Inborn Errors of Long-Chain Fatty Acid β-Oxidation Link Neural Stem Cell Self-Renewal to Autism

Highlights

- TMLHE controls the neural stem cell (NSC) pool in the embryonic mouse neocortex
- CPT1A and fatty acid mobilization from lipid droplets regulate the NSC pool
- TMLHE deficiencies lead to increased symmetric differentiating division of NSCs
- NSC defects under TMLHE deficiencies can be rescued by exogenous carnitine

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In Brief

The mechanisms underlying the association between inborn errors of fatty acid metabolism and developmental brain disorders such as autism remain unclear. Xie et al. find that TMLHE, a carnitine biosynthesis enzyme, and carnitine-dependent long-chain fatty acid β-oxidation control the neural stem cell pool during neocortical development by maintaining self-renewing divisions.
Inborn Errors of Long-Chain Fatty Acid β-Oxidation Link Neural Stem Cell Self-Renewal to Autism

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SUMMARY

Inborn errors of metabolism (IEMs) occur with high incidence in human populations. Especially prevalent among these are inborn deficiencies in fatty acid β-oxidation (FAO), which are clinically associated with developmental neuropsychiatric disorders, including autism. We now report that neural stem cell (NSC)-autonomous insufficiencies in the activity of TMLHE (an autism risk factor that supports long-chain FAO by catalyzing carnitine biosynthesis), of CPT1A (an enzyme required for long-chain FAO transport into mitochondria), or of fatty acid mobilization from lipid droplets reduced NSC pools in the mouse embryonic neocortex. Lineage tracing experiments demonstrated that reduced flux through the FAO pathway potentiated NSC symmetric differentiating divisions at the expense of self-renewing stem cell division modes. The collective data reveal a key role for FAO in controlling NSC-to-IPC transition in the mammalian embryonic brain and suggest NSC self-renewal as a cellular mechanism underlying the association between IEMs and autism.

INTRODUCTION

Inborn errors of metabolism (IEMs) affect approximately 1 in every 800 live births in humans (Pampols, 2010) and are commonly associated with developmental brain syndromes such as autism spectrum disorders and cognitive disabilities. Because those syndromes afflict ~1% and 2%–3% of children, respectively (van Karnebeek and Stockler, 2012; Ghaziuddin and Al-Owain, 2013), these clinical associations argue that understanding the mechanisms underlying these associations will translate into major improvements in treating developmental brain diseases. However, little is known about such underlying mechanisms.

Deficiencies in mitochondrial fatty acid β-oxidation (FAO) are particularly common IEMs. FAO pathways catabolize fatty acids of different chain lengths and represent a major metabolic engine for producing both ATP and reducing power (Houten and Wanders, 2010; Ito and Suda, 2014). The rate-limiting step for β-oxidation of long-chain fatty acids is their import from the cytoplasm into mitochondria (Figure S1A). This process requires carnitine as an acyl carrier and the action of several enzymes, including carnitine palmitoyltransferase I (CPT1), which catalyzes the rate-limiting reaction in this process. A distinct enzyme, TMLHE trimethyllysine hydroxylase, executes the first step of carnitine biosynthesis (Strijbis et al., 2010). Interestingly, recent studies have shown that TMLHE mutations occur with high frequency in human populations (Celestino-Soper et al., 2012; Nava et al., 2012).

Previous studies of mitochondrial FAO largely focused on extracerebral tissues (Houten and Wanders, 2010). However, there is evidence to suggest an association between FAO deficiencies and developmental brain disorders such as autism. Autistic children present altered circulating levels of carnitine or acyl-carnitine (i.e., phenotypes suggestive of deficiencies in long-chain FAO) (Clark-Taylor and Clark-Taylor, 2004; Filipek et al., 2004; Rossignol and Frye, 2012). Reciprocally, children identified as FAO-deficient by genetic screening commonly exhibit signature features of autism, such as developmental delay (Waisbren et al., 2013). Finally, clinical associations of TMLHE mutations with increased autism risk are now established (Celestino-Soper et al., 2012; Nava et al., 2012). However, the mechanisms underlying such associations remain unknown.

