Prenatal choline supplementation attenuates neuropathological response to status epilepticus in the adult rat hippocampus

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Prenatal choline supplementation (SUP) protects adult rats against spatial memory deficits observed after excitotoxin-induced status epilepticus (SE). To examine the mechanism underlying this neuroprotection, we determined the effects of SUP on a variety of hippocampal markers known to change in response to SE and thought to underlie ensuing cognitive deficits. Adult offspring from rat dams that received either a control or SUP diet on embryonic days 12–17 were administered saline or kainic acid (i.p.) to induce SE and were euthanized 16 days later. SUP markedly attenuated seizure-induced hippocampal neurodegeneration, dentate cell proliferation, and hippocampal GFAP mRNA expression levels, prevented the loss of hippocampal GAD65 protein and mRNA expression, and altered growth factor expression patterns. SUP also enhanced pre-seizure hippocampal levels of BDNF, NGF, and IGF-1, which may confer a neuroprotective hippocampal microenvironment that dampens the neuropathological response to and/or helps facilitate recovery from SE to protect cognitive function.

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Status epilepticus (SE), a period of prolonged seizures, produces a host of plastic changes in the hippocampus that are thought to contribute to the development of temporal lobe epilepsy. SE results in substantial neuronal loss (Cavazos et al., 1994; Haas et al., 2001; Gorter et al., 1988), γ-aminobutyric acid (GABA) system alterations (e.g., Houser and Esclapez, 1996), reactive gliosis (Jorgensen et al., 1993; Niquet et al., 1994a; Kang et al., 2006), mossy fiber innervation of the dentate gyrus (Sutula et al., 1988; Ben-Ari and Represa, 1990), changes in levels of growth factors (Khrestchatisky et al., 1995; Mudo et al., 1996; Schmidt-Kastner et al., 1996; Shetty et al., 2004), and a transient increase in cell proliferation and neurogenesis (Bengzon et al., 1997; Parent et al., 1997; Scharfman et al., 2000; Hattiangady et al., 2004). These SE-induced degenerative and regenerative changes in the hippocampus are also accompanied by deficits in hippocampal-dependent learning and memory (Stafstrom et al., 1993; Liu et al., 1994; Sarkisian et al., 1997; Hort et al., 1999; Mikati et al., 2001). Understanding the relationship between the hippocampal response to SE and its cognitive consequences may lead to new insights into the treatment of SE and epilepsy.

Remarkably, dietary choline supplementation has been shown to protect rats from seizure-induced spatial memory retention deficits normally observed after SE (Yang et al., 2000; Holmes et al., 2002). Prenatal and early postnatal choline supplementation has also been shown to protect against impairments in performance on hippocampal-dependent tasks caused by aging (Meck et al., 1988, 1989; Meck and Williams, 2003; McCann et al., 2006) or neonatal alcohol exposure (Thomas et al., 2004, 2007; Wagner and Hunt, 2006). While the mechanism by which prenatal choline supplementation confers neuroprotection is not fully understood, choline is a vital nutrient important for several biological functions: acetylcholine synthesis, building biological membranes, cell signaling, and methyl donation (Blusztajn, 1998; Zeisel, 2004, 2006). Prenatal choline supplementation also enhances several features of adult hippocampal plasticity known to influence learning and memory function, such as increased baseline levels of neurogenesis, brain-derived neurotrophic factor (BDNF) (Glenn et al., 2007) and nerve growth factor (NGF) (Sandstrom et al., 2002); a reduced threshold to induce long-term potentiation (Pyapali et al., 1998; Jones et al., 1999); and enhanced depolarization-induced mitogen-activated protein kinase (MAPK) and cAMP-response element binding protein (CREB) activation (Mellott et al., 2004). Enhanced hippocampal plasticity may underlie prenatal choline supplementation’s neuroprotection of...
the hippocampus, rendering the hippocampus better able to withstand or recover from the deleterious effects of a neural insult.

Given that prenatal choline supplementation has been shown to protect against seizure-induced memory loss on a hippocampal-dependent task (Yang et al., 2000; Holmes et al., 2002), we hypothesized that prenatal choline supplementation alters the neurophysiological response of the hippocampus to SE. To investigate this, we used a model of excitotoxic injury to examine whether prenatal choline availability modulates a variety of markers known to change shortly after SE, including hippocampal histopathology, glutamic acid decarboxylase (GAD) expression, dentate cell proliferation, neurogenesis, astrogliosis, and growth factor content.

Materials and methods

Animals

Twenty timed-pregnant Sprague–Dawley rats (CD strain, Charles River, Kingston, NY) were obtained on day 9 of gestation (ED9). All dams were individually housed in clear polycarbonate cages (27.9×27.9×17.8 cm) that were individually ventilated, and the colony was maintained at 21 °C on a 12-hour light/dark cycle with lights on at 7 a.m. Dams were fed a control diet ad libitum (AIN76-A from Dyets, American Institute of Nutrition, ICN, Nutritional Biochemical, Cleveland, Ohio; 1.1 g/kg choline chloride substituted for choline bitartrate). Prenatal diet treatments were the same as those used in studies showing memory-enhancing and memory-protecting effects of prenatal choline supplementation (Meck et al., 1988, 1989; Yang et al., 2000; Holmes et al., 2002; Meck and Williams, 2003). On the morning of ED11 to the morning of ED18 (ED 11–17), pregnant dams were either given ad libitum access to a control diet (n=14) or a choline-supplemented diet (n=6). Control dams were given the AIN76-A diet containing 7.9 mmol/kg choline chloride and water sweetened with 50 mM saccharine, resulting in an average daily choline intake of 0.59 mmol/kg/day. Choline-supplemented dams received the AIN76-A diet containing 7.9 mmol/kg choline chloride and water containing 25 mM choline chloride and sweetened with 50 mM saccharine, resulting in an average daily choline intake of 3.46 mmol/kg/day (approximately 6.8 times more choline than the control diet). On ED18, all dams were returned to normal drinking water. There were no significant differences in the amount of water intake, food consumed, or body weights on ED11–18 between control and choline-supplemented dams (ps>.05; data not shown). After birth, offspring from the control and choline-supplemented dams were toe clipped for identification and then were selected randomly and cross-fostered to dams that consumed the control diet throughout pregnancy to yield 10 pups per litter (5 males and 5 females, half from different control dams and half from supplemented dams). There were no significant differences between control and prenatally choline-supplemented litter size or pup birth weights (ps>.05). On postnatal day (P) 25, pups were weaned and pair-housed with a rat of the same sex and prenatal diet condition. All offspring were given ad libitum access to the control diet through the duration of the study. Male offspring were used as subjects. All animal procedures were in compliance with the Institutional Animal Care and Use Committee of Duke University.

