mNos2 Deletion and Human NOS2 Replacement in Alzheimer Disease Models

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Abstract

Understanding the pathophysiologic mechanisms underlying Alzheimer disease relies on knowledge of disease onset and the sequence of development of brain pathologies. We present a comprehensive analysis of early and progressive changes in a mouse model that demonstrates a full spectrum of characteristic Alzheimer disease–like pathologies. This model demonstrates an altered immune redox state reminiscent of the human disease and capitalizes on data indicating critical differences between human and mouse immune responses, particularly in nitric oxide levels produced by immune activation of the NOS2 gene. Using the APPSwDI+/−/mNos2−/− (CVN-AD) mouse strain, we show a sequence of pathologic events leading to neurodegeneration, which include pathologically hyperphosphorylated tau in the perforant path away at 6 weeks of age progressing to insoluble tau, early appearance of β-amyloid peptides in perivascular deposits around blood vessels in brain regions known to be vulnerable to Alzheimer disease, and progression to damage and overt loss in select vulnerable neuronal populations in these regions. The role of species differences between hNOS2 and mNos2 was supported by generating mice in which the human NOS2 gene replaced mNos2. When crossed with CVN-AD mice, pathologic characteristics of this new strain (APPSwDI+/−/ HuNOS2α+/−/mNos2−/−) mimicked the pathologic phenotypes found in the CVN-AD strain.

Key Words: Alzheimer disease, Magnetic resonance spectroscopy, Mouse model, Neuronal loss, NOS2, Tau pathology.

INTRODUCTION

Studies on humans have provided essential information on the type and extent of glial and neuronal damage found in Alzheimer disease (AD), but the mechanisms for the initiation of the neurodegenerative changes found in AD and for disease progression remain essentially unknown. Despite a large increase in brain imaging tools for use in humans with AD (1–3), the underlying mechanisms and causal factors that contribute to AD are currently not discoverable by relying solely on human studies. Animal modeling helps to fill this void in our knowledge base, and multiple types of models have been used to increase our understanding of AD pathogenesis (4, 5). To date, many of the available mouse models that have been used to study specific components of AD pathology, such as amyloid, can be described as “incomplete” (6). They are clearly useful, particularly when used to uncover molecular details on specific AD-related proteins and the response of a physiologic system to that protein. However, it can also be argued that models that recreate the full spectrum of AD-like pathology are critical to the field. Recreating the disease progression observed in humans with AD is essential for understanding how individual components, such as amyloid, are integrated into the pathophysiologic spectrum called AD. This type of model also enables analysis of mechanisms at disease onset and the sequence of events that are relevant to the development of the overall pathologic process. In this study, we have focused on a longitudinal analysis of disease progression in a mouse model that, like presymptomatic humans with AD, proceeds to the full spectrum of AD-like pathology, including amyloid deposition, tau hyperphosphorylation and aggregation, behavioral changes, and neuronal loss with age (7–9).

The APPSwDI/mNos2−/− (CVN; CVN-AD) mouse strain used in this longitudinal study has distinct advantages, including the following: 1) mutated human amyloid precursor protein (APP) is not overexpressed but is approximately 0.5-fold of the...
normal mouse APP level (10); 2) β-amyloid (Aβ) deposition is primarily caused by accumulation of Aβ around blood vessels and an apparent failure to clear Aβ from the brain, which is similar to humans with cerebral amyloid angiopathy (CAA) (11–13); 3) subsequent tau pathology is not dependent on other mutated genes such as mutated human tau (14); and 4) the difference between mouse and human NOS2 is addressed (15–17). This latter point is particularly crucial. Recent data have re-enforced the primary role of immunity in both the onset and the progression of AD in humans (18, 19) and have also underlined the differences between human and mouse immune responses (19, 20). We have now created an additional mouse strain that directly addresses the difference in immune regulation in mouse models of human disease and the role this difference plays in discovering disease onset and progression. Mice expressing the human NOS2 gene in a mouse Nos2 background (21) were crossed with the APPSwDI/mNos2−/− mouse model. This new strain of AD mice (APPswDI+/−/HuNOS2tg+/+mNos2−/−) shows similar pathology as the APPSwDI/mNos2−/− model. Thus, this strongly re-enforces the concept that the difference in human and mouse immune-mediated nitric oxide (NO) responses is a significant factor in AD-like disease progression.

**MATERIALS AND METHODS**

**Animals**

All animal experiments were performed in accordance with protocols approved by the Institutional Animal Care and Use Committee at Duke University Medical Center under the *NIH Guide for the Utilization and Care of Vertebrate Animals Used in Testing, Research, and Training* and at Charles River Finland under the *International Guiding Principles for Biomedical Research Involving Animals* (revised 2012).

**CVN-AD Mice**

Homozygous APPSwDI/mNos2−/− (cerebrovascular amyloid Nos2−/− or CVN-AD) mice were produced by crossing mice expressing the vasculotropic Swedish K670N/M671L, Dutch E693Q, and Iowa D694N human APP mutations under control of the Thy-1 promoter (10) with mNos2−/− (B6 129P2Nos2 tau1Lau2f) mice (22) (Jackson Laboratory, Bar Harbor, ME), as previously described (9). Mice were genotyped using standard polymerase chain reaction methods. Mice were also genotyped for the Rd1 mutation in the Pde68 gene that causes blindness (23) and for the Rd8 mutation in the Crb1 gene that, when expressed, causes retinal pathology but is not associated with blindness (24). CVN-AD mice did not express Rd1 but were heterozygous for Rd8. Animals were fed standard mouse chow and housed between 12-hour light/12-hour dark cycles at 21 ± 3°C in an Institutional Animal Care and Use Committee–approved barrier facility under approved animal protocols.

**APPSwDI+/−/Hu NOS2tg+/+mNos2−/− (APP/huNOS2tg) Mice**

A mouse strain that expressed mutated human APP in the context of expressing a human NOS2 gene was constructed by crossing the APPSwDI strain with mice expressing a human NOS2 transgene on a mouse Nos2 knockout background. The HuNOS2tg+/+mNos2−/− strain was developed and characterized by Vitek et al (25) as previously described. The induction pattern of inducible nitric oxide synthase (iNOS) protein and the level of NO production by activation of the human NOS2 gene in the HuNOS2tg+/+mNos2−/− were similar to those patterns for NOS2 in human macrophages (25, 26). Importantly, and unlike C57Bl/6J control mice, the HuNOS2tg+/+mNos2−/− mouse strain expresses the 3′ untranslated region motif found in the human NOS2 gene and that binds microRNA-939, resulting in posttranscriptional disruption of iNOS protein and significantly reduced NO production (Hoos et al, in press). Because triple homozygous mice unexpectedly died at early ages, the reconstituted mouse strain was kept heterozygous for APPSwDI and homozygous for HuNOS2tg+/+mNos2−/−. APP/ huNOS2tg mice were genotyped using standard polymerase chain reaction methods and for the presence of Rd1 and Rd8. Similar to the CVN-AD strain, the APPSwDI+/−/HuNOS2tg+/+mNos2−/− mice did not express Rd1 but were heterozygous for Rd8.