Given the growing recognition that intermediary metabolism is a central regulator of stem cell homeostasis (Ito and Suda, 2014) and that balanced neural stem cell (NSC) homeostasis is essential for proper brain development (Sun and Henner, 2014; Taverna et al., 2014), we investigated whether the association between IEMs and developmental brain disorders has an NSC component. Here we report a direct involvement of long-chain FAO in controlling the transition from NSCs to intermediate progenitor cells (IPCs) during brain development in the embryonic mouse. The collective data make a strong case for deranged NSC homeostasis as a significant mechanistic foundation for interpreting the clinical associations between IEMs of fatty acid metabolism and neuropsychiatric disorders.

RESULTS

Reduced TMLHE Expression Causes a Diminished NSC Pool in the Embryonic Neocortex

The identification of TMLHE as an autism risk gene motivated us to interrogate whether TMLHE regulates NSC...
homeostasis during development of the neocortex, the most recently evolved region of the mammalian brain and the one that houses higher brain functions. Both the TMLHE transcript and protein were detected readily in the mouse embryonic neocortex (Figures S1B and S1C). To determine whether and how TMLHE deficiencies affect NSCs, two independent small hairpin RNA (shRNA) plasmids for silencing Tmlhe expression were generated (Figures S1Da and S1E).

Adoption of loss-of-function strategies was motivated by reports that TMLHE mutations associated clinically with autism are expected to ablate, or strongly compromise, the catalytic activity of this enzyme (Celestino-Soper et al., 2012; Nava et al., 2012). TMLHE Catalytic Activity Is Required for NSC Pool Maintenance

To ensure that the NSC phenotypes observed resulted from specific knockdown of TMLHE rather than off-target effects, a functional complementation experiment was performed. Although an shRNA-resistant wild-type TMLHE rescued the shRNA-induced NSC depletion in the in utero electroporation system, co-expression of the catalytically dead TMLHE D244H did not (Figure 1).

This non-functional mutant is the product of the autism-associated TMLHE c.730G→C missense allele that alters D244, one of the three key residues of the TMLHE catalytic core (Monfregola et al., 2012; Nava et al., 2012). Mouse embryonic NSCs express the nuclear marker Pax6 and produce neurons primarily through IPCs that express the nuclear marker Tbr2 (Englund et al., 2005; Kowalczyk et al., 2009). Therefore, Pax6⁺Tbr2⁻ cells were scored as NSCs and Tbr2⁺ cells as IPCs. Nuclear markers were chosen over cytoplasmic NSC markers (e.g., Nestin) to assign cell identities because these enabled confident scoring of individual cells within the densely packed neocortical sections (Figure S1F). Quantifications based on Pax6 and Tbr2 nuclear marker labeling showed that embryonic neocortices electroporated with either of the Tmlhe shRNAs exhibited significant reductions in the fractional contribution of NSCs to total EGFP⁺ cell populations (Figure 1).

**Figure 1. TMLHE Regulates the Size of the NSC Pool in the Mouse Embryonic Neocortex**

Mouse embryos were co-electroporated with an EGFP plasmid and a control or Tmlhe shRNA plasmid at E12.5 and sacrificed at E15.5. In rescue groups, a plasmid for expressing shRNA-resistant, wild-type TMLHE (Tmlhe shRNA + TMLHE) or shRNA-resistant TMLHE D244H (Tmlhe shRNA + D244H) was co-electroporated with Tmlhe shRNA and the EGFP plasmid.

(A) Representative confocal images. Areas of the ventricular zone (Pax6⁺ layer) are shown at a higher magnification at the bottom. Scale bars, 50 μm. (B) Quantification and statistics. See also Figures S1 and S2.
These data demonstrate that TMLHE catalytic activity is required for maintaining NSC pools in the embryonic neocortex.

