T.C., and Sigurdsson, E.M. (2011). Postsynaptic degeneration as revealed by PSD-95 reduction occurs after advanced Abeta and tau pathology in transgenic mouse models of Alzheimer's disease. *Acta Neuropathol* 122(3), 285-292. doi: 10.1007/s00401-011-0843-x.
- Siwek, M.E., Muller, R., Henseler, C., Trog, A., Lundt, A., Wormuth, C., et al. (2015). Altered
  Theta Oscillations and Aberrant Cortical Excitatory Activity in the 5XFAD Model of
  Alzheimer's Disease. *Neural Plast* 2015, 781731. doi: 10.1155/2015/781731.
- Slemmon, J.R., Meredith, J., Guss, V., Andreasson, U., Andreasen, N., Zetterberg, H., et al.
  (2012). Measurement of Abeta1-42 in cerebrospinal fluid is influenced by matrix effects. *J Neurochem* 120(2), 325-333. doi: 10.1111/j.1471-4159.2011.07553.x.
- 971 Sperling, R.A., Aisen, P.S., Beckett, L.A., Bennett, D.A., Craft, S., Fagan, A.M., et al. (2011). 972 Toward defining the preclinical stages of Alzheimer's disease: recommendations from the 973 National Institute on Aging-Alzheimer's Association workgroups on diagnostic guidelines 974 Alzheimer's disease. for Alzheimers Dement 7(3), 280-292. doi: 975 10.1016/i.jalz.2011.03.003.
- Terry, R.D., Masliah, E., Salmon, D.P., Butters, N., DeTeresa, R., Hill, R., et al. (1991). Physical basis of cognitive alterations in Alzheimer's disease: synapse loss is the major correlate of cognitive impairment. *Ann Neurol* 30(4), 572-580. doi: 10.1002/ana.410300410.
- Toledo, J.B., Korff, A., Shaw, L.M., Trojanowski, J.Q., and Zhang, J. (2013a). CSF alphasynuclein improves diagnostic and prognostic performance of CSF tau and Abeta in
  Alzheimer's disease. *Acta Neuropathol* 126(5), 683-697. doi: 10.1007/s00401-013-1148z.
- Toledo, J.B., Xie, S.X., Trojanowski, J.Q., and Shaw, L.M. (2013b). Longitudinal change in CSF
  Tau and Abeta biomarkers for up to 48 months in ADNI. *Acta Neuropathol* 126(5), 659670. doi: 10.1007/s00401-013-1151-4.