Induction of status epilepticus and bromodeoxyuridine injections

Status epilepticus procedures were adapted from previous reports and were based on a model of chronic temporal lobe epilepsy that yields a low mortality rate (Hellier et al., 1998; Hellier and Duke, 1999; Hattiangady et al., 2004). Kainic acid (KA) was obtained from Ocean Produce International (Shelburne, Nova Scotia, Canada) and was dissolved in 0.9% sterile saline (Sigma, St. Louis, MO). On P60, a group of adult male control offspring (CON, n=11) and prenatally choline-supplemented offspring (SUP, n=11) were injected with KA (2.5 mg/kg, i.p.) every hour and observations of seizure behavior were recorded according to Racine’s scale (Racine, 1972). A separate group of CON (n=4) and SUP (n=4) male offspring was similarly treated with hourly injections of 1 mL/kg saline. All KA-treated animals initially showed wet dog shakes and head-nodding, followed by class III (unilateral forelimb clonus), class IV (bilateral forelimb clonus with rearing), and class V (bilateral forelimb clonus with rearing and falling over) motor seizures. KA treatment continued until rats reached status epilepticus, which was defined as at least one class IV/V seizure per hour for 3 consecutive hours. If a rat displayed ≥10 class IV/V seizures in an hour, administration of the next injection was delayed to the next hour. If a rat displayed bouncing seizures after reaching status epilepticus, it was given a low dose of diazepam (2.5 mg/kg) to reduce the risk of mortality. We have observed and others have demonstrated (Scharfman et al., 2000; Pitkanen et al., 2005) that administering a low dose of diazepam following status epilepticus does not compromise the development of spontaneous epileptic motor seizures. The total KA dosage administered was titrated for each rat according to each rat’s motor seizure activity. Because it has been found that seizures, as opposed to neuronal cell death per se, lead to increases in hippocampal cell proliferation and neurogenesis (Parent et al., 1997; Tooyama et al., 2002; Smith et al., 2005) our procedures were designed such that all KA-treated rats experienced similar duration and severity of seizure activity.

After KA treatment, rats were injected subcutaneously with 5 mL of lactated Ringer’s solution (Abbott Laboratories, Chicago, IL) to prevent dehydration and were monitored closely until all seizure activity subsided. Saline-treated rats were also given 5 mL of lactated Ringer’s solution to equate their post-treatment experience to that of KA-treated rats. For the next 3 to 5 days, all rats continued to receive daily subcutaneous injections of 1–5 mL of lactated Ringer’s solution and were provided with moistened chow and slices of fresh fruit to aid recovery. Mortality rates did not differ significantly between CON rats (2/11) and SUP rats (3/11), $\chi^2 (1, n=22)=.26, p=.61$.

Five days after being treated with KA or saline, all rats were administered daily injections of 5-bromo-2-deoxyuridine (BrdU; 100 mg/kg/day, i.p.; Sigma, St. Louis, MO) for 10 consecutive days to label dividing cells. This injection regimen was based on past research designed to capture the impact of a variety of manipulations on cell proliferation and survival in the hippocampus (Kempermann et al., 1997; Lee et al., 2002a; Rao et al., 2005; Glenn et al., 2007), and to parallel the time point that prenatal choline protects against seizure-induced memory deficits (Yang et al., 2000; Holmes et al., 2002).

Tissue harvesting and cresyl violet staining

Twenty-four hours after the last BrdU injection, rats were given an overdose of a ketamine/xylazine cocktail, decapitated, and brains were rapidly removed and midsagittally sectioned. The selection of this particular time point (16 days post-KA/saline treatment) was based on 1) the findings reported in Hattiangady et al. (2004) that showed robust increases in hippocampal neurogenesis 16 days...
following an analogous KA treatment, and 2) prenatal choline supplementation’s protection against memory deficits was observed 1–2 weeks after excitotoxin-induced status epilepticus (Yang et al., 2000; Holmes et al., 2002), which approximates the current time window we have chosen. The hippocampus from one half-brain was immediately dissected for mRNA and protein analyses and stored at −80 °C until assayed. The other half-brain from each rat was immediately post-fixed in 4% paraformaldehyde for 72 h at 4 °C and then cryoprotected in a 30% sucrose solution in 1 M phosphate buffer (PB). These half-brains were then sectioned coronally at 60 µm on a microtome through the rostral–caudal extent of the hippocampus and every fifth section was collected in 0.1% sodium azide in 1 M PB to yield five series of 15–20 sections each. The first and second series were processed for BrdU and doublecortin (DCX) immunohistochemistry, respectively, for subsequent cell counting. Representative samples of four to five 60 µm sections that captured the rostral–caudal extent of the hippocampus were randomly selected from the third series and processed for cresyl violet staining. Sections stained for cresyl violet were analyzed and scored for lesion severity by 2 observers blind to the treatment conditions. The following scale, adapted from Yang et al. (2000) and Holmes et al. (2002), was used to characterize each rat’s hippocampal cytopathology: a score of 0 indicated no cell loss, a score of 1 indicated minimal cell loss in CA1, CA3, and/or hilar regions, a score of 2 indicated moderate cell loss with relatively normal general cellular architecture, and a score of 3 indicated severe cell loss with considerable disruption of the general cellular architecture.

BrdU immunohistochemistry

Immunohistochemical procedures for BrdU-labeling were based on the methods of Kuhn et al. (1996). One series of free-floating sections was rinsed with tris-buffered saline (TBS: pH 7.3) followed by 30 min in 50% methanol and 30 min in 0.6% hydrogen peroxide in TBS at room temperature to reduce nonspecific staining. After rinsing again in TBS, tissue was treated for 2 h in 50% Formamide/2× SSC (0.3 M NaCl, 0.03 M sodium citrate) at 65 °C, rinsed in 2× SSC for 10 min, incubated in 2 N HCl for 30 min at 37 °C, and rinsed in 0.1 M boric acid (pH 8.5) for 15 min. Sections were rinsed in TBS, incubated in 0.1% Triton X-100 (TTX; Sigma) and 3% normal horse serum (Vector Laboratories, Burlingame, CA) in TBS for 30 min at room temperature, and then incubated with the primary antibody (monoclonal mouse anti-BrdU, 1:400; Boehringer Mannheim, Indianapolis, IN) for 24 h at 4 °C. Following this, the tissue was rinsed with TBS and incubated with the secondary antibody (biotinylated horse anti-mouse, 1:200; Vector Laboratories) for 2 h at room temperature. The tissue was then rinsed in TBS, incubated in an avidin–biotinylated peroxidase complex (ABC, Vector Laboratories) for 1 h at room temperature, rinsed again in TBS, and treated for peroxidase detection with diaminobenzidine (Vector Laboratories, nickel intensified) for 5 min. Stained sections were mounted on gelatin-coated slides, dehydrated, counterstained with cresyl violet, and coverslipped.

Doublecortin immunohistochemistry

An adjacent series of tissue sets was processed for doublecortin (DCX) immunohistochemistry. These procedures were based on the methods of Rao and Shetty (2004). Free-floating sections were rinsed in TBS, treated for 30 min in 50% methanol, treated for 30 min in 0.6% hydrogen peroxide, incubated in 0.1% TTX and 3% normal horse serum in TBS for 30 min at room temperature, and then incubated with the primary antibody (affinity purified polyclonal goat antibody raised against a peptide mapping at the carboxy terminus of human DCX, 1:200; Santa Cruz Biotechnology, Santa Cruz, CA) for 4 h at room temperature. Following this, the tissue was rinsed with TBS and incubated with the secondary antibody (biotinylated horse anti-goat, 1:200; Vector Laboratories) for 2 h at room temperature. The tissue was then rinsed in TBS, incubated in an avidin–biotinylated peroxidase complex (ABC, Vector Laboratories) for 1 h at room temperature, rinsed again in TBS, and developed with vector grey substrate (Vector Laboratories). Stained sections were mounted on gelatin-coated slides, dehydrated, and coverslipped.

