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Cell-autonomous regulation of astrocyte activation by the circadian clock protein BMAL1

Brian V. Lananna Washington University School of Medicine in St. Louis

Collin J. Nadarajah Washington University School of Medicine in St. Louis

Michelle R. Cedeno Washington University School of Medicine in St. Louis

David D. Xiong Washington University School of Medicine in St. Louis

Julie Dimitry Washington University School of Medicine in St. Louis

See next page for additional authors

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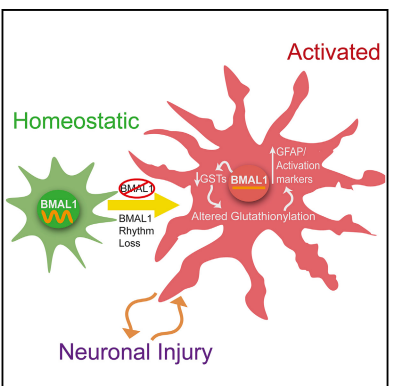
Authors

Brian V. Lananna, Collin J. Nadarajah, Michelle R. Cedeno, David D. Xiong, Julie Dimitry, Chak Foon Tso, Celia A. McKee, Percy Griffin, Patrick W. Sheehan, Jeffery A. Haspel, Erik S. Musiek, and et al

Cell Reports

Cell-Autonomous Regulation of Astrocyte Activation by the Circadian Clock Protein BMAL1

Graphical Abstract



Highlights

- Circadian disruption promotes astrocyte activation
- Astrocyte-specific deletion of the circadian clock gene BMAL1 induces activation
- BMAL1 regulates astrocyte activation by altering glutathione-S-transferase signaling
- Loss of astrocyte BMAL1 enhances neuronal cell death in a co-culture system

Authors

Brian V. Lananna, Collin J. Nadarajah, Mariko Izumo, ..., Joseph S. Takahashi, Ilia N. Karatsoreos, Erik S. Musiek

Correspondence

musieke@wustl.edu

In Brief

Lananna et al. show that the circadian clock protein BMAL1 regulates astrocyte activation via a cell-autonomous mechanism involving diminished glutathione-S-transferase signaling. This finding elucidates a function of the core circadian clock in astrocytes and reveals BMAL1 as a modulator of astrogliosis.



Cell-Autonomous Regulation of Astrocyte Activation by the Circadian Clock Protein BMAL1

Brian V. Lananna,¹ Collin J. Nadarajah,¹ Mariko Izumo,² Michelle R. Cedeño,¹ David D. Xiong,¹ Julie Dimitry,¹ Chak Foon Tso,³ Celia A. McKee,¹ Percy Griffin,¹ Patrick W. Sheehan,¹ Jeffery A. Haspel,⁴ Ben A. Barres,⁵ Shane A. Liddelow,^{6,7} Joseph S. Takahashi,^{2,8} Ilia N. Karatsoreos,⁹ and Erik S. Musiek^{1,10,*}

¹Department of Neurology and Hope Center for Neurological Disorders, Washington University School of Medicine, St. Louis, MO, USA ²Department of Neuroscience, University of Texas Southwestern Medical Center, Dallas, TX, USA

³Department of Biology, Washington University, St. Louis, MO, USA

⁴Department of Medicine, Washington University School of Medicine, St. Louis, MO, USA

⁵Department of Neurobiology, Stanford University School of Medicine, Stanford, CA, USA

⁶Neuroscience Institute, Department of Neuroscience and Physiology, NYU Langone Medical Center, New York, NY, USA

⁷Department of Pharmacology and Therapeutics, University of Melbourne, Melbourne, VIC, Australia

⁸Howard Hughes Medical Institute, University of Texas Southwestern Medical Center, Dallas, TX, USA

⁹Integrative Physiology and Neuroscience, College of Veterinary Medicine, Washington State University, Pullman, WA, USA ¹⁰Lead Contact

*Correspondence: musieke@wustl.edu

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SUMMARY

Circadian clock dysfunction is a common symptom of aging and neurodegenerative diseases, though its impact on brain health is poorly understood. Astrocyte activation occurs in response to diverse insults and plays a critical role in brain health and disease. We report that the core circadian clock protein BMAL1 regulates astrogliosis in a synergistic manner via a cell-autonomous mechanism and a lesser noncell-autonomous signal from neurons. Astrocytespecific Bmal1 deletion induces astrocyte activation and inflammatory gene expression in vitro and in vivo, mediated in part by suppression of glutathione-Stransferase signaling. Functionally, loss of *Bmal1* in astrocytes promotes neuronal death in vitro. Our results demonstrate that the core clock protein BMAL1 regulates astrocyte activation and function in vivo, elucidating a mechanism by which the circadian clock could influence many aspects of brain function and neurological disease.