**Collection of Tissue and Immunohistochemistry**

Mice used in this study were approximately 6, 12, 24, 36, 45, or 52 weeks of age. After injection with a lethal mixture of ketamine/xylazine, each mouse was intracardially perfused with approximately 25 mL of phosphate-buffered saline. Perfused brains were then rapidly removed and bisected in the midsagittal plane, and half was frozen in liquid nitrogen for use in enzyme-linked immunosorbent assay (ELISA) and Western blot analysis. The remaining half of the perfused brain was immersion-fixed in 4% paraformaldehyde and then cryoprotected by sequential passage through 10%, 20%, and 30% sucrose for 24 hours. Frozen sagittal or coronal sections (25 μm) were then cut using a sliding microtome and stored at 4°C in PBS with sodium azide to prevent microbial growth. Sections spaced either 12 or 24 slices apart (300 or 600 μm) were selected for free-floating immunohistochemistry. Immunohistochemistry was performed as previously described (9). Primary antibodies were as follows: anti–NeuN (neuron-specific nuclear) protein (MAB377, mouse monoclonal, 1:3000; Chemicon, Temecula CA), anti-Aβ (rabbit polyclonal anti-Aβ N-terminal, 1:7000; Life Technologies, Carlsbad, CA), anti-AT8 (mouse monoclonal recognizing phosphorylated Ser202/Thr205), 1:500; Thermo Scientific, Waltham MA), anti-AT180 (mouse monoclonal recognizing phosphorylated Thr231, 1:1000; Thermo Scientific), and anti–APP-A4 (MAB348, 1:500; Chemicon-Millipore, Billerica, MA). In addition, nonspecific binding of antibodies was assessed using the Innovex Biosciences Fc Receptor Blocker NB309-30 (Innovex Biosciences, Richmond, CA) following the manufacturer’s protocol or using the Vector MOM Immunodetection Kit BMK-2202 (Vector Laboratories, Burlingame CA) following the manufacturer’s protocol.

**Aβ ELISA**

Frozen brain samples were cryopulverized in liquid nitrogen, and the tissue was homogenized using glass beads and the Bullet Blender (NextAdvance, Averill Park, NY) in ice-cold Tris-buffered saline with 1% complete protease/phosphatase inhibitor (Thermo Scientific) and 2 mMol/L EDTA. The homogenate was then centrifuged at 16,000 × g for 1 hour at 4°C.
(Spin 1). The supernatant from Spin 1 containing the “soluble” Aβ40 and Aβ42 fraction was recovered to a clean tube, and the protein concentration was determined using bicinchoninic acid assay (Thermo Scientific). The pellets from Spin 1 were then rehomogenized in 70% formic acid using sonication and resupplied at 16,000 rpm (Spin 2) for 30 minutes at 4°C. The supernatants from Spin 2 (insoluble fraction) were removed to a new tube and slowly neutralized 1:20 with 1 mol/L Tris-buffered saline (pH 11). Protein levels in the insoluble samples were measured using the Bradford assay (Bio-Rad, Hercules, CA). The soluble and insoluble fractions were then assayed for levels of Aβ40 and Aβ42 using Covance BetaMark x-40 and x-42 ELISA kits (Covance, Dedham MA), as described by the manufacturer. Data were read on the SOFTMaxPro reader at 620 nm, and Aβ levels were expressed as picograms of Aβ per milligram of protein.

**Western Blot Analysis**

Frozen brain samples were cryopulverized, and the tissue was homogenized using glass beads and the Bullet Blender (NextAdvance) in ice-cold Tris-buffered saline with 1% complete protease/phosphatase inhibitor (Thermo Scientific) and 2 mmol/L EDTA. lysates were then centrifuged at 16,000 × g for 30 minutes at 4°C to pellet insoluble proteins. Supernatants were removed to clean tubes, and protein levels were measured using the BCA protein assay kit (Pierce, Rockford IL). Western blot analyses were performed as previously described (27); digital images were captured using the FluorChem Q imaging system (Cell Biosciences, Santa Clara, CA). Quantitative analysis was performed using Alpha View software (Cell Biosciences). Individual densitometry values were normalized to actin.

**Sarkosyl Extraction of Aggregated Tau**

Frozen cryopulverized brain samples were homogenized in buffer containing 50 mmol/L Tris (pH 8.0), 274 mmol/L NaCl, 5 mmol/L KCl, 1 mmol/L phenylmethyl-sulfonyl fluoride, and protease/phosphatase inhibitor cocktail. The homogenate was centrifuged at 1,300 × g for 15 minutes at 4°C, and the supernatant was further centrifuged at 100,000 × g for 25 minutes at 4°C. The supernatant was saved as soluble tau (S1) fraction. The pellet was homogenized in buffer containing 10 mmol/L Tris HCl (pH 7.4), 0.8 mol/L NaCl, 10% sucrose, 1 mmol/L ethylene glycol tetraacetic acid, 1 mmol/L phenylmethyl-sulfonyl fluoride, and protease/phosphatase inhibitor, and centrifuged at 100,000 × g for 25 minutes. The supernatant from this spin was incubated with 1% sarkosyl at 37°C for 1 hour and then centrifuged at 100,000 × g for 30 minutes. The supernatant was saved as sarkosyl-insoluble tau fraction (S2) fraction. The pellet, which contains sarkosyl-insoluble tau fraction, was suspended in Tris-EDTA buffer and subjected to Western blot analysis as described previously.

**[1H]Magnetic Resonance Spectrometry**

[1H]Magnetic resonance spectroscopy ([1H]MRS) analysis was performed in a horizontal 11.7-T magnet (bore size, 160 mm) equipped with a gradient set capable of a maximal gradient strength of 750 mT/m and interfaced with a Bruker Avance III console (Bruker Biospin, Ettlingen, Germany). A volume coil (Bruker Biospin) was used for transmission, and a surface phased array coil was used for reception (Rapid Biomedical, Rimpar, Germany). Isoflurane-anesthetized mice were fixed to a head holder and positioned in the magnet bore in standard orientation relative to gradient coils.