Quantification of BrdU- and DCX-labeled cells using unbiased stereology

BrdU- and DCX-labeled cells in each dentate gyrus were counted using the optical fractionator method (West, 1993, 1999; Mouton, 2002). We sampled every fifth section through the rostral–caudal extent of the dentate gyrus and used two separate sampling regions; one region encompassed the dorsal and ventral blades, including the granule cell layer and subgranular zone, and the other region included the hilus. StereoInvestigator (Microbrightfield Inc., Williston, VT) was used to sample systematically through the designated region and count numbers of labeled cells. For counting BrdU-labeled cells, we used a 40×40 µm counting frame and 14–41 sites per section were analyzed in 8 sections, yielding 146–232 frames per region for each rat. These parameters ensured adequate sampling of BrdU-labeled cells through the dentate gyrus. The same parameters were used for counting DCX-labeled cells with the exception of a larger counting frame (80×80 µm) to compensate for the more sporadic distribution of DCX-labeled cells, yielding 13–33 sites per section and 117–227 frames per region analyzed for each rat. For analysis we set an optical dissector height of 20 µm with a 2-µm guard zone and counted stained cells in each frame using a 40× objective lens. Finally, estimates of the volume of the region of dentate gyrus that was sampled for BrdU and DCX estimates were made using Cavalieri’s principle (Mouton, 2002). For each section examined, the area of the dentate gyrus was calculated by the StereoInvestigator software and was based on the boundaries of the contour tracings. Volume estimates were obtained by multiplying the section area estimates with the spacing between sampled areas. Spacing was derived by multiplying the measured, post-histology thickness of each sample by the number of sections examined.

ELISA for neurotrophic and growth factors

The entire hippocampus from the remaining half-brain from each rat was carefully dissected out, divided into 4–6 pieces along the rostral–caudal extent of the hippocampus, and half of the pieces were immediately stored at −80 °C until assayed. The remaining pieces were used for reverse transcriptase PCR assays (see Reverse transcriptase PCR section). For ELISA assays, all dissected samples were first weighed individually to get their wet weights. Whole tissue extracts were prepared by adding lysis buffer (50 mM Tris pH 7.5, 150 mM NaCl, 1% Nonidet NP-40, 10% glycerol, 2 mM 4-(2-aminoethyl)-benzenesulfonyl fluoride, 1 μg/mL leupeptin, 2 μg/mL aprotinin, 2 μg/mL pepstatin), followed by gentle sonication, incubation on ice for 15 min, and a brief centrifugation to clear. The supernatant from each sample was diluted 5 times with Dulbecco’s...
PBS and acidified to pH 2.6. After 15 min of incubation at room temperature, the diluted supernatants were neutralized to pH 7.6, aliquoted and frozen for subsequent measurement of brain-derived neurotrophic factor (BDNF), insulin growth-like factor-1 (IGF-1), fibroblast growth factor-2 (FGF-2), nerve growth factor (NGF), or neurotrophin-3 (NT-3) using ELISA. The above procedure was performed for all samples because a previous study indicated that acidification and subsequent neutralization with base increase the amount of detectable neurotrophins in extracts of CNS tissues (Okrably and Haak-Frendscho, 1997).

The Chemikine™ BDNF sandwich ELISA kit (Chemicon Int., Inc.) was used to assay the BDNF levels in hippocampal lysates. BDNF levels were measured according to manufacturer’s instructions. First, sample/standard diluent was added to each well of the microplate (100 µL/well). The standards were serially diluted (1:2) from 500–0 pg/mL. Standards and samples (100 µL/well) were added to wells in duplicate and were incubated overnight at 4°C. The wells were rinsed with wash buffer and then incubated with diluted biotinylated mouse anti-BDNF monoclonal antibody (1:1000) for 2 h at room temperature. After rinsing the plate again with wash buffer, diluted streptavidin–HRP conjugate solution (1:1000) was added to wells and incubated for 1 h at room temperature. Plates were again washed and then warm TMB buffer was added. After 15 min incubation at room temperature, stop solution was added.

The Quantikine® sandwich ELISA kits (R&D Systems) were used to assay IGF-1 and FGF-2 levels in the samples. ELISAs were performed according to manufacturer’s instructions. Briefly, microplates were pre-coated with the first primary antibody (monoclonal in IGF-1 and FGF-2 kits). Assay diluents were added to each well of the microplate (50 µL/well for IGF-1 assay and 100 µL/well for FGF-2 assay). Standard control samples for each neurotrophic factor were diluted serially (1:2) from either 6–0 ng/mL (IGF-1) or 32–0 pg/mL (FGF-2) or with their respective calibrator diluents and plated to two columns of wells (50 µL/well in IGF-1 assay and 100 µL/well in FGF-2 assay) designated for standard curve in every plate. The frozen ELISA samples (described above) were thawed on ice, and every sample plated in duplicate for measurement of each of the factors. Following a 2-h incubation at room temperature, wells were rinsed in wash buffer and treated with an enzyme-linked second primary antibody solution for 2 h. The second primary antibody was a polyclonal IGF-1 antibody conjugated to HRP in IGF-1 detection kit or a monoclonal FGF-2 antibody conjugated to alkaline phosphatase in FGF-2 detection kit. The wells were rinsed in wash buffer and a substrate solution was added to the wells and incubated in the dark for 30 min for IGF-1 or 45 min for FGF-2. The color reaction was stopped with 1 M hydrochloric acid in the IGF-1 assay or with 2 N sulfuric acid in the FGF-2 assay.

The Emax® immunoassay systems (Promega) were used to measure NGF and NT-3 in the samples. Flat-bottom 96 well plates (NUNC) were first coated with solution containing a polyclonal antibody against either NGF or NT-3 (the first primary antibody solution) prepared in carbonate coating buffer (100 µL/well, 1:1000 dilution) and incubated for 16 h at 4°C. Following a wash in TBST (Tris-buffered saline solution containing Tween 20), the coated wells were incubated with block and sample buffer (1X) for 1 h at room temperature and washed again with TBST. Standard control samples for NGF and NT-3 were diluted serially (1:2) from 250–0 pg/mL or 300–0 pg/mL, respectively, and plated in duplicate (100 µL/well). The frozen ELISA samples (described above) were thawed on ice, and every sample plated in duplicate for measurement of NGF or NT-3. Following a 6-hour incubation at room temperature, wells were washed in TBST. Diluted monoclonal antibody against either NGF or NT-3 (the second primary antibody solution; 1:4000) was added to each well and incubated overnight at 4°C. The wells were washed in TBST, incubated with appropriate secondary antibody conjugated to peroxidase for 2.5 h, washed again in TBST, and treated with tetramethyl benzidine (TMB) substrate for 10 min. The chromogen reaction was stopped by adding 100 µL of 1 N hydrochloric acid.