INTRODUCTION

Changes in behavioral circadian rhythms are common in many neurodegenerative diseases (Breen et al., 2014; Hatfield et al., 2004; Musiek and Holtzman, 2016) and can occur early in disease progression (Musiek et al., 2018). Circadian rhythms are generated by the suprachiasmatic nucleus of the hypothalamus (SCN) (Mohawk et al., 2012), which synchronizes cellular clocks throughout the body to the light-dark (L:D) cycle. The core molecular clock consists of a transcriptional-translational feedback loop with a positive transcriptional limb, consisting of heterodimers of the basic-helix-loop-helix (bHLH) transcription factor BMAL1 (also known as ARNTL [aryl hydrocarbon receptor nuclear translocator-like protein 1]) with either CLOCK (circadian locomotor output cycles kaput) or NPAS2 (neuronal PAS domain protein 2). The negative limb includes Period (PER), cryptochrome (CRY), and REV-ERB genes, which are transcriptional targets of BMAL1 and provide negative feedback inhibition of BMAL1 function (Mohawk et al., 2012). The core clock regulates transcription of thousands of genes in a highly tissue-specific manner, regulating cellular functions including metabolism, inflammation, and redox homeostasis (Bass and Takahashi, 2010). In addition to disrupted activity and sleep rhythms, dysregulated circadian gene expression patterns are also observed in neurodegenerative disease models and patients (Song et al., 2015; Breen et al., 2014; Cronin et al., 2017). However, the impact of clock dysfunction on neurological diseases of the brain remains poorly understood.

Astrocytes play a critical role in brain health and neurodegenerative disease, as dysfunctional astrocytes can drive neurodegeneration (Lian et al., 2015; Macauley et al., 2011; Yamanaka et al., 2008). Astrocyte activation is a ubiquitous response to brain injury, from neurodegeneration to trauma, which has historically been characterized by increased expression of the cytoskeletal protein glial fibrillary acidic protein (GFAP) (Sofroniew, 2014). Recent work has begun to elucidate the diversity of astrocyte activation phenotypes beyond GFAP, as astrocytes activated by different stimuli express unique transcriptional profiles that can be associated with divergent phenotypes, ranging from neurotoxic to neurotrophic (Liddelow et al., 2017; Zamanian et al., 2012). However, a further understanding of astrocyte activation mechanisms is needed in order to effectively target astrocyte function therapeutically.

Astrocyte activation and circadian clock dysfunction are two pervasive, often coexistent features of neurological diseases, though their interaction is unknown. We previously observed that deletion of the core clock gene *Bmal1*, which abrogates all circadian clock function, caused severe, spontaneous astrogliosis throughout the mouse brain, which was accompanied by increased oxidative stress, synaptic damage, and inflammation (Musiek et al., 2013). However, the cellular and molecular mechanisms linking BMAL1 to astrocyte activation and function

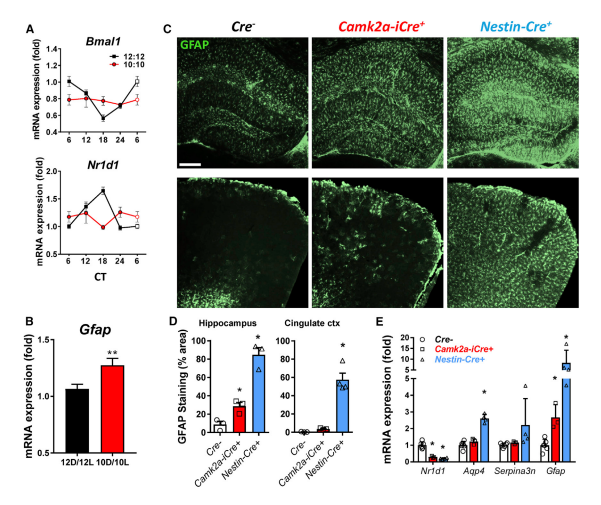


Figure 1. Light-Induced Circadian Disruption or Neuron-Specific *Bmal1* Deletion Partially Recapitulates the Astrocyte Activation Phenotype Observed following Brain-Specific *Bmal1* Deletion