To collect [1H]MRS data, we placed a voxel bilaterally in the hippocampus of the mouse based on the anatomic images collected, as described previously. Automatic 3-dimensional gradient echo shimming was initially used to adjust B0 homogeneity in the voxel. Water signal was suppressed using variable-power radiofrequency pulses with optimized relaxation delays to obtain B1 and T1 insensitivity. A STEAM sequence (echo time, 3 milliseconds), combined with outer volume suppression, was used for prelocalization. Three outer volume suppression blocks were used interleaved with water suppression pulses. Data were collected by averaging 512 excitations (frequency corrected for each average) with repetition time of 4 seconds, 4,096 points, and spectral width of 5 kHz. In addition, a reference spectrum without water suppression was collected from the identical voxel using the same acquisition parameters. Peak areas for metabolites (N-acetyl aspartate [NAA], alanine [Ala], choline, inositol, creatine [Cr] + phosphocreatine, glutamine, glutamic acid [GLU], and glutathione [GSH]) were analyzed using LCModel (Stephen Provencher Inc, Oakville, ONT, Canada); results were given relative to water content in tissue. Two-way analysis of variance (ANOVA) was used to determine differences between C57Bl/6J and CVN-AD mice across the 4 ages studied. Metabolic ratios were determined relative to total Cr, and then each metabolic ratio was analyzed separately, with genotype as between-subject factor and age as within-subject factor. Bonferroni posttests were performed to determine significant differences between CVN-AD and C57Bl/6J metabolic ratios at each age.

**Stereologic Analysis**

NeuN-immunopositive neurons were counted in the CA3 region of the left hippocampus using the optical fractionator method, as previously described (28). Every 12th sagittal section (25 μm thickness per section) was counted from the bregma to 2.0 mm lateral of the bregma on a Nikon 218912 light microscope (20× objective; Nikon, Tokyo, Japan) with computer-controlled mechanical stage interfaced with the Stereo Investigator 9 software package (MicroBrightField, Williston, VT). Only the CA3 region was counted, and animals with sections that were damaged were excluded from the analysis. All slides were precoded to ensure blinding by the observer. The number of neurons was calculated using the following equation: \( N = Q \times 1/ssf \times 1/asf \times 1/hsf \), where \( N \) is total neuron number, \( Q \) is the number of neurons counted, ssf is section sampling fraction, asf is area sampling fraction, and hsf is height sampling fraction. Data are presented as neuronal number per cubic millimeter.

**Radial Arm Water Maze**

CVN-AD, C57Bl/6J, and mNos2−/− mice were tested at the ages indicated using the 2-day radial arm water maze (RAWM) previously described in detail (9, 29). Briefly, a 6-arm maze is submerged in a pool of opaque water, and a platform is placed at the end of 1 arm below the surface. The
mouse receives 15 trials per day for 2 days and is started in a different arm each trial, whereas the arm containing the platform remains the same. Using static visual cues, the mouse learns the position of the escape platform. The first 10 trials are considered training and alternate between a visible platform and a hidden platform. The final trials for Day 1 and all trials on Day 2 use a hidden platform. The number of errors (incorrect arm entries) is counted across a 1-minute period. The errors are averaged over 3 trials per block, resulting in 10 blocks for the 2-day period. After completion of all hidden maze tasks, mice were tested for sensory or motor deficits using the open pool task with visible platform, as described by Alamed et al (29). Mice that did not perform successfully in the open pool task were excluded from data analysis. All behavioral tests were performed with the investigator blinded to treatment groups.

**Contextual Fear Conditioning Test**

The contextual fear conditioning test is modified from Comery et al (30). Training and testing were conducted on 2 consecutive days, using the Coulbourn FreezeFrame system (Coulbourn Instruments, Harvard Apparatus, Holliston MA). Training consisted of placing a subject in a chamber with bright house light on and allowing exploration for 2 minutes. Afterward, an auditory cue (1,700 Hz, 80 dB, conditioned stimulus) was presented for 15 seconds. A 2-second foot shock (1.5 mA; unconditioned stimulus) was administered for the final 2 seconds of the conditioned stimulus. This procedure was repeated, and the mouse was removed from the chamber 30 seconds later. At 2 hours after the training, mice were returned to the same chamber in which the training occurred (context), and freezing behavior was recorded by a computerized camera tracking system. The automated FreezeFrame system (which digitizes the video signal at 4 Hz and compares movement of the mouse frame by frame) was used to score the amount of freezing. At the end of the 5-minute context test, mice were returned to their home cage. Two hours later, freezing was recorded in a novel environment (altered context) and in response to the cue. The mouse was placed in the novel environment, and time sampling was used to score freezing for 3 minutes. The auditory cue (1,700 Hz, 80 dB, conditioned stimulus) was then presented for 3 minutes, and freezing was again scored. Freezing scores for each subject were expressed as a percentage for each portion of the test. Memory for context (contextual memory) for each subject was obtained by subtracting the percentage of freezing in the novel environment from that in the context.

**Statistical Analyses and Comparisons Between Groups**

Mean ± SE values were determined for each of the outcome measures described previously at 6, 12, 24, 36, 45, and/or 52 weeks of age in the CVN-AD and control mice. Except for the [1H]MRS analysis, mixed sexes were used in the analyses, and sex-based differences were not investigated. Significance with age within strain was determined using Student t-test or 1-way ANOVA; statistical significance between genotypes and age was determined using 2-way ANOVA (GraphPad Software, San Diego, CA). Significance was set at p ≤ 0.05.

**RESULTS**

**Expression of APP, Aβ, and Pathologic Tau With Age**

We used an antibody that recognizes the N-terminus of APP to detect immunostaining for holo-APP in neurons at the earliest stages of disease. As predicted for the Thy-1 promoter, holo-APP was observed within postmitotic neuronal populations and was found beginning at 6 weeks of age. These populations included neurons in the CA3 region of the hippocampus, in the mossy fiber region of the dentate gyrus (DG), in layer 5/6 of the cortex, and in the subiculum (Figs. 1A–C). Also beginning at 6 weeks of age, amyloid deposits were observed primarily around small-diameter blood vessels in the subiculum (Fig. 1D). β-Amyloid deposits in CVN-AD mice initially appeared as “caplike” structures reminiscent of CAA found in humans with AD (11, 31) (Fig. 1G). Immunoreactive Aβ deposits increased with age, appearing within the hippocampus and cortical layers beginning at 12 weeks of age and then spreading throughout other brain regions (Figs. 1E, F, H). The increase in Aβ levels with age was quantified, using ELISA, for soluble and insoluble Aβ40 and Aβ42. Figure 1 also shows the mean ± SE values for total Aβ40 (soluble plus insoluble) and total Aβ42 (soluble plus insoluble) at 6, 12, 24, 36, and 52 weeks of age. Both Aβ40 and Aβ42 were significantly increased, but total Aβ40 rose rapidly to a plateau by 12 weeks of age then rose again at 52 weeks, whereas total Aβ42 increased significantly at each age (Fig. 1I).