The optical density of each well was measured using the Victor3 microplate reader (PerkinElmer Life Sciences). The intensity of color was measured at a wavelength of 450 nm for all ELISAs. In order to correct for optical imperfections in the plate, readings at 540 nm were subtracted from readings at 450 nm. The standard curve was used to assess the validity of the protocol and to determine the relative concentrations of the growth factors. Values in all samples were normalized per gram of tissue assayed, and the average value for each sample was calculated separately before determining the group means.

**Western blot analysis for GAD65, GAD67, and GFAP**

For Western blot analysis, the lysates were prepared as described in the ELISA methods. However, a portion of the lysate was taken before dilution in Dulbecco’s PBS. The extracts were normalized for total protein and subjected to SDS-PAGE. After transfer of protein to an Immobilon P membrane (Millipore), the membrane was blocked with 5% nonfat dry milk in 1X Tris-buffered saline (TBS) containing 0.1% Tween 20 for 2 h and then probed overnight with either anti-gliaal fibrillary acidic protein (GFAP) monoclonal antibody GAS (1:1000) (Cell Signaling Technology), anti-GAD65/67 polyclonal antibody (1:1000) (Chemicon), or anti-beta-actin monoclonal A5441 (1:5000; Sigma). The antibody/antigen complexes on the membranes were detected using a peroxidase-conjugated anti-mouse IgG for GFAP and beta-actin (1:2000) or anti-rabbit IgG (1:5000) for GAD65/67 and visualized using the enhanced chemiluminescence method (Western Lightning, Perkin Elmer) and a Kodak Image Station 440. Digitized images of immunoblots were quantified using Kodak ID software. Protein levels were normalized with beta-actin values.

**Reverse transcriptase PCR**

The remaining pieces of the hippocampus dissected out from the remaining half-brain from each rat were immediately homogenized in cold guanidine isothiocyanate solution, frozen on dry ice, and stored at −80°C. Total RNA was extracted from tissues by phenol and chloroform method (Chomczynski and Sacchi, 1987) and precipitated. RNA was resuspended and its quantity was determined using Quant-iT™ RiboGreen® RNA assay kit (Molecular Probes) and the Victor™ multi-label plate reader (PerkinElmer Life Sciences). Hippocampal RNA was used for reverse transcriptase (RT)-PCR using Superscript One-Step RT-PCR with Platinum Taq (Invitrogen). First strand cDNA synthesis was performed with 25 ng of total RNA, oligo dT primers and reverse transcriptase at 48°C (45 min). Primers used for PCR include beta-actin (Forward: CAC AGC TGA GAG GAA AAT C, Reverse: TCA GCA ATG CCT GGG TAC), DCX (Forward: CGATCA AAC TGG AAA CCG C, Reverse: TTT GCG TCT TGG TCG TTA CC), BDNF (Forward: TGT AGC AGC TGA TCG AAC AGC, Reverse: GTT CGG CAT TGC GAG TTC CAG), IGF-1 (Forward: GTG GAC GCT CTT CAG TAC CT, Reverse: TGT GAC GCT CTT CAG TAC CT).
Revers: GCT TCT TCT TGT GTG TCG ATA G, FGF-2
Forward: GCC TAC CTG CGC ATC CA, Reverse: GCT CTT CAC TCA CCC AC, Reverse: AGG CCA CTG ACT AGG CTG AA, NT-3
Forward: CAC CCA GAG AAC CAG AGC AG, Reverse: TCT GAA GTC AGT GCT CGG AC), GAD65 (Forward: TCT TTT CTC CTG GTG GTG CC, Reverse: CCC CAA GCA GCA TCC ACA T), GAD67 (Forward: TACGTTCCGACAGGTC, Reverse: CCCCCAAGCACGATCCACAT), and GFAP (Forward: ACA TCG AGA TCG CCA CCT AC, Reverse: ACA TCA CAT CCT TGT GCT CC). PCR was performed using Platinum Taq DNA polymerase with a denaturing step for 2 min at 94 °C, followed by 28–40 cycles (36 cycles for β-actin; 32, DCX; 35, BDNF; 40, IGF-1; 30, GAD65; 38, GAD67; 38, GAD67; and 28, GAD65) of 1 min at 94 °C, 1 min at 55 °C and 2 min at 72 °C, and terminated by an elongation step at 72 °C for 7 min. PCR products were separated on a 10% TBE polyacrylamide gel and stained with ethidium bromide. PCR products were then visualized with the Kodak Image Station 440 (Rochester, NY) and product intensities were quantified using Kodak software. PCR product levels were normalized with β-actin values.

Statistical analyses

Numbers of BrdU-labeled and DCX-labeled cells estimated with the optical fractionator, the volumes of dentate gyrus that were estimated using Cavalieri’s principle, and protein and mRNA levels (expressed as percent of control levels) were all subjected to a 2 (Diet: CON vs. SUP) × 2 (Treatment: saline vs. KA) between-subjects ANOVA. Of particular interest were Diet × Treatment interactions. Where appropriate, a priori pairwise comparisons were used to evaluate differences between group means. A significance level of .05 was set for all statistical tests. Values are reported in the text as means ± SEM. Note that subjects contained within each experimental condition were randomly selected from different litters (n of 1/litter). Thus, we have taken the necessary precautions to be sure that our findings are not contaminated by a lack of within-litter variability.

Results

Seizure activity and hippocampal histopathology after KA-induced SE

All rats treated with KA reached SE and exhibited continuous class III–IV seizures for over 3 h. Careful observation of behavioral activity of rats receiving KA revealed that the pattern of seizure activity (progression, number, and severity of seizures) was similar for both CON and SUP rats. There were no significant differences between CON and SUP rats in the number of class III/IV/V or total number of motor seizures within the first 1-hour period of observable seizure activity (CON: class III = 0.25 ± 0.25, class IV = 7.00 ± 1.41, class V = 9.25 ± 3.61, total = 16.50 ± 4.91; SUP: class III = 4.00 ± 1.47, class IV = 10.00 ± 2.35, class V = 8.50 ± 3.52, total = 22.50 ± 6.81; all ps > .05). After exhibiting the first motor seizure, all rats entered a continuous class III–V seizure state in the second 1-hour period. To

Fig. 1. Histopathology of the hippocampus 16 days following KA-induced SE in CON (B) and SUP (C) rats. Panel A depicts representative sections of CA1 (A1), CA3 (A2), and dentate gyrus (A3) regions from an intact hippocampus from a saline-treated CON rat. Saline-treated SUP rats also did not show any lesions (histology data not shown). Note more severe degeneration (cell loss, disruption of cytoarchitecture, and gliosis) in KA-treated CON rats (B1, B2, B3). Areas of damage are indicated by arrows. Dentate hilar cell loss was observed in both CON (B3) and SUP rats (C3). Photomicrographs in each set were taken with a 10× objective. Bars indicate 50 µm. GCL, granule cell layer. SGZ, subgranular zone. H, hilus.