(A) Cortical qPCR from 6-week-old WT mice housed in 12 hr:12 hr or 10 hr:10 hr light-dark cycles for 6 weeks, harvested every 6 hr from CT6 to CT24, reveals blunting of circadian gene oscillations in 10 hr:10 hr mice.

(B) Total Gfap mRNA levels for mice from (A). **p < 0.01 by 2-tailed t test.

(C) Representative images showing GFAP immunostaining of hippocampus (top) and cingulate cortex (bottom) of neuron-specific *Bmal1* KO mice (*Cam2a-iCre+;Bmal1^{ff}*) and brain-specific (neurons+astrocytes) *Bmal1* KO mice (*Nestin-Cre+;Bmal1^{ff}*) at 4 months. Scale bar, 100 μm.

(D) Quantification of GFAP immunoreactivity in two brain regions of Cre-, *Cam2a-iCre+;Bma11^{ff}*, and *Nestin-Cre+;Bma11^{ff}* mice. n = 3–4 mice per genotype. Each data point represents 1 mouse. *p < 0.05 by 1-way ANOVA.

(E) Quantification of astrocyte activation transcripts by qPCR in the cortex of Cre-;*Bmal1^{flf}*, *Cam2a-iCre+;Bmal1^{flf}*, and *Nestin-Cre+;Bmal1^{flf}* mice. The clock-controlled gene *Nr1d1* is shown to illustrate loss of BMAL1-mediated transcription. *p < 0.05 by 2-way ANOVA with Dunnett's post-test for multiple comparisons. All data represent mean ± SEM.

remain uncertain. Thus, we sought to characterize the astrocyte activation induced by circadian clock disruption and evaluate whether the clock protein BMAL1 might regulate astrogliosis in a cell-autonomous manner.

RESULTS

We first asked if nongenetic disruption of circadian rhythms could influence *Gfap* levels in the brain. Exposure of wild-type (WT) C57BL/6 mice to a 10 hr:10 hr L:D cycle induces circadian desynchrony and eventually arrhythmicity, as well as loss of detectable circadian transcript oscillations in the cerebral cortex (Figures 1A and S1A). After 6 weeks of circadian disruption, *Gfap*

transcript levels were increased throughout the circadian cycle (Figure S1B), with an average increase of 19.7% (Figure 1B). Thus, light-induced behavioral circadian rhythm disruption promotes astrocyte *Gfap* expression.

We previously observed that abrogation of circadian function by deletion of *Bmal1* in mice induced striking increases in GFAP+ astrocytes throughout the brain (Musiek et al., 2013; Yang et al., 2016). We asked if the astrocyte activation observed following *Bmal1* deletion was simply a response to neuronal injury or a cell-autonomous process regulated by the core clock in astrocytes. To address this, we examined the brains of neuron-specific *Bmal1* knockout (KO) mice (*Camk2a-iCre;Bmal1^{fif}*). These mice express Cre recombinase in a pan-neuronal manner under the control of a BAC-CaMKIIa promoter, yielding widespread neuronal deletion of Bmal1 (Figure S1C) and loss of behavioral circadian rhythms (Izumo et al., 2014). At 4 months, Camk2a*iCre+;Bmal1^{f/f}* mice exhibited a modest increase in astrocyte activation, as assessed by GFAP immunostaining (Figures 1C and 1D) and qPCR (Figure 1E), in the cortex and hippocampus. However, similarly aged Nestin-Cre+;Bmal1^{f/f} (referred to as NBmal1 KO) mice, in which Bmal1 is deleted in neurons and astrocytes, have much more prominent astrogliosis throughout the brain and higher expression of the activation-related transcripts Agp4 and Serpina3n (Figures 1C-1E). While NBmal1 KO mice exhibit large increases in GFAP+ astrocytes in the cortex, this did not appear to be due to increased astrocyte division, as NBmal1 KO mice had no significant increase in dividing Ki67+, GFAP+ double-positive cells (Figure S1D) or immunoreactivity for the pan-astrocyte marker S100B (Figure S1E). This finding suggests that while a small component of astrogliosis observed following Bmal1 deletion may be a response to neuronal injury, BMAL1 within astrocytes appears to play a critical cell-autonomous role in astrocyte activation.

