Disorders of the Nervous System

**Obesity Accelerates Alzheimer-Related Pathology in APOE4 but not APOE3 Mice**

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**Abstract**

Alzheimer’s disease (AD) risk is modified by both genetic and environmental risk factors, which are believed to interact to cooperatively modify pathogenesis. Although numerous genetic and environmental risk factors for AD have been identified, relatively little is known about potential gene-environment interactions in regulating disease risk. The strongest genetic risk factor for late-onset AD is the e4 allele of apolipoprotein E (APOE4). An important modifiable risk factor for AD is obesity, which has been shown to increase AD risk in humans and accelerate development of AD-related pathology in rodent models. Potential interactions between APOE4 and obesity are suggested by the literature but have not been thoroughly investigated. In the current study, we evaluated this relationship by studying the effects of diet-induced obesity (DIO) in the EFAD mouse model, which combines familial AD transgenes with human APOE3 or APOE4. Male E3FAD and E4FAD mice were maintained for 12 weeks on either a control diet or a Western diet high in saturated fat and sugars. We observed that metabolic outcomes of DIO were similar in E3FAD and E4FAD mice. Importantly, our data showed a significant interaction between diet and APOE genotype on AD-related outcomes in which Western diet was associated with robust increases in amyloid deposits, β-amyloid burden, and glial activation in E4FAD but not in E3FAD mice. These findings demonstrate an important gene-environment interaction in an AD mouse model that suggests that AD risk associated with obesity is strongly influenced by APOE genotype.

**Key words:** Alzheimer’s disease; apolipoprotein E; β-amyloid; gliosis; obesity; transgenic

**Significance Statement**

The e4 allele of apolipoprotein E (APOE4) is the strongest genetic risk factor for Alzheimer’s disease (AD), but not all APOE4 carriers will develop the disease suggesting that APOE genotype interacts with other factors to modulate Alzheimer’s risk. Here, we show that diet-induced obesity (DIO) interacts with APOE4 genotype to increase Alzheimer’s-like pathology in an Alzheimer’s transgenic mouse model that contains human APOE3 versus APOE4 isoforms. Interestingly, mice with APOE3 do not show diet-induced increases in pathology, suggesting that the adverse effects of obesity on Alzheimer’s risk may be limited to APOE4 carriers. These findings identify an important gene-environment interaction that may have significant impact for understanding Alzheimer’s risk and etiology and promoting development of targeted therapeutic approaches that incorporate both obesity and APOE genotype.

**Introduction**

Alzheimer’s disease (AD) is a progressive neurodegenerative disorder, the underlying causes of which are currently incompletely understood. Both genetic and environmental factors are important in determining individual risk for AD. The strongest genetic risk factor for late-
onset AD is the ε4 allele of apolipoprotein E (APOE4; Strittmatter et al., 1993; Liu et al., 2013). In the United States, roughly 12% of the population carries the ε4 allele, but its frequency increases to ~60% in AD patients (Rebeck et al., 1993). APOE4 not only increases risk, but also accelerates the age of onset of AD (Corder et al., 1993; van der Flier et al., 2011). However, since homozygous carriers of APOE4 have a ~50% lifetime risk of AD, a significant number of APOE4 carriers never develop the disease (Genin et al., 2011). Thus, APOE4 likely interacts with other genetic and or environmental factors to drive AD risk.

A significant modifiable risk factor for dementia is obesity. Obesity has numerous adverse neural effects (Lee and Mattson, 2013) and increases the risk of dementia up to three-fold (Whitmer et al., 2008). Body mass index, a commonly used measure of obesity, has been shown to be associated with AD risk (Profenno et al., 2010) as well as with reduced brain volume in AD patients (Ho et al., 2010). Several studies indicate that obesity may be particularly problematic at midlife (Fitzpatrick et al., 2009; Profenno et al., 2010; Meng et al., 2014; Emmerzaal et al., 2015), suggesting that obesity contributes to the development of AD. Similar relationships have been observed in animal models. In particular, diet-induced obesity (DIO) accelerates AD-related pathology in mouse models of AD (Ho et al., 2004; Julien et al., 2010; Kohjima et al., 2010; Barron et al., 2013; Orr et al., 2014). Further, genetic models of obesity and type 2 diabetes exhibit features of AD-like neuropathology (Kim et al., 2009; Jung et al., 2013; Ramos-Rodriguez et al., 2013).

The extent to which APOE4 and obesity interact to regulate AD risk is unclear. Interestingly, APOE4 carriers can be more sensitive to metabolic consequences associated with obesity (de-Andrade et al., 2000; Kyperoos et al., 2009; Niu et al., 2009; Atabek et al., 2012; Zarkeh et al., 2012; Guan et al., 2013). Although some studies do not report an APOE4 bias in obesity-associated AD risk (Profenno and Faroane, 2008; Luchsinger et al., 2012), others have found that AD risk is increased by obesity (Peila et al., 2002; Ghebranious et al., 2011) and diets high in calories and fatty acids (Luchsinger et al., 2002) only in APOE4 carriers. Though the human literature suggests a gene-environment interaction between APOE and obesity in regulating development of AD, this question has not been addressed in experimental models. To study these relationships, we used EFAD transgenic mice, which combine AD transgenes with targeted replacement of mouse APOE with human APOE (Youmans et al., 2012). We compared metabolic and AD-related effects of Western diet in male APOE3 (E3FAD) and APOE4 (E4FAD) mice. Here, we report that DIO increases amyloid pathology and gliosis almost exclusively in E4FAD mice. Our data reveal a gene-environment interaction between APOE genotype and obesity, suggesting that APOE4 carriers may be more susceptible to obesity associated increases in AD risk.

**Materials and Methods**

**Animal procedures**

A colony of EFAD mice, which are heterozygous for the 5xFAD transgenes and homozygous for human APOE3 or APOE4 (Youmans et al., 2012), were maintained at vivarium facilities at University of Southern California from breeder mice generously provided by Dr. Mary Jo LaDu (University of Illinois at Chicago). All animals were housed under a 12-hour light/dark cycle with lights on at 6 A.M. and ad libitum access to food and water. At three months of age, male E3FAD and E4FAD mice were randomized to dietary treatment groups (N = 7–11/group): control diet (10% fat, 7% sucrose; #D12450J Research Diets) or Western diet (45% fat, 17% sucrose; #D12451, Research Diets). EFAD mice were maintained on experimental diets for 12 weeks, an exposure period previously established to yield obesity-induced metabolic impairments in APOE mice (Arbones-Mainar et al., 2010; Segev et al., 2016). Body weight and food consumption were recorded weekly.

At the end of the treatment period, mice were anesthetized with inhalant isoflurane and transcardially perfused with ice-cold 0.1 M PBS. The brains were rapidly removed and immersion fixed for 48 h in 4% paraformaldehyde/0.1 M PBS, then stored at 4°C in 0.1 M PBS/0.3% NaN3 until processed for immunohistochemistry. Gonadal and retroperitoneal fat pads were dissected and weighed as a measure of adiposity, and snap frozen for RNA extraction. All animal procedures were conducted under protocols approved by the University of Southern California Institutional Animal Care and Use Committee and in accordance with National Institute of Health standards.