**TMLHE Deficiencies Preserve Key NSC Features**

Several lines of evidence demonstrate that Tmlhe shRNA challenges levied specific and biologically interesting NSC deficits. First, Tmlhe shRNA-induced reductions in the NSC pool were not due to enhanced cell death or apoptosis in the stem cell population because incidences of EGFP⁺ cells exhibiting either fragmented nuclei or activated Caspase 3 were rare in the control and Tmlhe shRNA groups (~0.1%; Figure S1G). Second, no obvious defects in neuronal migration or early neuronal differentiation were induced (Figures S1G and S1H). Third, TMLHE deficiencies left intact other NSC properties critical for self-renewal. Sequential iododeoxyuridine (IdU)/chlordodeoxyuridine (CldU) injection experiments indicated that TMLHE deficiency did not alter the length of NSC S phase or the cell cycle (Figures S2A–S2C). Tmlhe silencing also did not compromise nuclear migration to the ventricular surface of mitotic NSCs (Figures S2D and S2E), indicating that TMLHE deficiency did not impair interkinetic nuclear migration. Similarly, no apparent defects were observed in NSC division cleavage plane (Figures S2F and S2G) or in the apical attachment (Figure S2H).

A bromodeoxyuridine (BrdU) labeling regimen, followed by analysis of actively cycling cells, showed that Tmlhe shRNA challenge led to only a modest increase in cells exiting the cell cycle (Figures S2H and S2J). Because the proliferating cell pool consists of both NSCs and IPCs, and given that most IPCs generate postmitotic neurons directly (most NSCs generate neurons via proliferating IPCs), this modest elevation in cell-cycle exit was accounted for by the elevated contribution of IPCs to the proliferating population of Tmlhe shRNA-expressing cells.

**CPT1 Deficiencies Compromise NSC Pools**

To buttress the TMLHE knockdown/rescue results, we tested whether interference with an independent step in the mitochondrial long-chain FAO pathway similarly depleted NSC pools in the embryonic neocortex. To that end, the rate-limiting enzyme for long-chain FAO (CPT1) was targeted for inhibition. Of the three CPT1 isoforms, CPT1A was chosen because Cpt1a transcripts are expressed in embryonic neocortex, particularly in the ventricular zone (Figure S3A; Genepaint ID: MH509). By contrast, CPT1C does not regulate mitochondrial FAO as a carnitine palmitoyltransferase (Wolfgang and Lane, 2011), and Cpt1b mutations in mice ablated for Cpt1a suffer an early uncharacterized embryonic lethality (Nyman et al., 2005), a Cpt1a-directed shRNA plasmid (Figure S3B) was co-electroporated with an EGFP plasmid into the embryonic neocortex at E12.5, and embryos were analyzed at E15.5. Indeed, Cpt1a shRNA challenge resulted in a significant decrease in the fractional representation of NSCs in EGFP⁺ cell populations, and this NSC depletion was reversed upon co-expression of an shRNA-resistant Cpt1a cDNA (Figures 2A and 2B). Moreover, as observed for TMLHE deficiencies, CPT1A silencing neither enhanced apoptosis nor did it overtly compromise neuronal migration or differentiation (Figures S3C and S3D).

CPT1A involvement in regulating NSC homeostasis was similarly evident upon NSC intoxication with etomoxir, a small-molecule inhibitor of the enzyme. Because of difficulties in controlling etomoxir exposure in vivo, E12.5 forebrain hemispheres were cultured in media with or without 100 μM etomoxir, a concentration at which etomoxir inhibited FAO activity in neocortical cultures dominated by NSCs (Figures 2C and 2D). The associated phenotypic consequences were thinning of the ventricular zone (Pax6⁺ layer) with accompanying increases in Pax6⁺/Tbr2⁺ IPC populations (Figures 2E and 2F), results congruent with the phenotypes evoked by Tmlhe and Cpt1a knockdown in the intact neocortex. Taken together, the complementary gene silencing and pharmacological intervention data demonstrated that reduced metabolic flux through the long-chain FAO pathway upsets NSC homeostasis.