To assess the potential cell-autonomous effects of Bmal1 on astrocyte activation, we generated primary astrocyte-enriched cultures from Bmal1 KO mice and WT littermates and examined activation markers in vitro. Bmal1 KO astrocytes were viable and appeared to have normal morphology but had significantly elevated expression of Gfap and Agp4 mRNA and GFAP protein (Figures 2A and 2B), suggesting spontaneous activation in the absence of neuronal influence. In order to exclude a developmental influence of BMAL1, we used small interfering RNA (siRNA) to knock down Bmal1 expression in WT primary astrocvte cultures. This method resulted in a 95% decrease in BMAL1 protein (Figures 2C and 2D) and a 66% decrease in expression of the BMAL1 target Nr1d1 (which encodes REV-ERBa) (Figure S2A). Bmal1 knockdown (KD) in astrocytes induced a highly consistent (across 14 independent experiments) 36% increase in Gfap mRNA expression (Figure S2A) as well as a 68% increase in GFAP protein (Figures 2C and 2D). We also saw significant increases in the astrocyte activation marker S100a6 (1.7-fold), the cytokine II6 (2.4-fold), and II33 (1.6-fold) (Figure S2A), an astrocytic cytokine implicated in astrocyte-microglia signaling (Wicher et al., 2017). Bmal1 KD in primary astrocytes from Per2-luciferase reporter mice caused a loss of circadian Per2 rhythms (Figure 2E). Thus, disruption of circadian function via KD of Bmal1 in WT astrocytes induces a cell-autonomous reactive phenotype in vitro.

We next sought to confirm these findings by specifically deleting *Bmal1* in astrocytes *in vivo*. We found that intracerebroventricular injection of an AAV8 viral vector expressing EGFP under a *Gfap* promoter (AAV8-GFAP-EGFP) on postnatal day 2 led to wide-spread EGFP expression in resting astrocytes throughout the cerebral cortex, with almost no neuronal labeling (Figures 2F, 2G, and S2B). Thus, we injected P2 *Bmal1*^{fif} mice either with AAV8-GFAP-GFP or with an identical virus expressing a Cre-GFP fusion protein (AAV8-GFAP-Cre^{GFP}). At 5 months, we observed *Bmal1* deletion and increased GFAP expression specifically in astrocytes expressing AAV8-GFAP-Cre^{GFP} (Figure 2F). The GFAP-Cre^{GFP} vector exhibits very weak nuclei-restricted fluorescence, making it more difficult to see than that of the AAV8-GFAP-EGFP vector.

Mice treated with the EGFP control vector had widespread astrocyte labeling throughout the cingulate and retrosplenial cortices, intact BMAL1 expression, and very little astrocyte activation as defined by GFAP immunoreactivity (Figures 2F and 2G). Mice injected with the Cre^{GFP} vector showed astrocytic nuclear GFP expression and 3.4- and 3.1-fold increases in GFAP immunoreactivity in the cingulate and retrosplenial cortices, respectively (Figure 2G). Finally, we generated tamoxifen-inducible astrocytespecific Bmal1 KO mice using an Aldh111-CreERT2 driver line (Aldh111-Cre^{ERT2}; Bmal1^{f/f} mice) (Srinivasan et al., 2016). We treated Cre+ and Cre- mice with tamoxifen at 1 month and examined astrocyte activation at 3 months. We observed specific deletion of Bmal1 in astrocyte nuclei of the hippocampus and cortex (Figures S2D and S2E). Aldh111-Cre+ mice had significant increases in astrogliosis, as assessed by GFAP immunostaining in the hippocampus (Figure 2H), and a nearly significant (p = 0.055) increase in the retrosplenial cortex (Figure S2C). While it is difficult to directly compare tamoxifen-inducible and constitutive Cre lines, Aldh111-Cre+ mice had similar levels of astrogliosis to tamoxifen inducible global Bmal1 KO mice (CAG-CreERT2+;Bmal1^{f/f}) (Yang et al., 2016) also harvested 2 months after TAM treatment (Figures 2H and S2C). Astrogliosis was not caused by Cre expression, as Cre+;Bmal1^{flox/wt} mice had no phenotype (Figures 2H and S2C). Ald111-Cre+ mice also had spontaneous and significant increases in a number of additional astrocyte activation transcripts including S100a6 (2.5-fold), C4b (2.6-fold), Cxcl5 (7.5-fold), and Mmp14 (3.1-fold) (Figure S2F). In total, these experiments demonstrate that Bmal1 deletion in astrocytes can induce activation in a cell-autonomous manner.