**Glucose, cholesterol, and triglyceride measurements**

Blood glucose readings were measured after overnight fasting (16 h) every four weeks beginning at week 0 of the 12-week treatment period. Blood was collected from the lateral tail vein and immediately assessed for glucose levels using the Precision Xtra Blood Glucose and Ketone Monitoring System (Abbott Diabetes Care).

Glucose tolerance testing (GTT) was performed at week 11. Fasting, baseline glucose readings were taken after which mice were administered a glucose bolus (2 g/kg body weight) via oral gavage. Blood glucose levels were recorded 15, 30, 60, and 120 min after the glucose bolus was given. Area under the curve (AUC) was calculated.

Plasma cholesterol and triglyceride levels were enzymatically determined at the conclusion of the experiment using commercially available kits (LabAssay Triglycerides #290-63701, Wako Chemicals; Total Cholesterol Colori-
metric Assay kit, #K603, BioVision). All samples were run in duplicate according to manufacturer’s instructions.

**Thioflavin-S (Thio-S) staining and quantification**

Fixed hemi-brains were fully sectioned in the horizontal plane at 40 μm using a vibratome (Leica Biosystems). Every eighth section was stained for Thio-S (#230456, Sigma-Aldrich) using standard methodology. Sections were mounted and allowed to dry overnight, after which they were washed three times in 50% ethanol for 5 min each, then washed in double-distilled H2O before being incubated for 10 min in 1% Thio-S dissolved in H2O. Stained slides were then rinsed in 70% ethanol before being dehydrated and coverslipped in aqueous anti-fade mounting medium (Vector Laboratories). Digital images were captured at 20× magnification using an Olympus BX50 microscope equipped with a DP74 camera and CellSens software (Olympus). The number of spherical thioflavin-positive deposits were counted using NIH ImageJ 1.50i (United States National Institutes of Health) with the cell counter plugin to mark stained plaque-like structures. Thioflavin-positive deposits were counted in entorhinal cortex (three fields/section), subiculum (two fields/section), and hippocampal subfields CA1 (three fields/section) and CA2/3 (three fields/section), across four sections per animal, for a total of ~44 fields per brain.

**Immunohistochemistry**

Immunohistochemistry was performed using a standard avidin/biotin peroxidase approach with ABC Vector Elite kits (Vector Laboratories). Aβ immunohistochemistry was performed on every eighth section using sections immediately adjacent to those processed for Thio-S. Briefly, sections were pretreated with 95% formic acid for 5 min, then rinsed in TBS before being treated with an endogenous peroxidase blocking solution for 10 min. After three 10 min washes in 0.1% Triton-X/TBS, sections were incubated for 30 min in a blocking solution consisting of 2% bovine serum albumin in TBS. Blocked sections were incubated overnight at 4°C in primary antibody directed against Aβ (#71-5800, 1:300 dilution, Invitrogen) that was diluted in blocking solution. Next, sections were rinsed and incubated in biotinylated secondary antibody diluted in blocking solution. Immunoreactivity was visualized using 3,3’-diaminobenzidine (Vector Laboratories). Additional sections were similarly immunostained without formic acid pretreatment using IBA-1 (#019-19741, 1:2000 dilution, Wako) and GFAP (#ab7260, 1:1000 dilution, Abcam).

To quantify the percentage area occupied by Aβ immunoreactivity (Aβ load), images of nonoverlapping fields were taken at 20× magnification in entorhinal cortex (three fields/section), subiculum (three fields/section), and hippocampal subfields CA1 (five fields/section) and CA2/3 (three fields/section) across 4 tissue sections, for a total of ~56 images per brain. Images were digitally captured using an Olympus BX50 microscope and DP74 camera paired with a computer running CellSens software (Olympus). The pictures were converted to grayscale images and thresholded using NIH ImageJ 1.50i to yield binary images separating positive and negative immunostaining.

Aβ load was calculated as the percentage of the total area that was positively immunolabeled.

Microglia and astrocyte activation was quantified using live imaging (Olympus BX50, CASTGrid software, Olympus) at 40× magnification. Each cell was categorized as either resting or reactive based on its morphology, as reported in previous studies (Ayoub and Salm, 2003; Wilhelmsson et al., 2006). Specifically, microglia were scored as resting (type 1) if they had spherical cell bodies, with numerous thin, highly ramified processes. Cells were scored as type 2 cells if they exhibited enlarged rod-shaped cell bodies with fewer processes that were shorter and thicker, and scored type 3 cells if they had very few or no processes or several filopodial processes. Both type 2 and type 3 morphologies were considered an activated microglia phenotype. Astrocytes were visualized with GFAP immunostaining and categorized as exhibiting either nonreactive (normally sized cell bodies with a few rather short projections) or reactive (both cell bodies and projections are enlarged) morphology phenotypes. Entorhinal cortex (four fields/section), subiculum (four fields/section), and hippocampal subfields CA1 (five fields/section) and CA2/3 (three fields/section) were quantified for both microglia and astrocytes. The number of cells across brain regions scored for each animal averaged ~700 microglia and ~600 astrocytes.

**RNA isolation and real-time PCR**

For RNA extractions, gonadal fat pads and hippocampi were homogenized using TRIzol reagent (Invitrogen), following the manufacturer’s protocol. The RNA pellet was treated with RNase-free DNase I (Epigen) for 30 min at 37°C, and a phenol/chloroform extraction was performed to isolate RNA. The iScript cDNA synthesis system (Bio-Rad) was used to reverse transcribe cDNA from 1 μg of purified RNA. Real-time quantitative PCR was performed on the resulting cDNA using SsoAdvanced Universal SYBR Green Supermix (Bio-Rad) and a Bio-Rad CFX Connect Thermocycler. All measurements were performed in duplicates. Quantification of PCR products was conducted by normalizing with a combination of corresponding hypoxanthineguanine phosphoribosyltransferase (HPRT) and succinate dehydrogenase [ubiquinone] flavoprotein subunit, mitochondrial (SDHA) expression levels from the gonadal fat samples, and with β-actin expression levels from hippocampus, using the ΔΔ-CT method to obtain relative mRNA levels. Gonadal fat was probed for levels of cluster differentiation factor 68 (CD68) and EGF-like module-containing mucin-like hormone receptor-like 1 (F4/80), while hippocampus was probed for β-secretase 1 (BACE1), neprylisin, insulin-degrading enzyme (IDE), CD68, glial fibrillary acidic protein (GFAP), and cluster of differentiation factor 74 (CD74). Primer pair sequences are shown in Table 1.

**Statistical analyses**

For the analysis of body weight and glucose tolerance data, two-way repeated measures ANOVAs were run using the Statistical Package for Social Sciences (SPSS; version 23, IBM). All other data were analyzed by two-way ANOVA using Prism (version 5, GraphPad Software). In
the case of significant main effects, planned comparisons between groups of interest were made using the Bonferroni correction. All data are presented as the mean ± SEM. Significance was set at a threshold of \( p < 0.05 \). Statistical results are presented in Tables 2, 3.