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ensure that differences in seizure severity or duration did not occur beyond the 3-hour time window we used to define SE, we continued to monitor both CON and SUP rats. No differences were apparent. All rats gradually dropped out of SE within 2–3 h of the last injection: continuous seizures subsided, followed by the emergence of 1–3 discrete Class III/IV/V seizures within a 1-hour period, and the absence of seizure activity by 5 h after the last injection. There were also no significant differences between CON and SUP rats in the total amount of KA needed to induce SE (range=3.75–16.25 mg/kg) or in the latency to the first motor seizure (range=62–331 min). This variability in dose and latency to the first behavioral seizure in our rats is consistent with previous reports showing that the Sprague–Dawley rats show a more variable convulsant response to KA than other rat strains (Golden et al., 1991, 1995). Thus, while the dose of KA needed to induce SE varied across rats, the SE produced was quite comparable between our CON and SUP rats.

We used histopathology scores adapted from a previous report showing protection against histopathology following KA treatment as a result of prenatal choline supplementation (Holmes et al., 2002) to confirm that our KA treatment lead to similar patterns of histopathology between CON and SUP rats. Comparable to Holmes et al. (2002), we also found evidence of considerable histopathology from CON rats that experienced KA-induced SE, and this damage also appeared to be attenuated in SUP rats. Fig. 1 shows photomicrographs of representative brain sections stained with cresyl violet from CON and SUP rats 16 days following KA treatment. While the granule cell layer of the dentate gyrus remained intact, areas CA1, CA3, and the hilus of KA-treated rats displayed cell loss, disrupted cytoarchitecture, and gliosis. The histopathology of the hippocampus was more severe in KA-treated CON rats when compared to KA-treated SUP rats (Figs. 1B, C). KA-treated SUP rats also had significantly lower histopathological scores than KA-treated CON rats (CON: 2.54±.40, SUP: 1.44±.37, t(6)=2.01, p<.05, one-tailed). To further examine the nature of the neuro-pathological response to KA-induced seizures in CON and SUP rats, we quantified a number of neurobiological markers in the hippocampus known to change as a result of SE and possibly contribute to the spatial learning and memory deficits observed shortly after seizures.

Prenatal choline supplementation protects against SE-induced decrease in hippocampal GAD65, but not GAD67, protein and mRNA expression

Excitotoxin-induced SE is known to cause a decrease in inhibition (Sloviter, 1987; Cornish and Wheal, 1989; Milgram et al., 1991; Kobayashi and Buckmaster, 2003) and degeneration of GABAergic neuronal function in the hippocampus (Obenaus et al., 1993; Houser and Esclapez, 1996; Morin et al., 1998; Esclapez and Houser, 1999; Shetty and Turner, 2001), which has been considered to be a significant contributor to hyperexcitability in the epileptic hippocampus. This seizure-induced disruption in GABAergic function can be measured via quantification of GAD65 and 67, enzymes important for local GABA synthesis at synaptic (GAD65) and cytoplasmic (GAD67) sites (Erlander and Tobin 1991; Esclapez et al., 1994), and may play a role in the packaging and release of GABA (Namchuk et al., 1997; Tian et al., 1999). Thus, as a marker of GABAergic function, we examined levels of hippocampal GAD65 and 67 proteins (via Western blot) and mRNA in saline- and KA-treated CON and SUP rats 16 days post-SE and these data, expressed as percent of control levels, are shown in Fig. 2. Analyses of mRNA levels revealed a significant Diet×Treatment interaction for GAD65 mRNA, F (1, 12)=17.19, p<.001. Baseline GAD65 mRNA levels were 33% lower in SUP rats than in CON rats (p<.05; Fig. 2B). Sixteen days following SE, GAD65 mRNA levels were reduced by more than half in CON rats (p<.001), which is consistent with previous reports that show decreases in hippocampal GAD65 mRNA 3–14 days following seizures (Houser and Esclapez, 1996; Kobayashi and Buckmaster, 2003). In contrast, there was no significant difference in GAD65 mRNA levels between saline-treated and KA-treated SUP rats (Fig. 2B). Although a Diet × Treatment interaction did not reach statistical significance for GAD65 protein levels, F(1, 10)=2.57, p=.14, planned comparisons revealed that KA-treated CON rats showed a significant decrease by more than half in GAD65 protein (p<.05) whereas GAD65 protein levels did not differ significantly between KA-treated and saline-
treated SUP rats (Fig. 2A). Taken together, these data suggest a preservation of hippocampal GABAergic GAD65-containing neurons, or a more rapid recovery of GAD65 in SUP rats. In contrast, neither prenatal diet nor KA treatment had an effect on levels of GAD67 mRNA 16 days following SE (all Fs < 1), as levels of GAD67 mRNA did not significantly differ across the four conditions (Fig. 2B). Analyses of GAD67 protein levels, however, revealed that KA-induced SE led to a reduction in GAD67 protein levels in both CON rats and SUP rats, $F (1, 10) = 4.78, p = .05$ (Fig. 2A). SUP rats also exhibited lower levels of GAD67 protein overall, $F (1, 10) = 8.43, p < .02$. These findings suggest that prenatal choline supplementation’s neuroprotective effect on GAD expression 16 days following seizures may be specific to the GAD65 isoform.

Prenatal choline supplementation attenuates SE-induced upregulation of hippocampal cell proliferation in the dentate gyrus

To detect changes in dentate cell proliferation as a result of KA-induced SE, we visualized cells immunopositive for the cell division marker, BrdU, administered 5–16 days post-SE to capture the time window during which savings in spatial learning and memory following excitotoxin-induced SE have been observed with prenatal choline supplementation (Yang et al., 2000; Holmes et al., 2002). The amount of cell proliferation after SE that we captured with our 10-day BrdU regimen yielded a very high density of immunostaining. Stereological procedures also require relatively thick sections for unbiased cell counting, which further contributes to our high density of BrdU labeling where many BrdU+ cells following SE were tightly clustered together and layered on top of each other, making the quantification of double labeled cells unreliable. Therefore, we quantified markers of neurogenesis and gliogenesis in the hippocampus (see additional results below). Fig. 3 shows photomicrographs of BrdU labeling in the hippocampus of representative sections from CON and SUP saline- and KA-treated rats. BrdU-labeled cells were expressed throughout the rostral-caudal extent of the dentate gyrus and in both the dorsal and ventral blades. BrdU-labeled cells were evident in the subgranular zone, indicating that many of the cells were newly generated. BrdU-labeled cells were also visible in the granule cell layer, indicating that some had survived for several days and were migrating from the subgranular zone. Moreover, a vast majority of BrdU+ cells displayed morphological characteristics of normal non-pyknotic cells (e.g., round or oval nuclei that did not appear highly condensed). In KA-treated rats, a considerable number of BrdU+ cells were also present in the hilus. BrdU labeling clearly showed that while KA treatment dramatically upregulated dentate and hilar cell proliferation in CON rats, which is consistent with previous reports (Bengzon et al., 1997; Parent et al., 1997; Jessberger et al., 2005), the SE-induced increase in cell proliferation was significantly attenuated in SUP rats (Fig. 3B, D). This observation was confirmed by quantification of BrdU+ cells. Using unbiased stereology, estimates of the number of BrdU+ cells were generated for both the dentate gyrus and hilus for each rat and these data are shown in Fig. 4. As expected, SE significantly increased the number of dentate and hilar BrdU+ cells in both CON and SUP rats (dentate gyrus: $F (1, 12) = 24.73, p < .001$; hilus: $F (1, 12) = 32.47, p < .001$). There were no main effects of Diet on dentate nor hilar cell proliferation but a significant Diet×Treatment interaction was found for dentate cell proliferation, $F (1, 12) = 7.91, p < .02$, which approached statistical significance for hilar cell proliferation, $F (1, 12) = 3.67, p = .07$. Remarkably, while CON rats exhibited a 780% increase in the total number of BrdU+ cells (dentate gyrus and hilus combined) following SE, SUP rats showed only a 207% increase. Planned comparisons revealed that KA-treated SUP rats had significantly more BrdU+ cells in the dentate gyrus than KA-treated SUP rats ($p < .01$; Fig. 4), confirming that SE-induced cell proliferation in the dentate gyrus was markedly attenuated in SUP rats.