While activated astrocytes generally exhibit morphologic changes and increased GFAP expression, the transcriptional profile of activated astrocytes is heterogeneous and can define their function (Liddelow et al., 2017; Zamanian et al., 2012). In order to examine the transcriptional profile in the brain following Bmal1 deletion, we performed microarray analysis of cortical tissue from 11-month-old NBmal1 KO mice, Cre- controls, and Per1^{Brdm}/Per2^{Brdm} (Per1/2^{mut}) mice placed in constant dark conditions for 36 hr and then harvested at clock time (CT) 6:00 and 18:00. Per1/2^{mut} mice were included because they have a dysfunctional circadian clock but no astrogliosis phenotype as well as tonically increased levels of BMAL1 transcriptional targets (Figures 3A and S3A). To understand astrocyte transcriptional changes in NBmal1 KO brain, we examined the overlap between the 200 most upregulated transcripts in NBmal1 KO with the 200 most astrocyte-enriched transcripts according to the Barres lab RNA-sequencing (RNA-seq) database (Zhang et al., 2014) (Figure 3A). Thirty genes overlapped, including Gfap, Aqp4 (encoding the astrocytic water channel aquaporin-4), and Megf10, a gene involved in astrocyte-mediated synaptic elimination (Chung et al., 2013). C4b, a complement cascade component also implicated in synapse elimination and a marker of astrocyte activation and aging (Boisvert et al., 2018; Clarke et al., 2018), was strongly upregulated (4-fold) in NBmal1 KO brain. Several other diseaseassociated astrocytic genes were increased in NBmal1 KO brain, including inflammatory mediators such as Cxcl5 and Ccr7 and the matrix metalloproteinases Mmp14 and Mmp2. Nearly all of these transcripts were either unchanged or decreased in Per1/2^{mut}

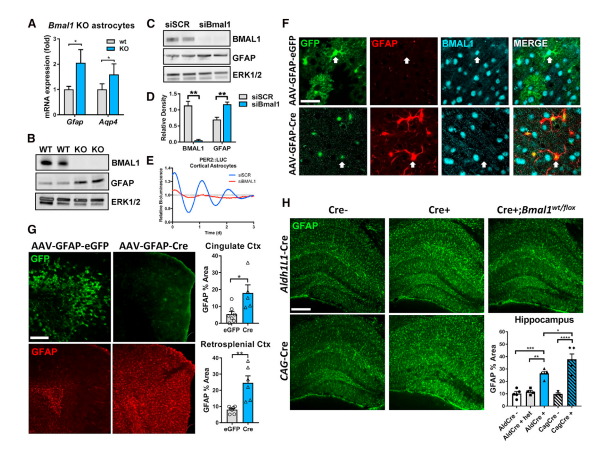


Figure 2. Bmal1 Deficiency Induces Cell-Autonomous Astrocyte Activation In Vitro and In Vivo

(A) qPCR of *Gfap* and *Aqp4* mRNA in primary astrocyte-enriched cultures from WT or global *Bmal1* KO mice. n = 8–12 samples from 3 independent experiments.
(B) Western blot showing increased GFAP protein levels in cultured astrocyte cell lysates from *Bmal1* KO mice. p42/44 ERK protein is shown as a loading control.
(C) Western blot showing knockdown of BMAL1 and increased GFAP protein in WT astrocyte cultures 7 days after treatment with non-targeting siRNA (siSCR) or siRNA targeting *Bmal1* (siBmal1). p42/44 ERK protein is shown as a loading control.