Data from our previous studies showed the presence of hyperphosphorylated and aggregated tau in CVN-AD mice aged 52 weeks or older. To understand the onset of tau pathology in this model, we used antibodies that target phospho-PHF tau. AT8 (pThr202/pSer205) and AT180 (pThr231) react with specific phosphoepitopes on the tau protein that are phosphorylated in AD patients. Positive immunoreactivity using these antibodies is commonly recognized as a marker of disease-relevant changes in the tau phosphorylation status (32–36). AT8-positive neuron fibers were observed in the mossy fibers of the hilus at 6 weeks of age (Figs. 2A, B) and in the outer layer of the cortex in CVN-AD mice around 12 weeks of age (Figs. 2C, D). AT8 immunoreactivity increased with age and was observed in multiple locations throughout the brain, including the hilus of the DG (Figs. 2I, J, N), in association with structures that resembled neuritic plaques (Figs. 2E, G) and around blood vessels (Figs. 2I, K, M, O). Although observed as early as 12 weeks, mice at 36 and 52 weeks demonstrated significant AT8 immunostaining around blood vessels, particularly in the subiculum and cortex (Figs. 2K, O). Immunostaining for AT180 followed the same regional patterns of immunostaining with age as for AT8 in CVN-AD mice (Figure, Supplemental Digital Content 1, http://links.lww.com/NEN/A608).

Although AT8-immunopositive tau has been observed in other APP-only mouse models of AD primarily around cortical neuritic plaques (37–39), we sought to confirm the
presence of hyperphosphorylated and aggregated tau in CVN-AD brain (Fig. 3). No detectable change in AT8-immunopositive tau was found for CVN-AD mice compared with C57Bl/6J mice by Western blot analysis using whole-brain lysates at 6 weeks (Figs. 3A, B), but increased levels of soluble hyperphosphorylated tau in whole-brain lysates were observed by Western blot analysis at 12 weeks of age (Figs. 3C, D). Although controversial, mouse tau has been shown to form insoluble aggregates in vitro and in vivo (40–43). To show aggregated tau and to follow its production with age, we assessed whole-brain lysates for the presence of sarkosyl-insoluble and hyperphosphorylated mouse tau at 6, 12, 24, and 45 weeks of age. No insoluble tau was found in CVN-AD mice at 6 or 12 weeks of age (data not shown), but strong sarkosyl-insoluble AT8-positive tau bands were observed in CVN-AD mice compared with C57Bl/6J mice at 24 and 42 weeks of age (Figs. 3E, F).

Neuronal Damage and Loss Increase With Age

To characterize neuron changes, we next used proton magnetic resonance spectroscopy ([1H]MRS) to compare live (isoflurane-anesthetized) CVN-AD and C57Bl/6J mice of the same ages. [1H]Magnetic resonance spectroscopy is commonly used to detect changes in metabolic substrates in vivo that reflect altered neuronal integrity and has been used as a biochemical imaging tool for the diagnosis of AD (44–47). Typical neuronal and metabolic substrates that are measured using this technique include neurotransmitters (glycine [GLY], g-aminobutyric acid [GABA], glutamate [GLU] and taurine [TAU]), metabolites (alanine [Ala], phospho-creatine [PCr],...
creatine [Cr], myo-inositol [INS], choline [CHO], phosphocholine [PCh]; glycerophosphocholine [GPC], N-acetyl aspartate [NAA]; N-acetylaspartylglutamate [NAAG]), and antioxidants (i.e. glutathione [GSH]).

Anesthetized CVN-AD male mice were imaged using an 11.7-T magnet at 12, 24, 36, and 52 weeks of age, and the metabolic substrate levels in the hippocampus were compared with levels of the same substrates in the hippocampus of age-matched C57Bl/6J mice. Figure 4 shows the mean levels of brain metabolites measured at 12, 24, 36, and 52 weeks of age; data are presented as the ratio of metabolite to Cr levels. Creatine has high intermeasurement stability and is generally considered to be a useful internal reference for measurement variability (45, 48). We observed small but statistically significant reductions in Ala/Cr ratio at 12 weeks and in GSH/Cr ratio at 24 weeks. However, these changes were apparently transient and were no longer evident at either 36 or 52 weeks of age. Myo-inositol levels, generally believed to be an indicator of glial “activation” (49–51), were significantly increased at 24, 36, and 52 weeks of age. Changes in neurotransmitter and neuronal indices were clearly shown at 52 weeks of age. Ratios of GLU, NAA, or NAAG to Cr were significantly reduced in CVN-AD mice. Importantly, the NAA/INS ratio in CVN-AD mice decreased from 1.18 at 12 weeks of age to 0.99 at 52 weeks of age, whereas the ratio in C57Bl/6J mice was 1.27 and 1.26 for 12 and 52 weeks, respectively. A declining

FIGURE 2. AT8 tau immunostaining in CVN-AD mice varies with age. Representative photomicrographs of CVN-AD mice at 6 weeks (A, B), 12 weeks (C, D), 24 weeks (E-G), 36 weeks (I-K), and 52 weeks (M-O) of age. mNos2−/− control mice stained with AT8 are shown in (H), (L), and (P) at 24, 36, and 53 weeks of age, respectively. Early tau pathology is observed in the DG, particularly in the mossy fibers of the polymorphic molecular layer (pml) (B is a magnified view of A; arrow denotes AT8-positive immunostain in the granule layer) and in the entorhinal cortex (Cx) (C; arrow denotes the region magnified in D). AT8-positive tau is also found around plaque-like structures in the cortex (E, G). Arrows in (E) show the regions of the cortex that are magnified in (F) and (G). With increasing age, tau AT8-positive immunostaining accumulates around blood vessels in multiple regions of the cortex and hippocampus (Hip) (I-K, 36 weeks; M-O, 52 weeks). Arrows in (J) and (M) denote the regions shown in the magnified views of the appropriate respective panels. For low-power views, scale bar = 500 μm. For higher-magnification views, scale bar = 100 μm. Sub, subiculum.
CA1 neuron loss was also readily observed in coronal sections of the hippocampus in CVN-AD mice using NeuN immunostaining (Fig. 7). Neuronal layers in the intermediate zone of the hippocampus were thinned or missing compared with either C57Bl/6j or mNos2−/− mice. This region has been shown to regulate both spatial and emotion-based memory and learning (53).