**Reduced Fatty Acid Mobilization from Lipid Droplets Diminishes the NSC Pool**

Lipid droplets (LDs) are primary fatty acid depots for mitochondrial FAO (Gross and Silver, 2014), and these structures are found in embryonic NSCs (Bush et al., 1992; Saito et al., 2009). To test whether interfering with fatty acid mobilization from LDs recapitulates the NSC derangements induced by direct compromise of mitochondrial FAO, a dominant-negative interference assay was developed. The experiment was grounded on demonstrations that PLIN1 inhibits LD lipolysis by prohibiting lipase access to LD surfaces. Phosphorylation of PLIN1 by protein kinase A (PKA) at six sites releases this lipolytic block, and PLIN1 mutants that cannot be so phosphor ylated are potent dominant-negative inhibitors of LD lipolysis (Figure S3E; Gross and Silver, 2014; Sztalryd and Kimmel, 2014). Co-electroporation of an EGFP plasmid with a vector driving expression of either a mutant PLIN1 with all six PKA phosphorylation sites mutated to alanine (PLIN1-6A) or a PLIN1 with five of the six PKA sites mutated to alanine (PLIN1-5A) evoked significant reductions in NSC pool size (Figures 2G and 2H). Those effects were again evident in the absence of enhanced cell death/apoptosis or overt defects in neuronal migration or neuronal differentiation (Figures S3F and S3G). The data show that withholding LD-derived fatty acids from an otherwise undulated mitochondrial FAO pathway elicits NSC homeostatic defects similar to those evoked by direct interference with activities of FAO enzymes.

**FAO Regulates the Mitochondrial Redox State**

FAO is a major supply for both ATP and reducing power (Houten and Wanders, 2010; Ito and Suda, 2014). To address how diminutions in FAO metabolic flux affect these two NSC parameters, TMLHE and CPT1A expression were knocked down individually or in combination (mtAT1.03), or a mitochondrially targeted AT1.03 fluorescence resonance energy transfer (FRET)-based ATP biosensor (Ismura et al., 2009), or a mitochondrially targeted AT1.03 (mtAT1.03) was co-electroporated with appropriate shRNA.

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Figure 2. Targeting CPT1 or Fatty Acid Mobilization Diminished the NSC Pool

(A and B) Silencing Cpt1a diminished the NSC pool. Mouse embryos were electroporated at E12.5 and sacrificed at E15.5. In the rescue group (Cpt1a shRNA + CPT1A), a plasmid for expressing shRNA-resistant mouse Cpt1a cDNA was co-electroporated with Cpt1a shRNA and the EGFP plasmid. (A) Representative confocal images. Areas of the ventricular zone (Pax6+ layer) are shown at a higher magnification at the bottom. Scale bars, 50 μm. (B) Quantifications.

(C–F) Etomoxir induces increased differentiation of NSCs.

(C) Neocortical cells used for the FAO assay. Dissociated neocortical cells of E11.5 mouse embryos were cultured overnight and then used in a [14C]-oleate oxidation assay to assess FAO activity. Most cultured cells express the NSC marker Nestin. DAPI labels the nuclei of all cells.

(D) The [14C]-Oleate oxidation assay confirms that etomoxir inhibits FAO activity in cultured NSCs.

(E and F) Etomoxir potentiated NSC differentiation in organotypic culture of forebrain hemispheres. (E) Representative images from three independent experiments showing reduced thickness of the ventricular zone (Pax6+ layer) in the etomoxir group and an increased fraction of Pax6+ cells that co-express Tbr2. The pial surface is outlined by dashed lines. Scale bar, 20 μm.

(*p<0.001 compared to control, one-way ANOVA)

(legend continued on next page)
vectors into the embryonic neocortex. Transfected cells exhibiting signature NSC morphologies (bipolar with an apical process contacting the ventricular surface) and locations (soma close to the ventricular surface) were analyzed by ratiometric YFP/CFP FRET imaging. No significant differences in FRET/CFP ratios were discerned between the control and Tmlhe shRNA or Cpt1a shRNA groups (Figures S3H–S3K), indicating that NSCs maintained substantially normal bulk and mitochondrial ATP levels in the face of TMLHE or CPT1A deficiency.