(D) Densitometric quantification of blots in (C). n = 4 samples/group.

(E) Representative trace depicting detrended circadian oscillations in luminescence seen in primary astrocyte cultures from *Per2-Luc* reporter mice treated with siRNA (siSCR, blue; or siBmal1, red). Similar results were obtained in 3 separate cultures.

(F) Representative confocal images from cerebral cortex of *Bmal1^{fff}* mice 5 months after intracerebroventricular (i.c.v.) injection at age P2 (postnatal day 2) with viral vectors inducing astrocyte-specific expression of EGFP (AAV8-GFAP-EGFP) or Cre (AAV8-GFAP-Cre^{GFP}). AAV8-GFAP-GFP-infected cells show whole-cell GFP expression and persistent colocalized nuclear BMAL1 staining, while AAV8-GFAP-Cre infected cells show nuclear GFP expression, increased GFAP expression, and loss of nuclear BMAL1. Scale bar, 30 μm.

(G) Representative images (left) and quantification (right) illustrating increased percent area covered by GFAP+ astrocytes in cortex in mice from (F) injected with AAV8-GFAP-Cre, as compared to AAV8-GFAP-GFP-injected controls. Note that AAV8-GFAP-Cre virus encodes a Cre^{GFP} fusion protein that is nuclear localized and more difficult to see (F). Scale bar, 100 μm. Each data point represents 1 mouse.

(H) Representative images and quantification of percent area of GFAP staining from hippocampus of *Aldh111-Cre^{ERT2}+;Bmal1^{fif}* and *Aldh111-Cre^{ERT2}+;Bmal1^{fif}* mice at 3 months (2 months after tamoxifen treatment), *CAG-Cre^{ERT2}+;Bmal1^{fif}* mice at 4 months (2 months after tamoxifen treatment), and *Bmal1^{fif}* controls. Scale bar, 100 µm.

All data represent mean + SEM. *p < 0.05, **p < 0.01, ***p < 0.001, and ****p < 0.0001 by 2-tailed t tests (C) and (G) or 1-way ANOVA (H) with Holm-Sidak correction for multiple comparisons when applicable.

mice, demonstrating the key role of decreased BMAL1 function in this phenotype. We next performed microfluidic qPCR profiling of global *Bmal1* KO hippocampal samples for markers of astrocyte activation and polarization (Liddelow et al., 2017), which demonstrated that *Bmal1* deletion increased most pan-reactive markers but did not clearly polarize astrocytes toward a neurotoxic or neurotrophic profile (Figure 3B). These genetic profiling studies suggest that *Bmal1* deletion induces a unique state of astrocyte activation. Although *Per1/2^{mut}* and NBmal1 KO mice are both behaviorally arrhythmic, expression of BMAL1 transcriptional targets *Nr1d1* and *Dbp* are elevated solely in *Per1/2^{mut}* mice because of loss of inhibition of BMAL1 (Figures 3A and S3A). *Gfap* is not elevated in *Per1/2^{mut}* mice (Figures 3A and S3A), suggesting that loss of BMAL1 activity, or at least blunting of BMAL1 oscillation (as in Figure 1A), is needed to induce astrocyte activation. In order to determine if *Gfap* and other selected activation markers are clock-controlled genes, we harvested cortex

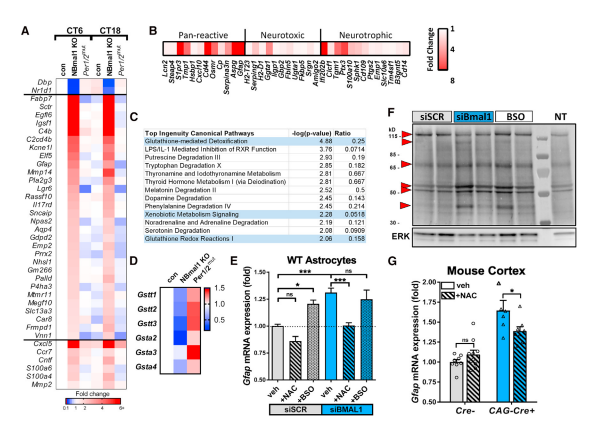


Figure 3. BMAL1 Regulates Astrocyte Activation via a Glutathionylation-Dependent Mechanism