### Results

#### Obesity-related outcomes of Western diet

To begin investigating whether there are gene X environment interactions between \( \text{APOE} \) and Western diet, we first compared measures of DIO in E3FAD versus E4FAD mice following the 12-week exposure to control and Western diets. The control diet was associated with \(<1\%\) gain in body weight in both E3FAD and E4FAD mice, whereas Western diet yielded a \( 39 \pm 7.7\% \) increase in body weight in E3FAD and a \( 24 \pm 7.21\% \) increase in E4FAD mice (Fig. 1A), such that the effects of diet did not vary significantly across genotypes (\( p = 0.112; \) Fig. 1A; Table 2). A \( 2 \times 2 \) repeated measures ANOVA revealed a significant main effect of diet on body weight (\( F = 10.51, p = 0.003; \) Fig. 1A) in which Western diet was associated with increased weight. \( \text{APOE} \) genotype did not significantly affect body weight (\( p = 0.759; \) Fig. 1A). Between group comparisons revealed that E3FAD mice fed a Western diet weighed significantly more than E3FAD mice fed a control diet at 4, 8, and 12 weeks (\( p < 0.05 \)). There were no statistically significant differences in body weights at any time point between control and Western diet groups in E4FAD mice.

We next examined plasma levels of cholesterol and triglycerides as measures of adverse effects of Western diet. We found that plasma cholesterol levels were significantly affected by neither genotype (\( p = 0.103 \)) nor diet (\( p = 0.221 \)), and we did not find an interaction effect (\( p = 0.119 \); Fig. 1B; Table 2). Likewise, there were no effects of either genotype (\( p = 0.46 \)) or diet (\( p = 0.102 \)), or an interaction effect (\( p = 0.179 \)) on plasma triglyceride levels (Fig. 1C).

Because metabolic impairments associated with obesity have been linked to adiposity, we assessed fat deposition across groups. We observed a significant interaction effect (\( F = 5.01, p = 0.033; \) Table 2), such that on the control diets, E4FAD mice had more gonadal fat than E3FADs (\( p = 0.027 \)), but there was no difference between E3FAD and E4FAD mice on Western diet (\( p = 0.230; \) Fig. 1D). Additionally, there was a significant main effect of diet (\( F = 37.04, p < 0.001 \)) on weight of the gonadal fat pads, so that both E3FAD and E4FAD mice had increased fat pads with Western diet (Fig. 1D). Parallel findings were observed in the retroperitoneal fat pads (data not shown). Because inflammation is an established hallmark of obesity, we examined gene expression of the macrophage markers CD68 and F4/80 by PCR in the adipose tissue. We found a significant main effect of diet on CD68 expression (\( F = 11.54, p = 0.003 \), although this effect reached statistical significance only in E3FAD but not in E4FAD mice (Fig. 1E). There was no statistically significant effect of genotype (\( p = 0.353 \)), nor was there an interaction between diet and genotype (\( p = 0.366 \) on CD68 expression. Diet had a main effect on adipose F4/80 expression (\( F = 7.02, p = 0.015 \)), and again, this effect reached statistical significance only in E3FAD mice (Fig. 1F). There was no statistically significant effect of genotype (\( p = 0.768 \)), and no interaction effect (\( p = 0.288 \)) on F4/80 expression (Table 2).

In addition to increasing body weight and adiposity, Western diet can induce metabolic impairments including dysregulation of glucose homeostasis. When examining glucose clearance in the GTT, we found a significant main effect of diet (\( F = 5.03, p = 0.033 \)), such that both E3FAD and E4FAD mice fed a Western diet were impaired at clearing glucose (Fig. 1G; Table 2). There was no main effect of genotype (\( p = 0.886 \)), or interaction effect between diet and genotype (\( p = 0.750 \)) on glucose clearance. We also calculated the area under the curve (AUC) for GTT, and found that there was a significant main effect of diet (\( F = 5.73, p = 0.023 \)), but not of genotype (\( p = 0.817 \)) on GTT AUC (Fig. 1H). However, the effect of diet failed to reach statistical significance when examined separately in E3FAD and E4FAD mice. There was no interaction between genotype and diet on GTT AUC (\( p = 0.737 \)). Changes in fasting glucose levels over the diet treatment period showed a trend toward a main effect of diet (\( F = 3.84, p = 0.059; \) Fig. 1I). There was no effect of genotype (\( p = 0.371 \)) nor was there an interaction between diet and genotype (\( p = 0.352 \)) on changes in glucose levels (Table 2).

**Western diet increases \( \beta \)-amyloid deposition in E4FAD but not in E3FAD mice**

The primary AD-related neuropathological change in EFAAD mice at this age is accumulation of \( \beta \)-amyloid protein, largely in the form of extracellular deposits, many of which exhibit positive Thio-S staining that is indicative of amyloid. Thus, to begin assessing AD-related neuropathology, Thio-S positive plaques were counted in entorhi-

| Target gene | Sequence |
|-------------|----------|
| CD68        | Forward: 5’-TTCTGCTTGAAAATGCAAG-3’  
Reverse: 5’-AGAAAAAGGCACAAGCGGAGA-3’ |
| F4/80       | Forward: 5’-TGCACTACAGAAGGACAGC-3’  
Reverse: 5’-GCCCTGCTGTTGTTTTGCT-3’ |
| HPRT        | Forward: 5’-AGGTTGTGTTGGAAAAAGGA-3’  
Reverse: 5’-TTGGGTCTGTCATTGGGCT-3’ |
| SDHA        | Forward: 5’-GACAGACCTGTTGGAAGC-3’  
Reverse: 5’-GGATGGTGGTTGGAGTAT-3’ |
| Neprilysin  | Forward: 5’-GAAAAAAGGCACTTGTTG-3’  
Reverse: 5’-GAAAAAGCAGATTGGCGAG-3’ |
| BACE1       | Forward: 5’-TGGTTGGTTGGAAAAAGGA-3’  
Reverse: 5’-ACCCTGCTGTTGTTTTGCT-3’ |
| IDE         | Forward: 5’-GTCTTCCACACAGCAGGT-3’  
Reverse: 5’-CTCTTCTGGTTGCTGCTTG-3’ |
| CD74        | Forward: 5’-AAATCGCGCAACAGGTGAC-3’  
Reverse: 5’-GCACTTGTGGTCACTTTATG-3’ |
| GFAP        | Forward: 5’-AACGGATATGCGCGCAACAGGT-3’  
Reverse: 5’-CTCTTCTGGTTGCTGCTTG-3’ |
| \( \beta \)-Actin | Forward: 5’-AGGCATTGAGTACGATCCAC-3’  
Reverse: 5’-CTTCTGACTGTGGTTGTA-3’ |