Memory and Learning

Reduced memory and learning behavior is a well-described characteristic of AD patients (54, 55) and has been observed in aged mouse models of AD, including the CVN-AD mouse at 52 weeks of age (56, 57). To examine hippocampal changes that are correlated with behavioral outcomes at each age and with disease progression, we used the RAWM, which tests spatial memory and learning (29, 58), and contextual fear conditioning test, which tests contextual integration and fear memories (59). CVN-AD mice showed a significantly increased number of errors in finding the escape platform during the RAWM compared with either mNos2−/− or C57Bl/6j control mice at 24, 36, and 52 weeks of age (Figs. 8A–C), but not at 12 weeks (data not shown). We also compared the number of errors for the last day of trials for CVN-AD mice at each age (Fig. 8D). These data show that although spatial memory was altered in 24- and 36-week-old CVN-AD mice versus control mice, there were no significant differences in memory between these 2 ages. There was, however, a significant increase in severity of memory loss at the 52-week time point versus the earlier time points in CVN-AD mice. mNos2−/− control mice did not show age-dependent differences in memory and learning from 12 to 52 weeks (Figure, Supplemental Digital Content 2, http://links.lww.com/NEN/A609). Failure to find the platform was not attributable to blindness. CVN mice were negative for the Rd1 gene that is known to cause blindness in mice, and all mice tested in the RAWM were additionally assessed using the open pool task with visible platform. Contextual fear memory was also significantly reduced beginning at 24 weeks of age and worsened with age and disease progression (Fig. 8E). Additional behavioral testing was performed. Figure, Supplemental Digital Content 3, http://links.lww.com/NEN/A610, shows Morris water maze performance data in CVN mice versus C57Bl/6j mice at 52 weeks of age. Figure, Supplemental Digital Content 4, http://links.lww.com/NEN/A611, shows open field performance data in CVN mice versus C57Bl/6j mice at 12 and 52 weeks of age.

Reconstitution of the Human NOS2 Gene in CVN-AD Mice

Although the lack of iNOS-mediated NO production in the CVN-AD model is a phenocopy of the low NO level observed in human immune responses, this model does not fully mimic the immune/redox environment that is typical of human brain that expresses human NOS2. To begin to address this problem, we generated an additional strain of mice that expresses the human NOS2 gene on a mNos2−/− background. When induced, NO levels that are more representative of human immune responses are produced. This transgenic mouse is termed the HuNOS2H/mNos2−/− strain, and the initial description of its characteristics has been
previously published (21). The HuNOS2tg+/+/mNos2j strain was then crossed with the APPSwDI/mNos2j strain to generate a triple transgenic that expresses a mutated human APP gene (APPSwDI) and 2 copies of the human NOS2 transgene on a disrupted mNos2 knockout background. Mice that were homozygous for APPSwDI died at variable ages between 30 and 50 weeks of age. Survival curves were not established. The new line was then kept heterozygous as for APPSwDI (APPSwDI+/-/HuNOS2tg+/+/mNos2j) and was termed APP/huNOS2tg mice. These mice were then examined for brain pathology at different ages.

Typical AD-like pathology was observed in the brains of APP/huNOS2tg mice, which closely resembled the neuropathologic patterns found in CVN-AD mice. As predicted by the expression of mutated APP genes, Aβ deposits were observed throughout the cortex and hippocampus (Figs. 9A, B). In addition, ELISA of total Aβ (soluble + insoluble Aβ40 and Aβ42) levels in whole-brain lysates from 36- and 52-week-old APP/huNOS2tg mice revealed a significant increase with age. Mean values rose from 79,050 ± 23,500 to 193,000 ± 48,000 pg/mg protein (p = 0.05; n = 4–7 mice), an approximately 2.3-fold increase. We also observed AT8-positive immunoreactivity to hyperphosphorylated tau in the cortex of APP/huNOS2tg mice at 52 weeks of age (Figs. 9C, D). Similar to CVN-AD mice, sarkosyl-insoluble tau was detected in brain lysates at 52 weeks of age (Fig. 9E).
To determine whether neuronal changes were observed in APP/huNOS2tg, we stained brain sections for NeuN and counted neurons using unbiased stereology in CA2/CA3 hippocampal regions. Representative views of the hippocampi and corresponding CA3 regions at 52 weeks of age are shown in Figures 10A, B for HuNOS2tg+/+ mNos2−/− control mice and in Figures 10C, D for APP/huNOS2tg mice. Using unbiased stereology, we detected a significantly decreased number of neurons in the CA3 region in APP/huNOS2tg mice compared with either mNos2−/− or HuNOS2tg+/+ mNos2−/− control mice (Fig. 10E). Behavioral changes were also observed in APP/huNOS2tg mice. Radial arm water maze tests showed a significantly increased number of errors in APP/huNOS2tg mice compared with either mNos2−/− or HuNOS2tg+/+ mNos2−/− control mice at 52 weeks of age (Fig. 10F). However, these deficits were only found on the second day of trials, which comprised the memory retention phase of the learning and memory behavioral task. This difference in response from CVN-AD mice may in part be caused by the expression of only 1 copy of the APPSwDI gene. APP/huNOS2tg mice are heterozygous for APP.

**DISCUSSION**

Our detailed longitudinal study on disease progression in the CVN-AD mouse model provides an integrated picture of brain response to the production of mutated human APP.

**FIGURE 5.** Loss of NeuN-immunopositive neurons in the hippocampus in CVN-AD mice starting at 36 weeks of age. (A–J) Representative photomicrographs depicting the CA2/CA3 subregion of the hippocampus of CVN-AD mice at 12 weeks (A, B), 36 weeks (E, F), and 52 weeks (I, J) of age. Panels (C, D) (12 weeks), (G, H) (36 weeks), and (K, L) (52 weeks) show representative age-matched mNos2−/− control mice. For low-power view, scale bar = (E) 500 μm. For high-power view, scale bar = (F) 100 μm. Sub, subiculum.