(A) Astrocyte-specific genes upregulated in Nestin-Cre⁺; Bmal1^{1/f} (NBmal1 KO) cortex and changes to the same genes in Per1/2^{mut} mice. Transcripts upregulated at least 2-fold in NBmal1 KO were compared to the top 200 astrocyte-specific genes (Zhang et al., 2014), and overlapping genes are displayed. Other notable genes implicated in astrocyte activation and function were upregulated in NBmal1 KO cortex (bottom). The clock-controlled genes *Dbp and Nr1d1* are direct *Bmal1* transcriptional targets.

(B) Expression of neurotoxic, neurotrophic, and pan-reactive astrocyte activation genes (Liddelow et al., 2017) in 5-month-old global *Bmal1* KO versus WT littermate hippocampus assayed by microfluidic qPCR. Each data point represents the average of 4 *Bmal1* KO mice normalized to the average of 4 WT littermates.

(C) Results from Ingenuity Pathway Analysis canonical pathway analysis of NBmal1 KO microarray data. Pathways related to glutathione homeostasis are colored blue.

(D) Expression levels for downregulated glutathione-S-transferases in NBmal1 KO cortex versus *Bmal1^{fif}* controls. Expression levels in *Per1/2^{mut}* cortex included for comparison (based on microarray data).

(E) qPCR illustrating effect of glutathione manipulation on *Gfap* transcript levels following *Bmal1* knockdown. WT primary astrocytes treated with control siRNA (siSCR, gray bars) or siBmal1 (blue bars), with or without N-acetyl-cysteine (NAC; striped bars) to increase glutathione levels, or buthionine sulfoxime (BSO; speckled bars) to deplete glutathione. n = 9 samples/condition from 3 separate experiments.

(F) WT astrocytes transfected with siSCR, siBmal1, or siSCR +BSO (BSO) were treated with biotin-linked glutathione ethyl ester (bioGEE) to assess glutathionylation. Increased band intensity indicates decreased endogenous glutathionylation. Arrows denote altered bands. NT, not treated with bioGEE, showing nonspecific labeling.

(G) qPCR depicting effects of NAC treatment on *Gfap* transcript levels in cortex of inducible *Bmal1* KO mice (*CAG-Cre*^{ERT2}+;*Bmal1*^{1//}). Inducible *Bmal1* KO mice and Cre– control littermates were treated with NAC before, during, and after 5-day tamoxifen treatment to delete *Bmal1*. Mice were harvested 9 days after the start of tamoxifen. n = 6-10 mice per group.

All data represent mean + SEM. *p < 0.05, ***p < 0.001 by t test with Holm-Sidak correction for multiple comparisons.

from 4-month-old NBmal1 KO and Cre- controls at 4-hr intervals across one circadian cycle in constant darkness. *Gfap* mRNA levels showed no circadian oscillation in control or Cre+ mice and were elevated across the circadian cycle in Cre+ mice. These findings were duplicated in other markers of astrocyte activation, including *Aqp4* and *Mmp14* (Figure S3B). Thus, the astrogliosis phenotype in *Bmal1* KOs is not likely due to direct regulation of *Gfap* or other activation markers by the clock. To address potential mechanisms linking BMAL1 to astrocyte activation, we performed pathway analysis on our NBmal1 KO transcriptomic data using two bioinformatics tools: Ingenuity Pathway Analysis (IPA) and Database for Annotation, Visualization and Integrated Discovery (DAVID) Bioinformatics Resources 6.8 (Huang et al., 2009). We analyzed an inclusive list consisting of all transcripts that differed between control and NBmal1 KO at 6 p.m. with uncorrected p values < 0.01 (Table S1). IPA canonical pathway analysis identified glutathione-mediated detoxification