Table 1. Gene targets for the PCR analyses are listed with their corresponding oligonucleotide sequences for the forward and reverse primers.
| Figure | Statistical test | Description | Significance |
|--------|-----------------|-------------|--------------|
| Figure 1A | Kolmogorov-Smirnov test for normality (p value) | Body weight | All groups at all time points are normally distributed (p > 0.05). |
| Figure 1B | | Plasma cholesterol | E3FAD CTL > 0.10, E3FAD WD > 0.10, E4FAD CTL > 0.10, E4FAD WD > 0.10. |
| Figure 1C | | Plasma triglycerides | E3FAD CTL > 0.10, E3FAD WD > 0.10, E4FAD CTL > 0.10, E4FAD WD > 0.10. |
| Figure 1D | | Gonadal fat weight | E3FAD CTL > 0.10, E3FAD WD > 0.10, E4FAD CTL > 0.10, E4FAD WD > 0.10. |
| Figure 1E | | CD68 | E3FAD CTL N/A, E3FAD WD > 0.10, E4FAD CTL > 0.10, E4FAD WD > 0.10. |
| Figure 1F | | F4/80 | E3FAD CTL N/A, E3FAD WD > 0.10, E4FAD CTL > 0.10, E4FAD WD > 0.10. |
| Figure 1G | | Glucose (GTT) | E3FAD WD 0 min = 0.002, E4FAD WD 15 min = 0.025, E3FAD WD 30 min = 0.011, E4FAD WD 30 min = 0.008. |
| Figure 1H | | GTT AUC | E3FAD CTL = 0.07, E3FAD WD = 0.097, E4FAD CTL = 0.10, E4FAD WD = 0.033. |
| Figure 1I | | Percent glucose change | E3FAD CTL > 0.10, E3FAD WD > 0.10, E4FAD CTL > 0.10, E4FAD WD > 0.10. |
| Figure 2B | | Thio-S: entorhinal cortex | E3FAD WD = 0.049, E4FAD CTL > 0.10, E4FAD WD > 0.10. |
| Figure 2C | | Thio-S: subiculum | E3FAD WD = 0.10, E4FAD CTL > 0.10, E4FAD WD > 0.10. |
| Figure 2D | | Thio-S: CA1 | E3FAD WD = 0.10, E4FAD CTL > 0.10, E4FAD WD > 0.10. |
| Figure 2E | | Thio-S: CA2/3 | E3FAD CTL > 0.10, E3FAD WD > 0.10, E4FAD CTL > 0.10, E4FAD WD > 0.10. |
| Figure 3B | | Aβ load: entorhinal cortex | E3FAD WD = 0.002, E4FAD CTL > 0.10, E4FAD WD > 0.10. |
| Figure 3C | | Aβ load: subiculum | E3FAD CTL > 0.10, E3FAD WD > 0.10, E4FAD CTL > 0.10, E4FAD WD > 0.10. |

(Continued)
Table 2. Continued

| Figure | Kolmogorov-Smirnov test for normality (p value) | Statistical significance |
|--------|-----------------------------------------------|-------------------------|
| **Figure 3D**<br>Alβ load: CA1 | E3FAD CTL > 0.10<br>E3FAD WD > 0.10<br>E4FAD CTL > 0.10<br>E4FAD WD > 0.036 | Genotype $F_{(1,29)} = 37.66, p < 0.001$<br>diet: $F_{(1,29)} = 2.91, p = 0.099$<br>interaction: $F_{(1,29)} = 2.71, p = 0.110$ |
| **Figure 3E**<br>Alβ load: CA2/3 | E3FAD CTL > 0.10<br>E3FAD WD > 0.10<br>E4FAD CTL > 0.10<br>E4FAD WD > 0.10 | Genotype $F_{(1,29)} = 47.27, p < 0.001$<br>diet: $F_{(1,29)} = 10.36, p = 0.003$<br>interaction: $F_{(1,29)} = 4.48, p = 0.043$ |
| **Figure 4B**<br>microglia number: entorhinal cortex | E3FAD CTL > 0.10<br>E3FAD WD > 0.10<br>E4FAD CTL > 0.10<br>E4FAD WD > 0.10 | Genotype $F_{(1,27)} = 9.78, p = 0.004$<br>diet: $F_{(1,27)} = 2.31, p = 0.141$<br>interaction: $F_{(1,27)} = 1.05, p = 0.316$ |
| **Figure 4C**<br>microglia number: subiculum | E3FAD CTL > 0.10<br>E3FAD WD > 0.10<br>E4FAD CTL > 0.10<br>E4FAD WD > 0.10 | Genotype $F_{(1,27)} = 42.77, p < 0.001$<br>diet: $F_{(1,27)} = 4.20, p = 0.050$<br>interaction: $F_{(1,27)} = 4.75, p = 0.038$ |
| **Figure 4D**<br>microglia number: CA1 | E3FAD CTL > 0.10<br>E3FAD WD > 0.10<br>E4FAD CTL > 0.10<br>E4FAD WD > 0.10 | Genotype $F_{(1,27)} = 51.42, p < 0.001$<br>diet: $F_{(1,27)} = 10.78, p = 0.003$<br>interaction: $F_{(1,27)} = 7.97, p = 0.009$ |
| **Figure 4E**<br>microglia number: CA2/3 | E3FAD CTL > 0.10<br>E3FAD WD > 0.10<br>E4FAD CTL > 0.10<br>E4FAD WD > 0.10 | Genotype $F_{(1,27)} = 21.64, p < 0.001$<br>diet: $F_{(1,27)} = 1.97, p = 0.172$<br>interaction: $F_{(1,27)} = 1.90, p = 0.180$ |
| **Figure 4F**<br>microglia reactivity: entorhinal cortex | E3FAD CTL > 0.10<br>E3FAD WD > 0.10<br>E4FAD CTL > 0.10<br>E4FAD WD > 0.10 | Genotype $F_{(1,27)} = 109.10, p < 0.001$<br>diet: $F_{(1,27)} = 1.64, p = 0.212$<br>interaction: $F_{(1,27)} = 5.52, p = 0.027$ |
| **Figure 4G**<br>microglial reactivity: subiculum | E3FAD CTL > 0.10<br>E3FAD WD > 0.10<br>E4FAD CTL > 0.07<br>E4FAD WD < 0.001 | Genotype $F_{(1,27)} = 19.70, p < 0.001$<br>diet: $F_{(1,27)} = 0.00, p = 0.995$<br>interaction: $F_{(1,27)} = 0.51, p = 0.480$ |
| **Figure 4H**<br>microglial reactivity: CA1 | E3FAD CTL > 0.10<br>E3FAD WD > 0.10<br>E4FAD CTL > 0.10<br>E4FAD WD = 0.04 | Genotype $F_{(1,27)} = 78.70, p < 0.001$<br>diet: $F_{(1,27)} = 5.00, p = 0.034$<br>interaction: $F_{(1,27)} = 11.58, p = 0.002$ |
| **Figure 4L**<br>microglial reactivity: CA2/3 | E3FAD CTL > 0.10<br>E3FAD WD > 0.10<br>E4FAD CTL > 0.10<br>E4FAD WD > 0.10 | Genotype $F_{(1,27)} = 165.70, p < 0.001$<br>diet: $F_{(1,27)} = 21.04, p < 0.001$<br>interaction: $F_{(1,27)} = 32.66, p < 0.001$ |
| **Figure 5B**<br>astrocyte number: entorhinal cortex | E3FAD CTL > 0.10<br>E3FAD WD > 0.10<br>E4FAD CTL > 0.10<br>E4FAD WD > 0.10 | Genotype $F_{(1,29)} = 3.82, p = 0.060$<br>diet: $F_{(1,29)} = 0.29, p = 0.593$<br>interaction: $F_{(1,29)} = 0.41, p = 0.528$ |
| **Figure 5C**<br>astrocyte number: subiculum | E3FAD CTL > 0.10<br>E3FAD WD > 0.10<br>E4FAD CTL > 0.10<br>E4FAD WD > 0.10 | Genotype $F_{(1,29)} = 9.95, p = 0.004$<br>diet: $F_{(1,29)} = 4.79, p = 0.037$<br>interaction: $F_{(1,29)} = 1.04, p = 0.316$ |
| **Figure 5D**<br>astrocyte number: CA1 | E3FAD CTL > 0.10<br>E3FAD WD > 0.10<br>E4FAD CTL > 0.10<br>E4FAD WD > 0.10 | Genotype $F_{(1,29)} = 5.88, p = 0.022$<br>diet: $F_{(1,29)} = 3.55, p = 0.069$<br>interaction: $F_{(1,29)} = 0.49, p = 0.489$ |
| **Figure 5E**<br>astrocyte number: CA2/3 | E3FAD CTL > 0.10<br>E3FAD WD > 0.10<br>E4FAD CTL > 0.10<br>E4FAD WD > 0.10 | Genotype $F_{(1,29)} = 1.82, p = 0.188$<br>diet: $F_{(1,29)} = 4.26, p = 0.048$<br>interaction: $F_{(1,29)} = 0.02, p = 0.894$ |
| **Figure 5F**<br>astrocyte reactivity: entorhinal cortex | E3FAD CTL > 0.10<br>E3FAD WD = 0.004<br>E4FAD CTL > 0.10<br>E4FAD WD > 0.10 | Genotype $F_{(1,29)} = 46.97, p < 0.001$<br>diet: $F_{(1,29)} = 5.75, p = 0.023$<br>interaction: $F_{(1,29)} = 4.82, p = 0.036$ |
| **Figure 5G**<br>astrocyte reactivity: subiculum | E3FAD CTL > 0.10<br>E3FAD WD > 0.10<br>E4FAD CTL = 0.045<br>E4FAD WD > 0.10 | Genotype $F_{(1,29)} = 27.72, p < 0.001$<br>diet: $F_{(1,29)} = 3.13, p = 0.088$<br>interaction: $F_{(1,29)} = 0.00, p = 0.989$ |
nal cortex and in subregions of the hippocampus. Visual inspection of stained sections qualitatively showed not only the expected increase in amyloid deposits in E4FAD mice, but also the surprising finding that Western diet increased Thio-S positive plaques only in E4FAD mice (Fig. 2A). Specifically, there were significant interaction effects between genotype and diet on Thio-S positive plaques in subiculum ($F = 9.75$, $p = 0.004$; Fig. 2C), CA1 ($F = 8.41$, $p = 0.007$; Fig. 2D), and CA2/3 ($F = 7.32$, $p = 0.011$; Fig. 2E), and a nonsignificant trend toward an interaction in entorhinal cortex ($F = 4.09$, $p = 0.053$; Fig. 2B; Table 2). Further analyses revealed that diet significantly increased Thio-S positive plaque counts in E4FAD but not E3FAD males across all brain regions sampled ($p < 0.01$). Additionally, there was a significant main effect of genotype even in the absence of diet, such that E4FAD mice had a greater number of Thio-S positive plaques in entorhinal cortex ($F = 50.30$, $p < 0.001$; Fig. 2B), subiculum ($F = 59.40$, $p < 0.001$; Fig. 2C), CA1 ($F = 80.58$, $p < 0.001$; Fig. 2D), and CA2/3 ($F = 46.39$, $p < 0.001$; Fig. 2E) than did E3FAD mice.