To determine whether a larger overall size of structure could account for the higher numbers of new cells detected in KA-treated CON versus KA-treated SUP rats, we estimated the volume of the dentate gyrus and hilus in each rat using Cavalieri’s principle (Mouton, 2002) and did not detect any statistically significant differences between any treatment groups ($F s < 1$).
SE-induced hippocampal neurogenesis is not modulated by prenatal choline supplementation

To determine whether prenatal choline supplementation decreased the characteristic proliferation of new neurons after SE, we used adjacent tissue sections to visualize cells immunopositive for the microtubule-associated phosphoprotein, doublecortin (DCX), that is transiently expressed in newly born neurons that are still in the process of migrating and differentiating (Brown et al., 2003; Rao and Shetty, 2004). Others have confirmed that DCX expression is a reliable indicator of neurogenesis in the adult brain (Couillard-Despres et al., 2005). As a secondary measure, we also quantified DCX mRNA levels via RT-PCR. Fig. 5 shows photomicrographs of DCX labeling in the hippocampus of representative sections from CON and SUP saline- and KA-treated rats. DCX+ cells were visible with processes in various stages of development and were evident along the subgranular zone and granule cell layer. In KA-treated rats, DCX+ cells were also detected in the hilus, which is consistent with previous studies that have shown that a portion of these newly generated neurons after SE aberrantly migrate to the hilus region (Parent et al., 1997; Hattiangady et al., 2004; Jessberger et al., 2005). In comparison to saline-treated rats many of the DCX+ neurons in KA-treated rats appeared displaced and exhibited abnormal morphological features, such as horizontally oriented cell bodies and processes (Figs. 5C, D).

Unbiased stereological estimates of the number of DCX+ neurons in the dentate gyrus and hilus were generated for each rat and are shown in Fig. 6. As expected, SE induced a dramatic increase in the number of DCX+ neurons in both CON and SUP rats (dentate gyrus: $F(1, 12)=11.93$, $p<.01$; hilus: $F(1, 12)=34.50$, $p<.001$). However, prenatal choline supplementation did not significantly diminish the proliferation of new neurons in response to SE; there was no significant main effect of Diet for both dentate and hilar neurogenesis or Diet x Treatment interactions ($F_s<1$). There was also no significant difference in the number of DCX+ neurons between saline-treated CON and saline-treated SUP rats. SE also produced a significant increase in DCX mRNA in both CON and SUP rats ($F(1, 12)=6.49$, $p<.05$), but prenatal choline availability did not alter baseline or SE-induced mRNA levels of DCX ($F_s<1$; Fig. 6). These data revealed that unlike the pattern of BrdU immunolabeling, seizure-induced upregulation of hippocampal neurogenesis was not modified by prenatal choline availability.

**Prenatal choline supplementation attenuates SE-induced upregulation of hippocampal GFAP protein and mRNA expression**

Prior work has shown that SE triggers astroglial activity as revealed by dramatic elevations of GFAP protein and mRNA expression (Jorgensen et al., 1993; Niquet et al., 1994a). GFAP is also expressed in progenitor cells that reside in the subgranular zone of the dentate gyrus and give rise to new neurons (Seri et al., 2001), which also makes GFAP a marker of interest given that SE...
produces robust increases in dentate cell proliferation. We quantified levels of hippocampal GFAP protein (via Western blot) and mRNA in saline- and KA-treated CON and SUP rats 16 days post-treatment and these data, expressed as percent of control levels, are shown in Fig. 7. As predicted, CON and SUP rats exhibited a significant increase in GFAP protein and mRNA levels as a result of KA treatment (protein: \( F(1, 12) = 16.60, p < 0.01 \); mRNA: \( F(1, 12) = 35.37, p < 0.05 \)). More importantly, there was a significant Diet \times Treatment interaction for GFAP mRNA levels, \( F(1, 12) = 5.88, p < 0.05 \), although a Diet \times Treatment interaction for GFAP protein levels did not reach statistical significance, \( F(1, 12) = 2.83, p = 0.12 \). Saline-treated SUP rats had significantly lower levels of hippocampal GFAP protein and mRNA than CON rats \( (p < 0.05) \). However, SE-induced increases in GFAP mRNA were significantly reduced in KA-supplemented rats. mRNA levels rose 3.5-fold in CON rats, but only doubled in SUP rats \( (p = 0.02) \). A similar, but not significant, pattern of change was observed for GFAP protein levels: a 3.1-fold increase following KA in CON rats, but only a 2.6-fold increase in SUP rats \( (p = 0.07) \). Collectively, these data suggest that SE-induced upregulation in GFAP levels may be attenuated in SUP rats.

**Growth factor expression in the intact and KA-lesioned hippocampus as a function of prenatal choline availability**

The expression of growth factors is thought to be an important mediator of hippocampal neurogenesis (Cameron et al., 1998; Anderson et al., 2002; Lee et al., 2002a,b) and some factors are known to change in response to status epilepticus (Gall and Isackson, 1989; Gall et al., 1991; Mudo et al., 1996; Schmidt-Kastner et al., 1996; Katoh-Semba et al., 1999; Shetty et al., 2004). We therefore quantified hippocampal protein levels (via ELISA) and mRNA levels (via RT-PCR) of the following growth factors known to influence neurogenesis: BDNF, IGF-1, FGF-2, NGF, and NT-3. Fig. 8 shows the relative protein levels and mRNA levels for each factor in CON and SUP saline- and KA-treated rats 16 days post-treatment (protein levels expressed as percent of control levels because protein concentrations varied greatly between growth factors).

Because we were ultimately interested in the potential effect of prenatal choline availability on hippocampal growth factor content in baseline conditions and in response to SE, we conducted planned comparisons between saline-treated CON and saline-treated SUP rats, as well as between saline-treated and KA-treated rats within each diet condition. In adult rats that did not experience seizures, protein levels of BDNF \( (p < 0.01) \), IGF-1 \( (p < 0.05) \), and NGF \( (p < 0.05) \) were significantly higher in prenatal SUP rats than saline-treated CON rats. Although differences between saline-treated CON and SUP rats in protein levels of FGF-2 \( (p = 0.29) \) and NT-3 \( (p = 0.13) \) failed to reach statistical significance, the respective means are in a similar direction (Fig. 8A). Collectively, these results suggest that SUP rats have a higher baseline hippocampal protein content of several neurotrophic factors.