**FIGURE 6.** Neuron numbers in the CA2/CA3 region of CVN-AD versus mNos2−/− mice. NeuN-immunopositive neurons in the CA2/CA3 region were counted using unbiased stereology for CVN-AD and mNos2−/− control mice at 12, 24, 36, and 53 weeks of age. Data are presented as mean ± SE. *** p < 0.001 for difference between mouse strains using 2-way ANOVA (n = 5–9 mice analyzed at each age).
FIGURE 7. Neuron loss in the CA1 intermediate zone is observed in CVN-AD mice. (A–I) Subregions of the mouse hippocampus are defined in (A). Representative views of neuronal layers stained for NeuN in the CA1 intermediate zone are shown for C57Bl/6J wild-type (WT) mice (B, C) and mNos2−/− mice (D, E). Control WT mice show an uninterrupted, dense CA1 pyramidal cell layer. CVN-AD mice demonstrate an aberrant CA1 pyramidal layer with an apparent altered density of neurons (F–I). Boxes on the left panels represent areas that are magnified in the adjacent right panels. Left: scale bar = (B) 500 μm. Right: scale bar = 100 μm.
protein across time. Although this model represents early-onset (familial) AD rather than late-onset (sporadic) AD, it provides us the opportunity to explore the vulnerability of key brain regions, such as the subiculum and hippocampus, when Aβ deposition is increased. Equally importantly, the CVN-AD model allows the study of how tau pathology may interplay with amyloid deposition in the same regions. Regionalism of amyloid deposition is characteristic of mouse models that use mutated human \textit{APP} or \textit{PS1} gene expression governed by the \textit{Thy-1} or prion protein promoter. For example, \textit{APP}^{Sw/Tg2576} (60), \textit{APP}^{Sw/PS1K1} (61), and \textit{5xFAD} (62) mice express mutated human \textit{APP} protein at the fifth and sixth layers of the cortex; in the CA3, CA1, and DG of the hippocampus; and in the subiculum early in the disease. Similar \textit{APP} expression and Aβ deposition patterns are found in CVN-AD mice even though the expression levels of mutated human \textit{APP} within neurons are much lower than those in other models. Aβ-Amyloid deposition in our model is found at the earliest time points around small blood vessels in the subiculum and seems to be restricted initially to this area. These Aβ deposits are consistent with the local release of Aβ from neurons in the region and with the vasculotropic nature of Dutch/Iowa mutations (13, 63, 64). The Dutch/Iowa mutations in \textit{APP} lead to abnormal Aβ aggregation around blood vessels at the blood-brain interface (i.e. CAA). Individuals expressing Dutch/Iowa \textit{APP} mutations can show hemorrhages, infarcts, and microbleeds in severe cases (64–66). The high degree of amyloid accumulation around the small-diameter vessels implies that the normal flux of Aβ peptides via the brain’s fluid drainage pathways may be obstructed as an early pathologic event (11, 67). In addition, restriction of critical transport mechanisms across the blood-brain barrier or the induction of an immune response by pericapillary deposits may rapidly alter the normal parenchymal microenvironment. Disrupted neuvascular unit function, including damaged glial end feet and loss of aquaporin and potassium channel function, has been reported for CVN-AD mice at 52 weeks of age (68). However, other mouse models of AD also show rapid and high levels of Aβ deposition in the subiculum, particularly when expression levels of the mutated \textit{APP} gene are extremely high (60, 61, 69). A preferential accumulation of amyloid in the subiculum, the involvement of Aβ in blood vessels, and the potential for damage to the neuvascular unit in these models are essentially equivalent to the CVN-AD model. How this profile of amyloid deposition leads to downstream AD pathology, including neuronal loss in humans, remains unclear. Unlike CVN-AD mice, the \textit{APP}^{SwDI} parent mouse strain and other mouse models that show accumulation of Aβ in the subiculum do not show neuronal loss in this region (70). Models that coexpress multiple \textit{APP} mutations and \textit{PS1} mutations show limited neuronal loss in the pyramidal layers of the CA1 region, but changes in subiculum neuronal number have not been reported (38, 71).

**FIGURE 8.** Spatial memory and contextual fear memory are altered in CVN-AD mice with age. (A–C) Mean \pm SE number of errors exhibited in the RAWM task during 2 days of trials (Materials and Methods section) in CVN-AD mice versus mNos2\textsuperscript{\textminus /\textminus} and control mice at 24 weeks (A), 36 weeks (B), and 52 weeks (C) of age. *** \( p < 0.001 \) by 2-way ANOVA (n = 10–20 mice per group). (D) Comparison of the memory retention phase of the RAWM across age and genotype. \( p < 0.001 \) for 52 weeks versus 36 or 24 weeks by 2-way ANOVA. (E) Mean \pm SE values for contextual memory for each mouse obtained by subtracting the percentage of freezing in the novel environment from that in the context. * \( p < 0.05 \), ** \( p < 0.01 \) (n = 10 mice per group). WT, wild type.
Involvement of Tau

An association of AT8-positive tau with Aβ-laden blood vessels in the subiculum is also found in CVN-AD mice at the earliest age studied and continues with increasing age. Perivascular accumulation of hyperphosphorylated tau has been observed in the presence of CAA in humans with AD (11, 31, 72). Using old (12 months) CVN-AD mice, Van Nostrand et al (73) found hyperphosphorylated tau within neurons, astrocytes, and microglia surrounding the small blood vessels. We speculate that the accumulated tau around the neurons, astrocytes, and microglia surrounding the small blood vessels “spreads” to other regions of the brain (74–77). The mechanisms underlying this “spreading” process remain unknown, but altered patterns of uptake may be generated around amyloid-laden blood vessels. How abnormal tau affects the function of the neurovascular interface is currently unknown. Again, however, the subiculum is implicated as one of the first regions of the brain to be impacted by tau pathology.