To monitor the mitochondrial redox status in transfected NSCs, a MitoTimer expression vector was co-electroporated with appropriate shRNA plasmids into an E12.5 embryonic mouse neocortex. MitoTimer is a mitochondria-targeted GFP whose emission wavelengths are sensitive to the redox state of the environment (Laker et al., 2014). Analyses of MitoTimer fluorescence properties were again restricted to transfected cells with morphologies and tissue distributions diagnostic of NSCs. In those cells, MitoTimer localized correctly to mitochondria, and the punctate MitoTimer localization profiles were similar in all experimental groups (Figure S3L). Therefore, neither Tmlhe nor Cpt1a deficiencies induced significant morphological derangements in the mitochondrial network. However, ratiometric analyses of MitoTimer green/red fluorescence emission spectra showed significant red shifts in NSCs expressing Tmlhe shRNA or Cpt1a shRNA relative to controls (Figures S3M and S3N), indicating that diminished flux through the FAO pathway resulted in oxidized NSC mitochondrial environments.

**TMLHE Regulates NSC Daughter Cell Fate**

Embryonic mouse NSCs undergo three exquisitely balanced modes of cell division: a symmetric self-expanding mode that produces two NSC daughters, an asymmetric division that produces one NSC and one differentiated daughter, and a symmetrical increase in the fractional representation of IPC/IPC cell pairs and that of NSC/NSC pairs. Because TMLHE generates a biodeliverable product (carnitine), the effect of TMLHE deficiency on NSC daughter fate was recapitulated in a different assay where Attractene, a nonliposomal transfection reagent, was used to transfet a mouse embryonic neocortex in vivo. When an EGFP plasmid was co-injected with Attractene into lateral ventricles of E13.5 mouse embryos, EGFP expression was already apparent 10 hr after injection and became robust by 15 hr. In samples with low transfection efficiencies, EGFP+ cells were scattered around the lateral ventricle as isolated single cells or cell pairs that were interpreted to be derived from the division of transfected single cells. Immunostaining analyses 15 hr after transfection revealed that all EGFP+ cells (>300 cells examined) were either NSCs or IPCs. Moreover, most of the cell pairs (>90%) were either NSC/IPC or NSC/NSC pairs, whereas the remainder scored as IPC/IPC pairs (Figures S4D and S4E). Because IPCs divide to produce IPCs or postmitotic neurons (but not NSCs), whereas NSCs divide to produce both NSCs and IPCs, these data indicated that the Attractene regimen preferentially transfected NSCs.

To determine the effect of TMLHE deficiency on daughter cell fate during NSC division, a mixture of EGFP plasmid and control or Tmlhe shRNA was co-injected with Attractene into the lateral ventricle of E13.5 mouse embryos, and the cell fates of EGFP+ cell pairs (but not EGFP+ single cells or clones of more than two EGFP+ cells) were analyzed after 24 hr when Tmlhe shRNA-induced NSC defects were already detectable (Figures S4F and S4G). Tmlhe shRNA challenge led to a dramatic increase in the fractional representation of IPC/IPC cell pairs, and this increase was again corrected by co-expression of silencing-resistant, wild-type TMLHE but not silencing-resistant TMLHED244H (Figures 3C and 3D). Both the fractional representation of NSC/IPC pairs and that of NSC/NSC pairs decreased in the Tmlhe shRNA group (Figures S4H and S4I). Although the phenotypes in the Attractene experiments were somewhat milder than those in the two-step electroporation experiments, this difference was expected because it is the likely consequence of less efficient depletion of endogenous TMLHE protein in the Attractene regimen (the interval between shRNA transfection and cell fate analysis was shorter). Therefore, the two-step electroporation and the Attractene transfection data converge to provide compelling evidence that TMLHE deficiencies promote symmetric differentiating NSC divisions that produce two IPCs at the expense of the symmetric NSC/NSC and asymmetric NSC/IPC cell division modes that support stem cell self-renewal.