- Townsend, M., Shankar, G.M., Mehta, T., Walsh, D.M., and Selkoe, D.J. (2006). Effects of
   secreted oligomers of amyloid beta-protein on hippocampal synaptic plasticity: a potent
   role for trimers. *J Physiol* 572(Pt 2), 477-492. doi: 10.1113/jphysiol.2005.103754.
- Viola, K.L., and Klein, W.L. (2015). Amyloid beta oligomers in Alzheimer's disease pathogenesis,
   treatment, and diagnosis. *Acta Neuropathol* 129(2), 183-206. doi: 10.1007/s00401-015 1386-3.
- Viola, K.L., Sbarboro, J., Sureka, R., De, M., Bicca, M.A., Wang, J., et al. (2015). Towards noninvasive diagnostic imaging of early-stage Alzheimer's disease. *Nat Nanotechnol* 10(1),
  91-98. doi: 10.1038/nnano.2014.254.
- Wang, C., Xiong, M., Gratuze, M., Bao, X., Shi, Y., Andhey, P.S., et al. (2021). Selective removal of astrocytic APOE4 strongly protects against tau-mediated neurodegeneration and decreases synaptic phagocytosis by microglia. *Neuron* 109(10), 1657-1674 e1657. doi: 10.1016/j.neuron.2021.03.024.
- Wang, H.W., Pasternak, J.F., Kuo, H., Ristic, H., Lambert, M.P., Chromy, B., et al. (2002).
  Soluble oligomers of beta amyloid (1-42) inhibit long-term potentiation but not long-term depression in rat dentate gyrus. *Brain Res* 924(2), 133-140.
- Wipke, B.T., Wang, Z., Kim, J., McCarthy, T.J., and Allen, P.M. (2002). Dynamic visualization of
   a joint-specific autoimmune response through positron emission tomography. *Nat Immunol* 3(4), 366-372. doi: 10.1038/ni775.
- Xiao, C., Davis, F.J., Chauhan, B.C., Viola, K.L., Lacor, P.N., Velasco, P.T., et al. (2013). Brain
   transit and ameliorative effects of intranasally delivered anti-amyloid-beta oligomer
   antibody in 5XFAD mice. *J Alzheimers Dis* 35(4), 777-788. doi: 10.3233/JAD-122419.
- Yang, T., Dang, Y., Ostaszewski, B., Mengel, D., Steffen, V., Rabe, C., et al. (2019). Target
  engagement in an alzheimer trial: Crenezumab lowers amyloid beta oligomers in
  cerebrospinal fluid. *Ann Neurol* 86(2), 215-224. doi: 10.1002/ana.25513.
- Yang, Y., Kim, J., Kim, H.Y., Ryoo, N., Lee, S., Kim, Y., et al. (2015). Amyloid-beta Oligomers
  May Impair SNARE-Mediated Exocytosis by Direct Binding to Syntaxin 1a. *Cell Rep*12(8), 1244-1251. doi: 10.1016/j.celrep.2015.07.044.
- Yuan, Q., Liu, X., Zhang, Y., Xian, Y.F., Zou, J., Zhang, X., et al. (2021). Established Beta
  Amyloid Pathology Is Unaffected by TREM2 Elevation in Reactive Microglia in an
  Alzheimer's Disease Mouse Model. *Molecules* 26(9). doi: 10.3390/molecules26092685.
- 1017 Zhang, M., Zhong, L., Han, X., Xiong, G., Xu, D., Zhang, S., et al. (2021a). Brain and Retinal
   1018 Abnormalities in the 5xFAD Mouse Model of Alzheimer's Disease at Early Stages. *Front* 1019 *Neurosci* 15, 681831. doi: 10.3389/fnins.2021.681831.
- Zhang, T., Liao, Q., Zhang, D., Zhang, C., Yan, J., Ngetich, R., et al. (2021b). Predicting MCI to
   AD Conversation Using Integrated sMRI and rs-fMRI: Machine Learning and Graph
   Theory Approach. *Front Aging Neurosci* 13, 688926. doi: 10.3389/fnagi.2021.688926.
- 1023Zhao, Y., Wu, X., Li, X., Jiang, L.L., Gui, X., Liu, Y., et al. (2018). TREM2 Is a Receptor for β-1024Amyloid that Mediates Microglial Function. Neuron 97(5), 1023-1031.e1027. doi:102510.1016/j.neuron.2018.01.031.
- 1026 Zhong, L., Xu, Y., Zhuo, R., Wang, T., Wang, K., Huang, R., et al. (2019). Soluble TREM2
  1027 ameliorates pathological phenotypes by modulating microglial functions in an Alzheimer's
  1028 disease model. *Nat Commun* 10(1), 1365. doi: 10.1038/s41467-019-09118-9.
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#### 1033 Figure 1