As a second measure of AD-like pathology, we assessed total $\beta$-amyloid burden by immunohistochemistry. This provides a measure of complete $\beta$-amyloid, as the antibody recognizes intra- and extracellular accumulations of $\beta\beta$, even those that have not progressed to Thio-S positive amyloid deposits. Results repeated the same general pattern observed with Thio-S staining. That is, (1) E4FAD mice exhibit greater $\beta$-amyloid burden, and (2) E4FAD but not E3FAD mice show increased $\beta$-amyloid accumulation with Western diet (Fig. 3A). We found significant interaction effects between genotype and diet in entorhinal cortex ($F = 4.91$, $p = 0.035$; Fig. 3B) and in CA2/3 ($F = 4.48$, $p = 0.043$; Fig. 3E), but not in subiculum ($F = 0.11$, $p = 0.742$; Fig. 3C) or in CA1 ($F = 2.71$, $p = 0.110$; Fig. 3D; Table 2). Bonferroni post hoc tests showed that Western diet significantly increased $\beta\beta$ load in E4FAD but not in E3FAD mice across all brain regions surveyed.

### Table 3. Relative gene expression in hippocampus

| Gene | Mean ± SEM | Kolmogorov-Smirnov test for normality ($p$ value) | Statistical significance |
|------|------------|--------------------------------------------------|-------------------------|
|      |            | E3FAD CTL > 1 ± N/A                             | Genotype $F_{(1,28)} = 1.10$, $p = 0.304$ |
|      |            | E3FAD WD > 1.53 ± 0.31                           | diet $F_{(1,28)} = 3.44$, $p = 0.074$ |
|      |            | E4FAD CTL > 1.32 ± 0.19                          | interaction $F_{(1,28)} = 0.03$, $p = 0.874$ |
|      |            | E4FAD WD > 1.76 ± 0.41                           | Genotype $F_{(1,28)} = 0.02$, $p = 0.902$ |
|      |            | E4FAD CTL > 1.61 ± 0.79                          | diet $F_{(1,28)} = 2.49$, $p = 0.126$ |
|      |            | E4FAD WD > 1.94 ± 0.30                           | interaction $F_{(1,28)} = 0.06$, $p = 0.802$ |
|      |            | E4FAD CTL > 1 ± N/A                              | Genotype $F_{(1,28)} = 0.08$, $p = 0.785$ |
|      |            | E4FAD WD > 1.27 ± 0.39                           | diet $F_{(1,28)} = 0.00$, $p = 0.955$ |
|      |            | E4FAD CTL > 1 ± N/A                              | interaction $F_{(1,28)} = 0.49$, $p = 0.489$ |
|      |            | E4FAD WD > 1.12 ± 0.35                           | Genotype $F_{(1,28)} = 10.75$, $p = 0.003$ |
|      |            | E3FAD CTL > 1 ± N/A                              | diet $F_{(1,28)} = 1.91$, $p = 0.178$ |
|      |            | E3FAD WD > 1.21 ± 0.29                           | interaction $F_{(1,28)} = 0.40$, $p = 0.532$ |
|      |            | E4FAD CTL > 1.74 ± 0.30                           | Genotype $F_{(1,28)} = 14.26$, $p < 0.001$ |
|      |            | E4FAD WD > 2.30 ± 0.29                           | diet $F_{(1,28)} = 0.23$, $p = 0.634$ |
|      |            | E3FAD CTL > 1 ± N/A                              | interaction $F_{(1,28)} = 0.14$, $p = 0.712$ |
|      |            | E3FAD WD > 1.02 ± 0.11                           | Genotype $F_{(1,28)} = 16.98$, $p < 0.001$ |
|      |            | E4FAD CTL > 1.56 ± 0.21                           | diet $F_{(1,28)} = 1.86$, $p = 0.184$ |
|      |            | E4FAD WD > 2.70 ± 0.04                           | interaction $F_{(1,28)} = 0.96$, $p = 0.335$ |
|      |            | E3FAD CTL > 1 ± N/A                              | Genotype $F_{(1,28)} = 23.82$, $p < 0.001$ |
|      |            | E4FAD CTL > 1.61 ± 0.21                           | interaction $F_{(1,28)} = 2.08$, $p = 0.160$ |
|      |            | E3FAD WD > 1.02 ± 0.11                           | Genotype $F_{(1,28)} = 11.68$, $p = 0.002$ |
|      |            | E4FAD CTL > 1.61 ± 0.21                           | diet $F_{(1,28)} = 2.83$, $p = 0.099$ |
|      |            | E4FAD WD > 1.02 ± 0.11                           | interaction $F_{(1,28)} = 2.405$, $p = 0.132$ |