**Exogenous Carnitine Rescues Tmlhe shRNA-Induced NSC Defects**

Because TMLHE generates a biodeliverable product (carnitine), it was of interest to assess whether depletion of NSCs from the embryonic neocortex in the face of TMLHE deficiencies could...
be remediated by providing carnitine to the tissue. Control or Tmlhe shRNAs were co-electroporated with an EGFP plasmid into a mouse embryonic neocortex at E12.5, and electroporated embryos were allowed to develop in utero for a further 24 hr before being sacrificed. Forebrain hemispheres were then dissected and incubated in physiological glucose media with or without carnitine supplementation for 18 hr (Figure 4A). Under those conditions, Tmlhe shRNA induced a 2-fold decrease in the fractional representation of NSCs in EGFP+ cell populations, and this reduction was rescued by inclusion of carnitine in the incubation medium (Figures 4B and 4C). Therefore, derangements in NSC pools caused by TMLHE deficiencies are carnitine-remediable. Although the NSC pool was not recovered completely, we noted that Tmlhe shRNA had already reduced the NSC pool size by 24 hr after electroporation (i.e., prior to carnitine supplementation; Figures S4F and S4G). Further analyses revealed that, although carnitine supplementation significantly enlarged the NSC pool in the Tmlhe shRNA group, it did not expand the NSC pool in the control group (Figure 4C). This result indicated that exogenous carnitine modulates NSC pools only when endogenous carnitine biosynthesis is compromised.

**DISCUSSION**

Here, we demonstrate a critical requirement for long-chain FAO in maintaining NSC homeostasis in the mammalian embryonic neocortex. The key evidence includes gene silencing and chemical inhibition approaches that interfere with distinct enzymatic reactions in the mitochondrial FAO pathway and dominant-negative approaches that restrict substrate entry into a biochemically uncompromised mitochondrial FAO pathway. These results demonstrate that reduced flux through this catabolic pathway directly disturbs NSC homeostasis by inappropriately enhancing the transition of NSCs to IPC lineages. Our findings offer a mechanistic framework for interpreting the associations between FAO deficiencies and developmental brain disorders and provide a biochemical basis to explain why TMLHE scores as an autism-linked gene.

Figure 3. TMLHE Inhibits Symmetric NSC Division that Produces Two IPCs
(A and B) Analysis of NSC division by two-step electroporation assay.
(A) Representative images showing vicinal EGFP+mCherry+ cell pairs of NSC/NSC, NSC/IPC, and IPC/IPC. White and yellow arrows indicate NSCs (Pax6+/Tbr2+) and IPCs (Tbr2+), respectively. Scale bars, 20 μm.
(B) Quantification and statistics.
(C and D) Analysis of NSC division using Attractene-mediated transfection.
(C) Representative images showing vicinal EGFP+ cell pairs of NSC/NSC, NSC/IPC, and IPC/IPC. White and yellow arrows indicate NSCs and IPCs, respectively. Scale bars, 20 μm.
(D) Quantification and statistics.
See also Figure S4.
Lineage tracing experiments demonstrate that an active FAO pathway supports NSC pool maintenance in the embryonic neocortex by dampening the frequency of symmetric differentiating cell divisions that give rise to lineage-restricted cells (IPCs) at the expense of NSC self-renewal. The relevance of this activity to autism is underscored by growing evidence showing that established autism-linked genes or pathways are key regulators of the NSC-to-IPC transition (Mutch et al., 2010; Saffary and Xie, 2011; Bian et al., 2013; Lv et al., 2013; Krumm et al., 2014; O’Roak et al., 2014). Our data further show that flux through the mitochondrial FAO pathway controls a metabolic trigger that regulates daughter cell fate during NSC division. The nature of this trigger remains unclear. Although in vitro data suggest that FAO maintains energy levels in adult subventricular zone (SVZ) neural stem and/or progenitor cells (Stoll et al., 2015), our in vivo experiments measured no significant derangements in bulk or mitochondrial ATP pools in FAO-deficient NSCs. However, because ATP levels are highly buffered in cells, we cannot discount the possibility that mitochondrial FAO insufficiencies are accompanied by compensatory metabolic adaptations that marshal other metabolic resources in support of bulk intracellular ATP. Such compensatory metabolic adaptations might be accompanied by wider pleiotropic effects that could influence daughter cell fate during NSC division.