Along with the CA1 and CA3 regions of the hippocampus, the subiculum is a vulnerable region in AD patients and shows pathologic alterations, including atrophy, early in the disease process (78–80). Damage to this region would impact brain function because the subiculum is a critical relay between the entorhinal cortex and the hippocampus. Neurons displaying hyperphosphorylated and aggregated tau protein are well known to pass through the subiculum in AD patients. The exact relationship of hyperphosphorylated (i.e., AT8-positive) and aggregated tau with neuronal loss, however, has been controversial (6, 81–88). This relationship has been particularly difficult to study in mouse models of AD because of the inability to show significant tau pathology in amyloid models unless mutated human tau proteins (e.g., P301L tau mutation) are coexpressed (89). Because mutations may dictate the development of pathologic tau differently when compared with the absence of mutations, the CVN-AD mice provide unique insights into changes in mouse tau as the disease progresses with age. The earliest appearance of hyperphosphorylated tau, as judged by AT8 immunostaining, was found at 6 and 12 weeks of age in the subiculum associated with blood vessels but was also found in neurons in the entorhinal cortex, the DG, the polymorphic layer and mossy fibers in the hilus, and the CA3 neuronal targets of these fibers. These expression patterns closely match those in AD (84, 90) and, with the exception of the subiculum and CA3 region of the hippocampus, do not seem to correlate directly with marked APP expression or amyloid deposits at the same age. Interestingly, the deposition of hyperphosphorylated tau in a specific subset of CA3 neurons has been recently demonstrated in AD patients and is proposed to be instrumental to the loss of the interconnecting neuronal networks between the entorhinal cortex and the hippocampus (91). Also as in AD, our data from CVN-AD mice clearly show that tau hyperphosphorylation increases with age in neurons throughout the hippocampus and cortex and in association with amyloid plaques. Detection of sarkosyl-insoluble tau in CVN-AD mice confirms the presence of disease modifications to tau that result in aggregate formation. These data further support earlier studies by Kamper et al (43) and Adams et al (92), who have shown that mouse tau can form insoluble aggregates. Finally, the hyperphosphorylated and aggregated tau we observed in the CVN-AD mice is also found in APP/huNOS2mice. Reconstitution of human NOs2 into a mouse Nos2−/− background (as in APP/huNOS2mice) produces the same general phenotype as the CVN-AD mice and a different phenotype from mice that express mutated APP or mNos2 alone. These data reinforce the concept that a difference between human and mouse NO production contributes to the different patterns of pathology found in mouse models of AD and humans with AD.
FIGURE 10. Neuron loss and behavioral deficits in APP/huNOS2tg mice. (A–D) Representative brain sections from HuNOS2tg/mNos2j/j, mNos2j/j, and APP/huNOS2tg mice were immunostained for NeuN to identify neurons. Typical views of the CA2/CA3 region of the hippocampus in HuNOS2tg/mNos2j/j mice at low magnification (A) and higher magnification (B). Typical views of the same region in 52-week-old APP/huNOS2tg mice (C, D). (E) Numbers of neurons per cubic millimeter are significantly decreased in APP/huNOS2tg mice versus either HuNOS2tg/mNos2j/j or mNos2j/j mice. p < 0.001 using 2-way ANOVA (n = 5–9 mice per group). (F) The mean number of errors made in finding the platform in the RAWM for APP/huNOS2tg mice versus either HuNOS2tg/mNos2j/j or mNos2j/j mice, p < 0.001 using 2-way ANOVA (n = 10–15 mice per group). Scale bars = (A, C) 500 μm; (B, D) 100 μm. ***p < 0.001.
Detecting Neuronal Damage in Young Mice

Although the pathologic interaction of Aβ and hyperphosphorylated tau is implicated in neuronal loss in AD, the mechanisms that lead to cell death remain unclear. Atrophy in the hippocampus and subiculum is well-described, and critical features of disease progression in AD are associated with neuron loss (28, 93). Thus, mouse models that show full progression to neuronal loss are critical. As previously reported, CVN-AD mice aged 12 months or older show extensive neuron loss in the hippocampus and subiculum (9). Using in vivo imaging with [1H]MRS, we compared changes in hippocampal metabolites in CVN-AD mice at 12, 24, 36, and 52 weeks of age with those in C57Bl/6J mice at the same age. APP/huNOS2tg mice were not studied in this manner. The high field strength we used facilitated the reliable detection of a wide variety of metabolites at each of time point, thus allowing us to detect early changes in brain functioning. We observed changes in specific metabolites that reflect the altered metabolic states of the hippocampus throughout disease progression, beginning with reduced levels of Ala found only in 12-week-old mice. Interestingly, this abnormal [1H]MRS progression, beginning with reduced levels of Ala found only in the hippocampus and subiculum (9). Using in vivo imaging with [1H]MRS, we compared changes in hippocampal metabolites in CVN-AD mice at 12, 24, 36, and 52 weeks of age with those in C57Bl/6J mice at the same age. APP/huNOS2tg mice were not studied in this manner. The high field strength we used facilitated the reliable detection of a wide variety of metabolites at each of time point, thus allowing us to detect early changes in brain functioning. We observed changes in specific metabolites that reflect the altered metabolic states of the hippocampus throughout disease progression, beginning with reduced levels of Ala found only in 12-week-old mice. Interestingly, this abnormal [1H]MRS finding corresponds in time to a significant increase in the overall activity levels (hyperactivity) of CVN-AD mice compared with C57Bl/6J control (WT) mice at this age. It did not correspond to altered contextual memory or spatial memory.

A transient decrease in GSH levels was observed at 24 weeks of age in CVN-AD mice. Glutathione is an antioxidant found primarily in astrocytes that functions as a critical regulator of cellular and tissue redox state (94). Altered ratios of reduced GSH to oxidized GSSG are believed to be a principal factor in the oxidative stress associated with AD (95). Therefore, it is likely that the reduced GSH levels observed at this age signal worsening oxidative stress before measurable neuronal loss. The change in antioxidant profile was associated in time and disease stage with increased brain INS levels. Similar to GSH, INS is thought to represent glia parameters, and increased hippocampal INS levels have been clearly shown in AD patients (44, 47, 48, 51). These changes are not consistently observed in other mouse models of AD, such as the APP/PS1 mice (96). CVN-AD mice at 24 weeks of age also displayed significantly reduced contextual memory and increased errors in the RAWM task. Later stages of disease progression demonstrated changes in metabolites that are consistent with increasing glial and neuronal damage. Increased INS levels were found at both 36 and 52 weeks of age, whereas decreased levels of NAA and GLU + GLN were found at 52 weeks of age. These [1H]MRS data from CVN-AD mouse brains are highly reminiscent of changes in metabolites observed in AD. Humans diagnosed as having either mild cognitive impairment or more advanced AD show progressively decreased levels of NAA/Cr and GLU + GLN/Cr ratios compared with control age-matched individuals (44, 97).

Neuronal Loss in the Critical Relay Between the Entorhinal Cortex and the Hippocampus

Here, we specifically focused on neuronal loss in the hippocampus, particularly in the CA2/CA3 subregion. The importance of this subregion has been underscored by Braak et al (84), who hypothesized that the appearance of hyperphosphorylated tau in the entorhinal cortex to the DG to the CA2/CA3 circuitry serves as an indicator of AD progression in humans. Furthermore, interconnections between CA2/CA3 neurons and neurons in the DG and entorhinal cortex play critical roles in memory and learning behaviors in humans and mice (54, 98). Thus, loss of CA2/CA3 neurons is likely to disrupt these circuits. Our results demonstrated no significant change in CA2/CA3 neuronal number in 12- or 24-week-old CVN-AD mice versus control mice. By 36 weeks, however, there was a 30% loss of neurons, with no further increased loss at 52 weeks. These data suggest that a specific subset of CA2/CA3 neurons may be particularly vulnerable in CVN-AD mice. The same area is damaged in old APP/huNOS2tg mice, which also show significant neuronal loss. Interestingly, using magnetic resonance imaging, Hanseu et al (99) found a significant increase in CA2 atrophy compared with other hippocampal subregions in mild cognitive impairment patients with amnesia.