Data are presented as mean fold differences (±SEM) relative to E3FAD mice on a control diet. The Kolmogorov-Smirnov test for normality was performed, with $p > 0.05$ indicating a normal distribution. Genes related to $\beta$-amyloid production (BACE-1) and clearance (neprilysin, IDE) showed no significant changes with either diet or genotype, while genes related to glial activation (CD68, GFAP, and CD74) were increased in E4FAD mice on both control and Western diets.
There was a significant main effect of genotype with E4FAD mice having greater Aβ load than E3FAD mice in entorhinal cortex ($F = 21.38$, $p < 0.001$; Fig. 3B), subiculum ($F = 25.40$, $p < 0.001$; Fig. 3C), CA1 ($F = 37.66$, $p < 0.001$; Fig. 3D), and CA2/3 ($F = 47.27$, $p < 0.001$; Fig. 3E).

Western diet increases gliosis more strongly in E4FAD than in E3FAD mice

Gliosis is an important neuropathological feature of AD that is also associated with both obesity and APOE4. To assess gliosis, we compared both the relative cell numbers and morphologic activation state of microglia and astrocytes across groups. We found that, in comparison to E3FAD mice, E4FAD mice consistently had a higher total number of glial cells as well as a higher percentage of glial cells with reactive versus resting phenotypes. Moreover, the effects of diet on glial number and reactivity were stronger in E4FAD than in E3FAD mice.

We first examined microglia number and morphology by IBA-1 staining. Figure 4A shows a resting microglial cell with thin, ramified processes (type 1), and activated cells with rod-shaped cell bodies and fewer, thicker processes (type 2), and amoeboid cells (type 3). We found significant interactions between genotype and diet when examining the total number of microglia per mm$^2$ in subiculum ($F = 4.75$, $p = 0.038$; Fig. 4C) and in CA1 ($F = 7.97$, $p = 0.009$; Fig. 4D), with Bonferroni post hoc tests showing that Western diet increased microglia number in E4FAD but not in E3FAD mice in these brain regions ($p < 0.05$; Table 2). There were no interaction effects on microglia number in entorhinal cortex ($p = 0.316$; Fig. 4B), or in CA2/3 ($p = 0.180$; Fig. 4E). There was a significant effect of genotype on the total number of microglia per
mm² in entorhinal cortex ($F = 9.78, p = 0.004; \text{Fig. 4B}$), subiculum ($F = 42.77, p < 0.001; \text{Fig. 4C}$), CA1 ($F = 51.42, p < 0.001; \text{Fig. 4D}$), and CA2/3 ($F = 21.64, p < 0.001; \text{Fig. 4E}$), such that E4FAD mice had a greater total number of microglia across these brain regions than did E3FAD mice. However, in entorhinal cortex, the effect of genotype was significant only in animals on a Western diet.

Measures of microglial reactivity showed similar results as microglial number. Significant interaction effects between genotype and diet were observed in entorhinal cortex ($F = 5.52, p = 0.027; \text{Fig. 4F}$), CA1 ($F = 11.58, p = 0.002; \text{Fig. 4H}$), and CA2/3 ($F = 32.66, p < 0.001; \text{Fig. 4I}$), but not in subiculum ($p = 0.480; \text{Fig. 4G; Table 2}$). Bonferroni post hoc tests revealed that Western diet increased the percentage of reactive microglia in entorhinal cortex, CA1, and CA2/3 of E4FAD, but not E3FAD, male mice. There was a significant main effect of genotype even in the absence of diet, such that E4FAD mice had a greater percentage of reactive microglia than E3FAD mice.

**Figure 2.** Accumulation of amyloidogenic deposits assessed by Thio-S staining in E3FAD and E4FAD mice across dietary treatments. A, Representative images of Thio-S staining in the subiculum of E3FAD and E4FAD males fed control and Western diets. Scale bar, 50 µm. Numbers of Thio-S positive plaque numbers in E3FAD and E4FAD mice maintained on control and Western diets were quantified in B entorhinal cortex, and hippocampal subregions (C) subiculum, (D) CA1, and (E) CA2/3. Data are presented as mean (±SEM) values; $n = 7–11/group$. E3FAD mice are shown as circles, E4FAD mice are shown as squares; control diet groups are indicated as open symbols, and Western diet groups as filled symbols. *, $p < 0.05$ relative to genotype-matched mice in control diet condition. #, $p < 0.05$ relative to E3FAD mice in same diet condition.
in entorhinal cortex ($F = 109.10$, $p < 0.001$; Fig. 4F), subiculum ($F = 19.70$, $p < 0.001$; Fig. 4G), CA1 ($F = 78.70$, $p < 0.001$; Fig. 4H), and CA2/3 ($F = 165.70$, $p < 0.001$; Fig. 4I).

We next examined astrocyte number and activation by GFAP staining. Figure 5A shows examples of a nonreactive astrocyte with a normally sized soma versus a reactive phenotype with enlarged soma and projections. For the measure of astrocyte number, the effects of diet did not differ across genotype for any of the brain regions sampled (Table 2). We found significant main effects of genotype on the total number of astrocytes in subiculum ($F = 9.95$, $p = 0.004$; Fig. 5C), although this effect was only statistically significant in animals on a Western diet. There was a main effect of genotype on astrocyte number in CA1 ($F = 5.88$, $p = 0.022$; Fig. 5D), but this did not reach statistical significance when examined separately in control and Western diet-fed animals. There was a trend
Figure 4. Microglia number and morphologic status assessed by IBA-1 immunohistochemistry in E3FAD and E4FAD mice across dietary treatments. A, Representative images of microglial morphology associated with resting (type 1) and reactive (types 2 and 3) phenotypes. Scale bar, 40 μm. B–E, Densities (cells/mm²) of IBA-1-immunoreactive cells in E3FAD and E4FAD mice on control and Western diets were quantified in (B) entorhinal cortex, and hippocampal subregions (C) subiculum, (D) CA1, and (E) CA2/3. F–I) Percentages of all IBA-1-immunoreactive cells scored as having reactive phenotype (types 2 and 3) were quantified in (F) entorhinal cortex, and hippocampal subregions (G) subiculum, (H) CA1, and (I) CA2/3. Data are presented as mean (±SEM) values; n = 7–11/group. E3FAD mice are shown as circles, E4FAD mice are shown as squares; control diet groups are indicated as open symbols.
and Western diet groups as filled symbols. *, p < 0.05 relative to genotype-matched mice in control diet condition. #, p < 0.05 relative to E3FAD mice in same diet condition. toward a significant effect of genotype in entorhinal cortex ($F = 3.82, p = 0.060$; Fig. 5B), but no effect in CA2/3 ($p = 0.188$; Fig. 5E). Diet had significant main effects on astrocyte number in subiculum ($F = 4.79, p = 0.037$; Fig. 5C), and CA2/3 ($F = 4.26, p = 0.048$; Fig. 5E), with a trend toward a main effect in CA1 ($F = 3.55, p = 0.069$; Fig. 5D), although this effect did not reach statistical significance when examined separately in E3FAD and E4FAD mice in any brain region. There was no effect of diet on astrocyte number in entorhinal cortex ($p = 0.593$; Fig. 5B).