As expected, SE significantly increased protein levels of BDNF \( (p < 0.01) \), IGF-1 \( (p < 0.01) \), FGF-2 \( (p < 0.01) \), NGF \( (p < 0.01) \), and NT-3 \( (p < 0.05) \) in CON rats. In contrast, SUP rats exhibited a significant increase only in BDNF \( (p < 0.01) \) and IGF-1 \( (p < 0.01) \) in response to SE (Fig. 8A). There was a similar trend for a SE-induced increase in FGF-2 in SUP rats \( (p = 0.06) \). It is important to note that because protein levels of BDNF and IGF-1 were significantly elevated in saline-supplemented rats, the increase in these neurotrophic factor levels following SE in supplemented rats (51% increase in BDNF; 59% increase in IGF-1) was not as large as it was in CON rats (BDNF: 72% increase; IGF-1: 81% increase). Together, these data suggest that prenatal choline supplementation either prevents or attenuates protein expression of some growth factors in response to SE.

We also quantified via RT-PCR the relative hippocampal mRNA levels of each growth factor (Fig. 8B). These data revealed a slightly different pattern of results than that of protein levels. While baseline mRNA levels were not significantly altered by prenatal choline supplementation \( (p > 0.05) \), prenatal choline availability had striking effects on SE-induced alterations in BDNF, IGF-1, and NT-3. BDNF mRNA levels were significantly elevated in KA-treated SUP rats in comparison to KA-treated CON rats \( (p < 0.01) \); SUP rats had a 29% increase in BDNF mRNA levels following SE whereas CON rats showed virtually no change in BDNF mRNA levels. Similarly, IGF-1 mRNA levels were also significantly higher in KA-treated SUP rats than in KA-treated CON rats \( (p < 0.05) \). There was an unexpected Diet \times Treatment interaction for NT-3 mRNA levels where CON rats showed an

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**Fig. 6.** Mean (±SEM) numbers of DCX-labeled cells (A) and mean (±SEM) mRNA expression levels (B) detected in the dentate gyrus and hilus of CON (white bars) and SUP (grey bars) rats 16 days following saline treatment (open bars) or KA-induced SE (hatched bars). KA-induced SE significantly increased the number of DCX-labeled neurons and levels of DCX mRNA in both CON and SUP rats \( (p < 0.05) \). Prenatal choline supplementation did not alter the number of DCX-labeled neurons or levels of DCX mRNA in response to KA-induced SE. * statistically different from within-diet saline-treated group.
increase in NT-3 mRNA following SE, whereas the opposite pattern was revealed for SUP rats, \( F(1, 12) = 11.75, p < .01 \) (Fig. 8B). These data suggest that in addition to affecting levels of growth factor protein expression, prenatal choline availability also modulates levels of growth factor mRNA following SE.

**Discussion**

We have shown that increased availability of choline during early development produced a long-lasting change in the fetal brain that led to a dramatically reduced hippocampal response 16 days following KA-induced SE. Compared to control rats, adult offspring of dams that were fed a choline-enriched diet for just 1 week, mid-gestation, responded to SE with: 1) reduced hippocampal histopathology, 2) robust protection against loss of GAD65 protein and mRNA expression, 3) marked attenuation of dentate cell proliferation, 4) dramatic reduction of GFAP mRNA expression (with a tendency for GFAP protein to be reduced as well), and 5) an altered pattern of growth factor expression. These data are particularly important because prior work has shown savings in hippocampal-dependent learning and memory at a similar time point following excitotoxin-induced SE in prenatally choline-supplemented rats (Yang et al., 2000; Holmes et al., 2002), suggesting that increased choline availability to the fetus may alter neural development such that the adult brain is protected from SE-induced damage and cognitive impairment. While these data do not rule out the possibility that choline supplements may have neuroprotective actions at other periods, we know that choline has profound memory-enhancing effects if it is supplemented on ED12-17 or PD15-30, but not at other developmental or adult timeframes (Meck and Williams, 2003).

Consistent with previous findings (Holmes et al., 2002), we report that prenatally choline-supplemented rats (SUP) showed reduced cell loss and gliosis in CA1, CA3, and hilus compared to...
control-fed rats even though all rats experienced the same severity and duration of seizure activity. While widespread neurodegeneration of the hippocampus is a notorious consequence of prolonged seizures (Niquet et al., 1994a; Fujikawa et al., 2000; Kotloski et al., 2002), GABAergic neurons are particularly vulnerable (Obenauer et al., 1993; Houser and Esclapez, 1996; Esclapez and Houser, 1999; Shetty and Turner, 2001). Thus, our finding that prenatal choline supplementation prevented the loss of hippocampal GAD65 protein and mRNA (though not GAD67 response to SE) is particularly interesting. Our findings may represent a protection of GABAergic neurons following SE, although loss of functional inhibition may not be due to a loss of inhibitory neurons per se (Sloviter, 1987; Davenport et al., 1990; Esclapez and Houser, 1999). Rather, a loss in GAD65 expression may contribute to a disruption in GABAergic function, as mice deficient in GAD65 are susceptible to epileptic seizures (Kash et al., 1997). Alternatively, our SUP rats may show an early restorative upregulation in GAD65 production. Indeed, Houser and Esclapez (1996) have found that remaining GABAergic hippocampal neurons upregulate GAD65 mRNA 2–4 months, but not 3–14 days, following seizure-induced neuronal damage. Future studies are needed to determine if the savings seen in our SUP rats are in numbers of GABAergic neurons or in altered recovery of GAD expression or both.

SE produces a robust reactive astrogial response in the hippocampus, yet our data reveal that seizure-induced increases in GFAP mRNA and dentate cell proliferation were significantly reduced in SUP rats. There was also a tendency for GFAP protein levels and hilar cell proliferation levels to be attenuated as well. SE-induced reactive astrogliosis plays a significant role in the aberrant synaptic remodeling of the hippocampus thought to underlie the development of epilepsy (Jorgensen et al., 1993; Niquet et al., 1994a,b; Represa et al., 1995). Reactive astrocytes that proliferate in response to seizures express several genes that promote neurite outgrowth (Represa et al., 1993; Niquet et al., 1994a,b; Stringer, 1996) and the increase in the expression of these genes coincides with both the emergence of mossy fiber sprouting and the period when both GFAP and hippocampal cell proliferation are elevated (Khrestchatisky et al., 1995; Represa et al., 1995; Bengzon et al., 1997; Parent et al., 1997; Borges et al., 2006). An attenuation of seizure-induced astrogliosis in SUP rats may thus indicate a reduced progression of epilepsy following SE.