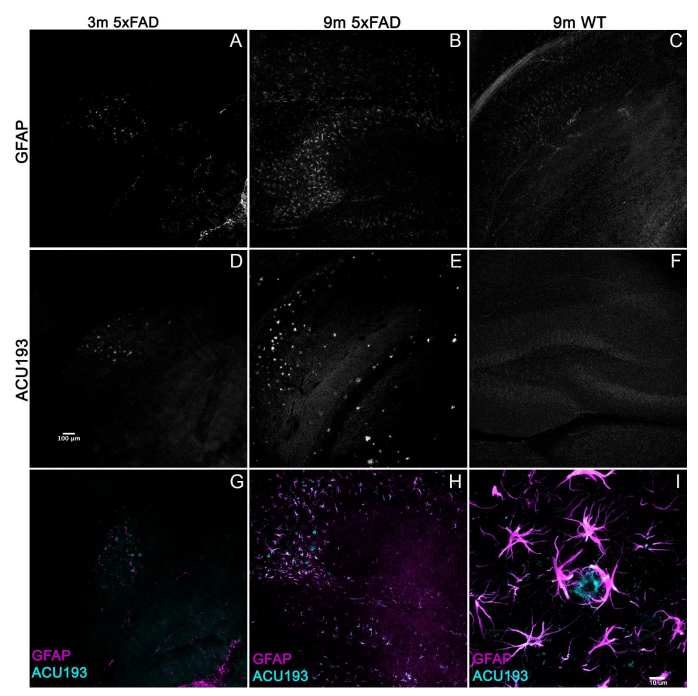
1034 Memory dysfunction in 5xFAD mice is substantial by 4 months and is preceded by A $\beta$ O pathology, 1035 detectable by 3 months of age. (Left) 5xFAD mice and wild-type littermates were assessed for memory 1036 dysfunction using novel location recognition (NLR; hippocampal-dependent task) and novel object 1037 recognition tasks (NOR; cortical-dependent task). Ages ranged 2-12 months. Data shown here are for 1038 the hippocampal-dependent NLR assay (similar results with NOR, not shown). In 5xFAD mice, memory 1039 impairment was negligible at 2-3 months, substantial by 4-5 months, and fully penetrant by 6 months of 1040 age. Statistical analysis shows that there was no significant difference between the behaviors of the WT 1041 mice and the 5xFAD mice at ages 2-3 months, but a statistically significant difference was evident 1042 between the recognition task behaviors of the WT mice and 5xFAD mice for ages 4-5 months (p<0.001) 1043 and 6-7 months (p<0.0001). (Right) Sagittal brain sections were obtained from 5xFAD and WT mice at 1044 ages 2, 3, 4, 6, and 8 months and probed for ABO pathology using a humanized ABO monoclonal 1045 antibody. Fluorescent signal was barely detectable at 2 months of age (not shown), was more readily 1046 detectable by 3 months, and robust by 6 months. Wild-type littermates presented no signal. Scale bar = 1047 100 µm.



#### 1049

#### 1050 Figure 2

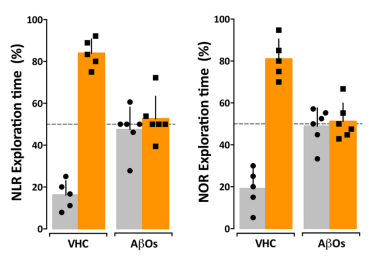
1051 **ACU193 and NU4 detect A\betaOs ex vivo.** Sagittal sections from 9-month-old 5xFAD mice were 1052 immunolabeled with 2 different anti-A $\beta$ O antibodies, NU4 and ACU193, to determine the extent to which 1053 A $\beta$ O pathology is detected by both antibodies. Data show that A $\beta$ Os accumulate and that ACU193 and 1054 NU4 show very similar detection of A $\beta$ Os.



#### 1056

#### 1057 Figure 3

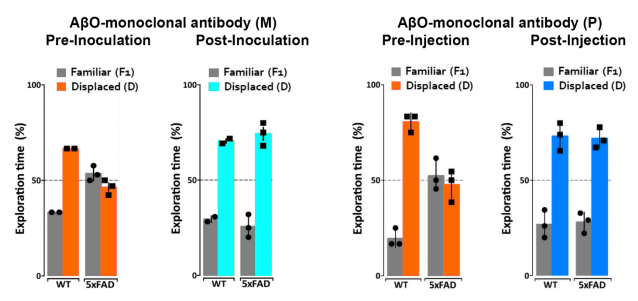
GFAP is activated in a developmental manner in the 5xFAD mouse. Sagittal sections from 5xFAD mice, aged 3-9 months, and their wild-type littermates were immunolabeled with antibodies against GFAP and ACU193, then imaged on the Leica SP5 confocal microscope at 10x and 100x. Data show that, like the ACU193, GFAP positive glial cells accumulate in an age dependent manner. Sale bar = 100 μm for panels A-H ad 10 μm for panel I.