When examining astrocyte reactivity, we found similar trends as with microglial reactivity. That is, there was a significant interaction effect between genotype and diet on astrocyte reactivity in entorhinal cortex ($F = 4.82, p = 0.036$; Fig. 5F), with Western diet increasing reactivity only in E4FAD mice (Table 2). There were no significant interaction effects between genotype and diet in subiculum ($p = 0.989$; Fig. 5G), CA1 ($p = 0.160$; Fig. 5H), or CA2/3 ($p = 0.132$; Fig. 5I). Moreover, in the absence of diet, genotype had a significant effect on astrocyte reactivity, with E4FAD mice having a greater percentage of reactive astrocytes in entorhinal cortex ($F = 46.97, p < 0.001$; Fig. 5I), subiculum ($F = 27.72, p < 0.001$; Fig. 5G), CA1 ($F = 87.49, p < 0.001$; Fig. 5H), and CA2/3 ($F = 11.68, p = 0.002$; Fig. 5I). In CA2/3 the effect of genotype was only significant in Western diet-fed animals. Furthermore, Western diet significantly increased astrocyte reactivity in CA1 ($F = 23.82, p < 0.001$; Fig. 5H), and CA2/3 ($F = 7.83, p = 0.009$; Fig. 5I), although this effect was only significant in E4FAD mice in CA2/3. There was a nonsignificant trend toward an effect of diet in subiculum ($F = 3.13, p = 0.088$; Fig. 5G).

### E4FAD mice have increased gene expression of inflammatory markers

To begin addressing possible mechanisms underlying the interactive effects of APOE4 and Western diet, we examined hippocampal gene expression of several markers related to Aβ production and clearance, as well as inflammation. Overall, our results indicate that gene expression of factors involved in Aβ clearance and production are not significantly altered by genotype or diet, and that inflammatory gene expression is increased in E4FAD mice, without being altered by Western diet (Table 3).

For BACE1, relative mRNA levels did not show evidence of an interaction between the diet and APOE genotypes ($p = 0.874$). There was no significant main effect genotype ($p = 0.304$), but there was a nonsignificant trend of increased BACE1 levels with Western diet ($p = 0.074$). Expression of the Aβ clearance factor neprilysin was not significantly affected by genotype ($p = 0.902$) or diet ($p = 0.126$), and there was no interaction between genotype and diet ($p = 0.802$). Likewise, gene expression of IDE was not altered by genotype ($p = 0.785$), diet ($p = 0.955$), or the interaction between genotype and diet ($p = 0.489$).

In assessing gene expression of inflammatory markers we found that E4FAD mice had significantly greater levels of the microglial markers CD68 ($F = 10.75, p = 0.003$), the astrocyte marker GFAP ($F = 14.26, p < 0.001$), and the innate immune marker CD74 ($F = 16.98, p < 0.001$), than did E3FAD mice. However, there were no significant effects of diet on levels of CD68 ($p = 0.178$), GFAP ($p = 0.634$), or CD74 ($p = 0.184$). Moreover, there were no significant interactions between genotype and diet on levels of CD68 ($p = 0.532$), GFAP ($p = 0.712$), or CD74 ($p = 0.335$).

### Discussion

The goal of this study is to examine whether APOE genotype and obesity interact to promote AD pathogenesis. Comparing E3FAD and E4FAD mice maintained on standard versus Western diets, we demonstrate a significant gene-environment interaction whereby DIO drives AD-related pathology primarily in APOE4 mice. Our results are consistent with previous findings in humans (Fitzpatrick et al., 2009; Provenno et al., 2010), and confirm studies in rodent models (Ho et al., 2004; Julien et al., 2010; Kohjima et al., 2010; Barron et al., 2013) that obesity increases risk for development of AD. Similarly, our findings replicate prior rodent data (Fryer et al., 2005; Castellano et al., 2011; Youmans et al., 2012; Rodriguez et al., 2014; Cacciottolo et al., 2016) that model the human observation that APOE4 increases the risk and or accelerates the onset of AD pathology (Corder et al., 1993; Saunders et al., 1993; Strittmatter et al., 1993; Morris et al., 2010; Jack et al., 2015). Importantly, our data indicate that the effects of DIO and APOE4 are not strictly additive. Although APOE4 status is associated with greater AD-like pathology on both control and Western diets, obesity increased AD-like pathology in E4FAD but not E3FAD mice. Our finding that E3FAD mice did not show a diet-induced increase in AD-related pathology is similar to null findings in some rodent models of obesity (Zhang et al., 2013; Knight et al., 2014; Niedowicz et al., 2014), suggesting that deleterious effects of obesity can be regulated by genetic factors besides APOE4. Thus, these data suggest an important gene X environment interaction in which APOE4 carriers are more susceptible to the AD-promoting effects of obesity.

How neural outcomes in human populations are impacted by the relationship between APOE genotype and metabolic risk factors remains incompletely defined. Many studies simply control for APOE genotype rather than considering its potential moderating role in the relationship between obesity and AD risk (Vanhanen et al., 2006; Luchsinger et al., 2012). When APOE status has been considered as a modulator of AD risk associated with metabolic factors, the results have been mixed. In some studies, APOE4 carriers showed significantly more cognitive impairment in association with adverse metabolic conditions including atherosclerosis, peripheral vascular disease, type 2 diabetes (Haan et al., 1999), and high systolic blood pressure at midlife (Peila et al., 2001).
Figure 5. Astrocyte number and morphologic status assessed by GFAP immunohistochemistry in E3FAD and E4FAD mice across dietary treatments. 

A, Representative images of astrocyte morphology associated with resting and reactive phenotypes. Scale bar, 50 μm. B–E, Densities (cells/mm²) of GFAP-immunoreactive cells in E3FAD and E4FAD mice on control and Western diets were quantified in (B) entorhinal cortex, and hippocampal subregions (C) subiculum, (D) CA1, and (E) CA2/3. F–I, Percentages of all GFAP-immunoreactive cells scored as having reactive phenotype (type 2) were quantified in (F) entorhinal cortex, and hippocampal...
Further, levels of senile plaques and neurofibrillary tangles were highest in obese men that were also APOE4 carriers (Peila et al., 2002). However, several other studies reported that the AD risk associated with obesity and metabolic syndrome is stronger in APOE3 carriers (Dixit et al., 2005; Leiva et al., 2005; Singh et al., 2006; Profenno and Faraone, 2008).