The neuronal damage and loss observed in the hippocampus of CVN-AD mice and APP/huNOS2tg mice are clearly significant factors in the learning and memory deficits observed in these mice. However, the behavioral profiles of CVN-AD and APP/huNOS2tg mice resemble the behavioral outcomes observed in other models of AD that do not show neuronal death (100). As in other AD models, CVN-AD mice show behaviors that include decreased contextual fear conditioning and increased spatial memory loss. These behavioral changes are found as early as 12 weeks of age and do not correlate with our data on neuronal loss at this age. Rather, any disease-related damage to the memory circuitry is likely to represent the disruption of synapses and neuronal plasticity, resulting in abnormal hippocampal network circuitry before overt measurable neuronal loss. In this regard, Hoos et al (27) recently published a detailed analysis of unbiased protein expression levels in CVN-AD mice at 6, 12, 24, and 42 weeks. They used liquid chromatography–tandem mass spectrometry to quantitate protein levels and the accompanying phosphorylation profiles of the same brain lysate samples. Multiple proteins that are related to regulation of neuronal plasticity (e.g., α-internexin or ninein) (101, 102), to the structure and function of synapses (e.g., bassoon, Shank2, and syntaxin B) (103), and, importantly, to regulation of visual spatial memory (e.g., disks large 2 [DLG2]) (104) are significantly altered in CVN-AD mice. With continued disease progression in CVN-AD mice, however, regional neuronal loss is likely to add to the behavioral deficits generated by synaptic damage, as observed in the RAWM results at 52 versus 36 weeks of age.

The behavioral changes in CVN-AD mice and mouse models of AD include both spatial memory and emotional memory deficits. Recent studies by Fanselow and Dong (53), Gallagher and Koh (54), and Bast (105) have provided a more detailed view of the anatomic regional connections in the hippocampus that lead to changes in memory. In general, spatial memories are formed as part of the synaptic interconnections of the dorsal and intermediate hippocampi (CA1/CA3d), whereas the ventral hippocampus (CA1v/CA3v) is associated with emotional memories. As discussed by Fanselow and Dong (53) and Bast (105), it is likely that the neurons in the intermediate zone between the dorsal region and the ventral...
region play the important task of integrating and linking the emotional/motivational aspects of forming memory with the visual spatial information. Importantly, we show disruption of CA1 intermediate zone neurons in CVN-AD mice. Damage to this area is predicted to impact both cognitive and motivational behavioral outcomes, as we have observed in CVN-AD mice. Gene mapping has also now provided identification of genes in specific regions of the dorsal, intermediate, and ventral zones (106). Candidate genes that overlap with the intermediate zone of the CA1 region were identified by Thompson et al (106) and include kctd4 (potassium channel tetramerization domain containing-4), pkp2 (plakophilin gene family) and dkk3 (dickkopf gene family). The roles of these genes in AD are unknown, but further study of their functional outcomes may ultimately provide insights into the cognitive deficits that occur in AD.

**Relationship With NO: Adding Back the Human NOS2 Gene**

The unique nature of the CVN-AD mouse model is attributable, at least in part, to the primary role of NO in the redox regulation of the brain environment and to NO-mediated regulation of immune response in neurodegenerative diseases (107). The immune response is clearly a critical event in neurodegenerative processes in AD. Zhang et al (18) used an integrated network analysis of genes associated with late-onset AD to demonstrate that immune/microglia gene networks have the strongest degree of disease association and the highest correlation with AD neuropathology compared with other gene subsets. We have made mice that represent a phenotypic copy of the immune/redox environment by lowering iNOS-mediated NO production to zero (APPSwDI/mNos2−/− CVN-AD strain) and now show a second mouse model where the NO levels are not zero but instead represent more human-like NO levels. This was accomplished by reconstituting the human NOS2 gene into the APPSwDI/mNos2−/− strain. APP/huNOS2−/− mice contain gene DNA-regulatory sites associated with NO production in humans (Hoos et al, in press). Our data from APP/huNOS2−/− mice show characteristic pathologic features similar to those observed in CVN-AD mice, including accumulation of amyloid deposits throughout the parenchyma and around capillaries, the presence of hyperphosphorylated and sarkosyl-insoluble aggregates of tau, neuronal loss, and behavioral deficits. Both mouse strains display pathologic features that are not typically observed in other models and may be used to improve our understanding of AD.

The failure to demonstrate accompanying tau pathology when Aβ is overexpressed and the lack of neuronal death in the most commonly used AD models have been problematic. For example, despite abundant Aβ deposition in the subiculum and hippocampus, neuronal loss in the parent APPSwDI mouse strain and in many other AD mouse models is not measurable. Thus, extracellular Aβ deposition alone is apparently not a primary requirement for significant neuronal death. It is likely, however, that excess intraneuronal Aβ does damage neurons. This exception is found in the APP23, APP-PS1 double-transgenic, and APP-PS1K1 transgenic strains that coexpress high levels of mutated APP and PS1 (38, 61, 71). The mechanism of damage in these mice likely involves changes in Aβ handling within neurons and is clearly different from the CVN-AD strain, which shows lower-than-endogenous expression levels of APP. Accompanying tau pathology that is characteristic of AD is also not observed in these other mouse models and consequently limits further understanding of the sequence of events in disease progression. In contrast, our data from the CVN-AD model confirm that immune activation is a critical event and that NO participates in both tau and neuronal pathology. Changing NO levels may shift the vulnerability to tau aggregate formation, potentially by direct nitration of the tau protein (108, 109) and/or by indirect mechanisms such as differential regulation of caspase-3 activation (7). Low NO, as found in CVN-AD and APP/huNOS2−/− mice, is predicted to promote aggregate formation and is consistent with the increased level of sarkosyl-insoluble tau in both CVN-AD and APP/huNOS2−/− mice strains. The production of pathologic tau proteins in the entorhinal cortex to the subiculum to the hippocampus circuitry and the critical role of these pathways as an information transfer hub between the entorhinal cortex and the hippocampus may be a causative factor in the death of neurons in the CA1 and CA2/CA3 regions (106, 110–112). The maintenance of the distinct AD-like disease phenotype with the reconstitution of human NOS2 into the mouse Nos2 knockout background provides direct supportive data for the concept that the differences between human and mouse immune-based regulation of NO may be at least one critical factor that impacts the level and type of disease-related pathologies observed in mouse models of AD.

**REFERENCES**