#### 1065 Figure 4

1066 **Intraventricular A\betaO injection caused memory impairment in wild type mice within 24 hours.** Wild 1067 type mice were tested for performance in recognition tasks beginning 24 hours after receiving A $\beta$ O 1068 injections (10 pmols in 3 µI) into the right lateral ventricle. Mice first were assessed for novel location 1069 recognition (NLR; 24 hr post-injection) and subsequently for novel object recognition (NOR; 48 hr post-1070 injection). A $\beta$ O-injected mice were unable to perform either recognition task. Statistical analysis shows 1071 that there is a statistically significant difference between the recognition task behaviors of the WT mice 1072 and the A $\beta$ O injected mice (p<0.0001).

1073

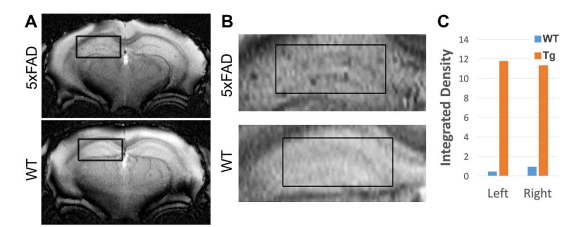


#### 1075 Figure 5

1074

#### 1076 ACUMNS delivered intranasally and ACUPET given iv both rescue memory function in 6- to 7-1077 month-old mice.

1078 Tg and WT mice, aged 6 and 18 months (Left set of bar graphs), were tested by NLR and NOR assays 1079 to ensure predicted behavioral deficits. Mice were then intranasally inoculated with ACUMNS and imaged 1080 for probe distribution and detection of AβO pathology *in vivo*. After imaging, animals were monitored for 1081 30 days for signs of adverse reactions to the probe (none detected), then re-tested by NOR. The 6-1082 month-old animals showed a significant recovery of memory impairment 30 days after inoculation. This 1083 recovery was not observed in the very old animals, although some memory improvement was seen (data 1084 not shown). To test the impact of the ACUPET probe on memory function, Tg and WT mice, aged 7 1085 months, were tested by NLR and NOR assays prior to imaging as before. Mice were then injected, via 1086 tail vein, with ACUPET or non-specific IgGPET and imaged for up to 24 hours to monitor probe 1087 distribution. After imaging, animals were monitored for 40 days for signs of adverse reactions to the 1088 probe. Animals were re-tested by NOR at 40 days recovery. 5xFAD animals injected with ACUPET 1089 showed a persistent recovery of memory impairment that was not seen in the 5xFAD animals injected 1090 with IgGPET. Data support the long-term benefits of these antibody-based probes on memory.

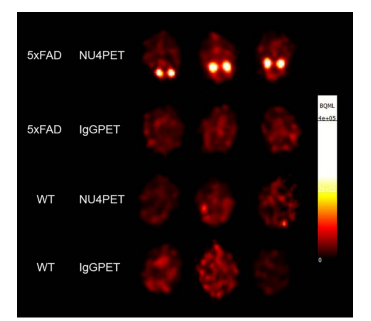


### 1092

### 1093 Figure 6

1094 ACUMNS gives AD-dependent MRI signal in hippocampus of 12-month-old 5xFAD mice.

1095 *In vivo* studies with ACUMNS probe show robust AD-dependent MRI signal in the hippocampus of 12 1096 month-old mice.



#### 1098

#### 1099 Figure 7

#### 1100 **NU4PET** probe gives 5xFAD- specific CNS signal that is maintained even 44 hr after iv injection.

1101 Signal obtained after IV injection of NU4PET showed probe accumulation in the hippocampus of 5xFAD 1102 mice (aged 5-7 months). Controls (IgGPET in AD mice; NU4PET in wild type littermates; IgGPET in wild

1102 The final system and the