With regard to redox status, enhanced oxidative environments were registered in NSC mitochondria with diminished FAO activity, suggesting that enhanced oxidative microenvironments impair NSC self-renewal in the mouse embryonic neocortex. Although reactive oxygen species are reported to regulate NSC self-renewal in the adult mouse brain (Le Belle et al., 2011), in that case, oxidative stress promoted NSC self-renewal rather than diminishing that capacity; i.e., the conclusion we arrive at here. Perhaps NSC self-renewal is regulated differentially as a function of magnitude of the oxidative stimulus. Alternatively, embryonic NSCs may fundamentally differ from adult brain NSCs in their metabolic wiring.

Why has a direct role for FAO in NSCs and brain development been largely ignored? First, classical studies of brain energy metabolism focused on neurons, glial cells, or the brain at the organ level (Bélanger et al., 2011). Because NSCs constitute only a small fraction of brain cells, NSC metabolic activity does not contribute significantly to the bulk-averaging measurements that define many analyses. Second, characterization of NSC defects in animal models of FAO deficiencies is confounded by broader physiological consequences of these metabolic defects. For instance, mice ablated for Cpt1a suffer an early embryonic lethality presumed to reflect systemic metabolic failures (Nyman et al., 2005). Third, previous studies of NSC metabolism relied on dissociated NSC cultures. Those strategies, although convenient, ignore niche parameters that regulate NSC self-renewal. By contrast, the experimental paradigms used here imposed TMLHE and CPT1A insufficiencies acutely, in situ, and in a manner restricted to defined subsets of cells within the embryonic neocortex. Such manipulation of NSCs, in an authentic physiological context, exposed the critical role played by long-chain FAO in NSC self-renewal. Although our analyses were restricted to mitochondrial β-oxidation of long-chain fatty acids, inherited deficiencies in medium- and short-chain FAO pathways are also recognized to predispose affected patients to developmental challenges (Waisbren et al., 2013). Our findings now raise the possibility that those diseases also reflect derangements in NSC homeostatic control.
Finally, linkage of NSC homeostatic mechanisms with IEMs of developmental brain disorders has clinical implications. TMLHE deficiencies represent the most frequent IEMs (Pampols, 2010; Celestino-Soper et al., 2012; Nava et al., 2012) and are associated with an increased autism risk. The carnitine rescue experiment suggests actionable prophylactic measures for minimizing developmental brain deficits associated with TMLHE deficiencies. We forecast that elevation of the fetal carnitine supply will prove to be an effective strategy for alleviating developmental brain deficits associated with inborn TMLHE mutations.

The commercial availability of carnitine as a nutritional supplement notwithstanding, this compound is not on the list of dietary supplements recommended by the Food and Drug Administration (FDA) for pregnant women. We suggest that genetic screening of prospective parents for TMLHE mutations, coupled with inclusion of carnitine as a dietary supplement upon initial diagnosis of pregnancy, promises mental health benefits for newborns otherwise at significant risk for developmental brain disorders.

EXPERIMENTAL PROCEDURES

Detailed experimental procedures are provided in the Supplemental Experimental Procedures.

Animals and In Utero Electroporation

Swiss Webster female mice were purchased from Taconic, Charles River Laboratories, or Harlan Laboratories. Mice were handled in accordance with NIH and institutional guidelines regarding the care and use of animals. In utero electroporation was performed as described previously (Saffary and Xie, 2011).

Tissue Preparation and Immunostaining

Electroporated mouse embryos were sacrificed by decapitation. Forebrain hemispheres were fixed in 2% paraformaldehyde (prepared in PBS) for 20–30 min. Cryosections (30–40 μm) were prepared from the hemispheres and used for immunostaining.

Statistical Analysis

Statistical analysis was performed using GraphPad Prism software.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures and four figures and can be found with this article online at http://dx.doi.org/10.1016/j.celrep.2016.01.004.

AUTHOR CONTRIBUTIONS

Z.X. conceptualized, designed, and performed the research; analyzed the data; and wrote the manuscript. V.A.B. designed the research, analyzed the data, and edited the manuscript. A.J. and J.T.D. designed, performed, and analyzed FAO metabolic flux measurements. S.K.H. contributed to in utero electroporation assays.

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