We did not detect any differences between diet groups in dentate and hilar neurogenesis following SE. Previous work has shown that dentate neuronal and astroglial proliferation following SE likely arises from independent progenitor pools: inhibiting hippocampal neurogenesis before SE does not prevent astrogliosis or mossy fiber sprouting (Parent et al., 1999). Thus, prenatal choline supplementation may have a preferential effect on just one type of proliferative hippocampal response following SE. Although we did not detect changes in the number of DCX+ dentate and hilar neurons following seizures as a function of prenatal choline availability, it remains to be determined whether prenatal choline supplementation alters specific properties (e.g., migration pattern, morphology, neuronal functionality) of newborn neurons following seizures that are hypothesized to contribute to the development of epilepsy (Scharffman, 2004; Scharffman and Gray, 2007).

In order to further investigate the possible mechanisms of prenatal choline-induced neuroprotection, we examined expression of neurotrophic factors that are known to support hippocampal plasticity and are regulated by seizure activity. It is well established that 1 h to 4 days after SE, hippocampal NGF and BDNF are transiently increased while NT-3 levels are downregulated (e.g., Gall et al., 1991; Rocamora et al., 1994; Marcinkiewicz et al., 1997; Shetty et al., 2004). Less is known about growth factor responses beyond one week after SE, but hippocampal NGF and NT-3 proteins are still upregulated 45 days following SE while BDNF returns to baseline (Shetty et al., 2003). Our study adds to this timeline by demonstrating large increases in hippocampal BDNF, IGF-1, FGF-2, NGF, and NT-3 proteins 16 days post-SE in control-fed rats. Thus, the hippocampal growth factor response to SE may be characterized by periods of elevation and/or reduction in levels that are specific for each factor. Levels of BDNF and IGF-1 protein were significantly elevated in both SUP and CON rats following KA treatment, although the percent increase in these growth factors following SE compared to baseline levels was reduced in the SUP rats.

These data may suggest that the response of these growth factors to SE is attenuated, or that some ceiling prevents further increases in the SUP rats. The latter hypothesis seems somewhat unlikely because we find that the NGF and NT-3 protein response to SE is completely attenuated in the KA-treated SUP rats compared to KA-treated CON rats, suggesting that prenatal choline supplementation reduces the dramatic elevation of several growth factors following SE. Previous work has shown that blocking NGF’s actions prior to kindling both retards kindling and prevents mossy fiber sprouting (Rashid et al., 1995; Van der Zee et al., 1995); therefore, the attenuated NGF response to KA-induced in SUP rats may confer protection against mossy fiber sprouting. Additionally, BDNF and IGF-1 hippocampal mRNAs following SE were greater in SUP than CON rats, while NT-3 mRNA was lower in SUP rats following SE. This difference between protein and mRNA responses to SE may occur because mRNA is present within the intrinsic cells of the tissue, whereas protein may also be additionally contributed by the axons and terminals of theafferent projections to the hippocampus. Whether the muted response of some growth factors to SE in our supplemented rats underlies our finding of reduced cell proliferation following SE remains to be determined.

What may be key to understanding the neuroprotection conferred by prenatal choline supplementation that we report here, as well as the choline-induced protection against memory deficits following seizures reported previously (Yang et al., 2000; Holmes et al., 2002), is an altered hippocampal microenvironment inSUP rats prior to SE. SUP rats showed enhanced baseline levels of several growth/ neurotrophic factors (BDNF, NGF, IGF-1) in the hippocampus, compared to CON rats. This finding extends our previous demonstrations of higher BDNF protein in 8-month old (Glenn et al., 2007) and NGF protein in 20- and 90-day old (Sandstrom et al., 2002) female offspring of choline-supplemented dams. Infusions of BDNF or bFGF into the hippocampus prior to SE (Liu et al., 1993; Liu and Holmes, 1997; Reibel et al., 2000) as well as exercise (Setkowicz and Mazur, 2006), which stimulates BDNF, NGF, and IGF-1 (Gomez-Pinilla et al., 1998; Cotman and Berchtold, 2002), have been shown to ameliorate the progression of excitotoxin-induced SE. Prior exercise also mitigates learning and memory deficits shortly after KA-induced seizures (Gobbo and O’Mara, 2005). These data support our view that higher baseline growth factor levels in our SUP rats prior to SE may confer a neuroprotective hippocampal microenvironment that would make SUP rats more resilient to the deleterious effects of seizures. The larger growth factor pool in the SUP hippocampus may be due to several factors. Prenatal choline-supplemented rats may have enhanced cholinergic neurotransmission (Blusztajn et al., 1998) and increased respon-
siveness to cholinergic stimulation (Montoya et al., 2000), factors that are known to upregulate hippocampal BDNF and NGF expression (Lindefors et al., 1992; Knipper et al., 1994). Prenatal choline supplementation also enhances the phosphorylation of CREB (Mellott et al., 2004), and CREB-related factors regulate BDNF transcription (Tao et al., 1998).

Although we did not detect a significant increase in baseline dentate neurogenesis with prenatal choline supplementation in our young males, previous work from our laboratory has demonstrated higher baseline neurogenesis in 8-month old SUP female rats compared to age-matched controls (Glenn et al., 2007). Our inability to detect differences in baseline neurogenesis in the present study may be due to a high baseline rate of neurogenesis in the 2-month old rat (Kuhn et al., 1996; Rao et al., 2006), which may mask any effects of prenatal choline.

The mechanism by which choline supplementation to the embryo causes long-lasting changes in the adult brain and behavior is not yet fully understood, but one likely mechanism relates to choline’s role as a methyl donor (via its conversion to betaine) used for DNA methylation. Prenatal choline availability during ED12–18 determines ED 18 brain betaine concentration (Garner et al., 1995), alters both global and gene-specific DNA methylation (Niculescu et al., 2004, 2006; Kovacheva et al., 2007), and leads to changes in adult expression of genes relevant for hippocampal plasticity (Mellott et al., 2007). Enhanced hippocampal plasticity likely plays a significant role in both the memory-enhancing effects of prenatal choline supplementation (Meck and Williams, 2003; McCann et al., 2006) and its neuroprotective actions. Much like our prenatal choline-supplemented adult rats, juvenile rats that experience SE also show resistance to seizure-induced hippocampal cell loss, disinhibition, and mossy fiber sprouting (Sperber et al., 1991; Haas et al., 2001), and exhibit savings in spatial learning and memory (Stafstrom et al., 1993; Sarkisian et al., 1997) when compared to adult rats that experience SE. The retention of juvenile-like neuroplasticity and response to seizures into adulthood may contribute to prenatal choline supplementation’s preservation of cognitive function in the face of a neural insult.

Our findings may have important clinical implications. While this study examined only the neuroprotective effects of prenatal choline supplementation, some reported neuroprotective actions of choline appear to be postnatal (Thomas et al., 2007); thus, infants at risk for seizures might benefit from added choline to formula or choline supplementation to mother’s diet. These data also suggest that prenatal choline deficiency may make offspring more vulnerable to the deleterious consequences of seizures, though this remains to be determined. While estimates of normal human consumption of choline is about 425–550 mg/day, intake by pregnant woman is frequently less than half this amount (Zeisel, 2004, 2006), and menopausal status and some genetic polymorphisms of choline metabolism may even further compromise choline availability in humans (da Costa et al., 2006), suggesting that our findings may be of clinical importance to women who may be more vulnerable to choline deficiency.

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