An important consideration in interpreting these seemingly discordant findings is the potential role of sex differences. Although the impact of sex differences in the interactions among obesity, APOE, and AD risk has not been thoroughly addressed, AD is characterized by numerous sex differences (Li and Singh, 2014; Pike, 2017). Further, the AD-associated risk of APOE4 appears to disproportionately affect women (Payami et al., 1994; Farrer et al., 1997; Altmann et al., 2014). Additionally, there are sex differences in various aspects of obesity (Lovejoy et al., 2009; Mauvais-Jarvis, 2015; Moser and Pike, 2016), including observations that women exhibit relative protection against obesity until menopause (Meyer et al., 2011; Sugiyama and Agellon, 2012; Bloor and Symonds, 2014). Given that sex differences have been found in each of these factors, future studies should address sex as a possible mediator in the relationship between APOE4 and obesity. Ongoing projects in our lab have begun to address this issue using female E3FAD and E4FAD mice.

How obesity and APOE interact to regulate AD pathogenesis remains to be determined. One candidate mechanism linked to both factors is metabolic impairment. Obesity is strongly associated with development of impaired glucose and insulin metabolism (Kahn et al., 2006; Singla et al., 2010), which are also characteristic of AD patients and have been proposed as possible mechanisms driving AD pathogenesis (Craft, 2005; Martins et al., 2006; Craft, 2009). Notably, APOE genotype affects metabolic responses to diet (Snoek et al., 1999; Barberger-Gateau et al., 2011), and several studies show that APOE4 carriers are at increased risk for a number of metabolic disturbances (de-Andrade et al., 2000; Oh and Barrett-Connor, 2001; Elousa et al., 2003; Marques-Vidal et al., 2003; Sima et al., 2007; Kypreos et al., 2009; Niu et al., 2009; Aatabek et al., 2012; Zarkesh et al., 2012; Guan et al., 2013), although some studies find no effect of APOE genotype on metabolic outcomes (Meigs et al., 2000; Ragogna et al., 2011). Our findings suggest that E3FAD mice may be more susceptible to some metabolic effects of Western diet, although E4FAD mice trend toward metabolic disturbances even in the absence of a Western diet. Specifically, relative to E4FAD mice, E3FAD mice showed greater diet-induced body weight gain, gonadal fat inflammatory cytokine expression, and higher glucose levels on Western diet. Conversely, E4FAD mice had higher gonadal fat pad weight and a trend toward higher fasting glucose levels than E3FAD mice under the control diet condition. These findings are consistent with several previous reports showing that mice with human APOE3 gain more weight in response to a high fat diet than mice with either human APOE4 (Arbones-Mainar et al., 2008; Segev et al., 2016) or mouse APOE (Karagiannides et al., 2008). It is important to note that the Western diet used in this study has elevated levels of saturated fats, cholesterol, and sucrose, all of which have been independently associated with increased AD-related pathology (Refolo et al., 2000; Oksman et al., 2006; Cao et al., 2007; Takechi et al., 2010). Understanding how APOE genotype interacts with various dietary components should be one target of future studies. Though metabolic factors may have a role in AD pathogenesis, our findings that metabolic outcomes of DIO were greater in E3FAD than E4FAD mice argue against the possibility that metabolic impairment significantly contributes to the observed APOE4 bias in diet-induced increases in AD-like pathology.

There are several other mechanisms besides metabolic impairment that may contribute to the observed interactions among obesity, APOE, and AD-like pathology. One established consequence of obesogenic diets is pro-amyloidogenic alteration in the expression and or activity of factors that regulate generation and clearance of Aβ, including BACE1, neprilysin, and IDE (Standeven et al., 2010; Maesako et al., 2012; Brandimarti et al., 2013; Wei et al., 2014; Maesako et al., 2015). Although we cannot exclude a significant role of such pathways in our observations, we did not observe that mRNA levels of BACE1, neprilysin, and IDE were significantly altered by either the simple or interactive effects of Western diet and APOE. Another compelling candidate mechanism is neuroinflammation, which is widely implicated as a significant regulator of AD risk and development of AD pathology (Glass et al., 2010; Wyss-Coray and Rogers, 2012; Heneka et al., 2015). Notably, both obesity and APOE4 are associated with increased inflammation in brain and systemically. For example, obesity is linked with increased immune cell infiltration into brain (Buckman et al., 2014), as well as increased glial activation (Koga et al., 2014; Dorfman and Thaler, 2015; Douglass et al., 2017). In addition, obesity increases inflammation in peripheral organs including adipose tissue (Weisberg et al., 2003; Zeyda and Stulnig, 2009) and liver (Park et al., 2010). APOE4 is also associated with greater levels of inflammation in the brain (Ophir et al., 2005; Vitek et al., 2009) and throughout the body (Colton et al., 2004; Gale et al., 2014). Moreover, stimulating innate inflammation in the presence of apoE4 increases cell death and damage in macrophages (Cash et al., 2012), and in microglia and neurons (Maezawa et al., 2006a; 2006b). In the context of AD pathology, APOE4 is associated with greater glial activation in EFAD mice (Rodriguez et al., 2014). Similarly, we found that both the total number and the relative level of morphologic...
activation of microglia and astrocytes were higher in E4FAD than E3FAD mice. Further, we observed that E4FAD mice expressed significantly higher mRNA levels of glial markers than E3FAD mice under both control and Western diets. These glial markers were significantly increased across several brain regions in response to DIO in E4FAD but not E3FAD mice. Perhaps in contrast to our results, middle-aged female APOE4 mice showed higher levels of neuroinflammation in hippocampus under control diet but decreased neuroinflammation with high-fat diet, relative to age- and sex-matched wild-type mice (Janssen et al., 2016). Though the presence of familial AD transgenes and Aβ pathology in the EFAD model may account for these divergent findings, there may also be age and sex differences in inflammatory responses to both diet and APOE4. Further, because reactive astrocytes and microglia are associated with Aβ plaques, the changes in gliosis we observe with APOE4 and DIO may be a consequence of, rather than a contributor to, Aβ pathology. Thus, additional research is needed to directly assess the potential mechanistic role of gliosis in the interaction between APOE4 and obesity in AD.

To our knowledge, this is the first experimental investigation examining the interaction between APOE4 and obesity in the context of AD. Interactions among genetic risk factors like APOE4 and environmental and modifiable lifestyle risk factors in AD have thus far not been well studied, although there are some epidemiological studies consistent with this possibility (Dufouil et al., 2000; Hanson et al., 2013; Rajan et al., 2014; Wirth et al., 2014; Ishioka et al., 2016; Zheng and Li, 2016). Our findings suggest that APOE genotype affects the relationship between obesity and AD, such that APOE4 carriers may be more susceptible to obesity-associated risks than APOE3 carriers. This illustrates an important gene-environment interaction and points to the need for additional research exploring such relationships in the context of AD, as well as identifying underlying mechanisms. Additionally, these findings identify a large population that may be at increased risk of AD, but whose chance of developing the disease may be reduced by preventative lifestyle changes.

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