as the top hit (Figure 3C; p = 0.000013). Functional annotation clustering with DAVID showed that the most enriched gene cluster focused on glutathione-S-transferase (GST) activity, and 4 of the top 5 Kyoto Encyclopedia of Genes and Genomes (KEGG) pathways identified were related to glutathione metabolism (Figure S3C). GST enzymes catalyze the phase II detoxification of reactive intermediates in cells via adduction to glutathione and can alter cellular signaling via glutathionylation of proteins (Grek et al., 2013). We found that expression of several GST isoforms was suppressed in NBmal1 KO cortex and increased in Per1/2^{mut} cortex (Figure 3D), suggesting reciprocal regulation by the positive and negative limbs of the circadian clock. In support of our microarray analysis, we observed significant decreases in 2 of the 3 GST isoforms probed in cortex from 3-month-old CAG-Cre+;Bmal1^{f/f} mice 9 days after tamoxifen treatment (Figure S3D) and in 1 of the 3 GST isoforms probed in hippocampus from 3-month-old Aldh1l1-Cre+ mice 2 months after tamoxifen treatment (Figure S3E). We next asked if altered glutathione homeostasis within astrocytes linked Bmal1 deficiency with astrocyte activation. Our data clearly show that Bmal1 deletion induced changes in both Gfap mRNA and GFAP protein, which are very closely correlated across all models (see Figure S4A), so Gfap mRNA was used as a marker for astrogliosis in subsequent experiments. Treatment with N-acetyl cysteine (NAC), a glutathione precursor, completely prevented increased Gfap expression in astrocytes following Bmal1 KD (Figure 3E). Conversely, incubation of WT astrocytes with buthionine sulfoxime (BSO), which blocks glutathione production, lowered cellular glutathione levels (Figure S3F) and caused an increase in Gfap mRNA similar to that observed with Bmal1 KD (Figure 3E). Treatment of Bmal1-deficient astrocytes with BSO did not induce a further increase in Gfap, suggesting that BSO and Bmal1 KD are causing increases in Gfap through overlapping pathways. Interestingly, neither Bmal1 KO nor KD of Bmal1 in cultured primary astrocytes altered glutathione levels (Figure S3F). Additionally, the oxidation state of glutathione (ratio of reduced GSH to oxidized GSSG) was not altered in cortex from NBmal1 KO or Per1/2^{mut} mice (Figure S3G). As glutathione levels and oxidation state remained unchanged with loss of Bmal1, we next examined glutathione signaling via glutathionylation, which has been shown to oscillate in a circadian manner in the SCN (Wang et al., 2012). Western blot analysis of WT astrocytes treated with biotin-linked glutathione ethyl ester (bioGEE) revealed widespread changes in the pattern of S-glutathionylated proteins in Bmal1 KD cells, which were similar to changes seen in BSO-treated controls (Figure 3F). In general, numerous bands showed increased biotin labeling, indicating decreased glutathionylation, as would be expected with decreased GST expression. These data suggest that the observed astrogliosis may be due at least in part to altered protein glutathionylation resulting from impaired expression of specific GSTs.

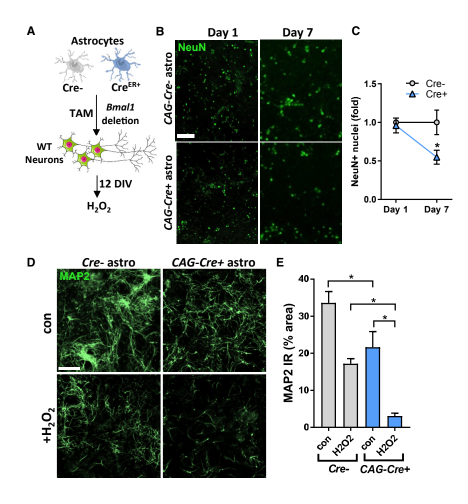
We next examined whether bolstering glutathione signaling can prevent astrogliosis following *Bmal1* deletion *in vivo*. We previously showed that tamoxifen-inducible global *Bmal1* deletion in mice (*CAG-Cre+;Bmal1*^{f/f} mice) causes astrocyte activation (Yang et al., 2016). We treated *CAG-Cre+;Bmal1*^{f/f} mice with NAC in their drinking water (40 mM beginning 5 days before tamoxifen) and via intraperitoneal injection (150 mg/kg daily beginning 2 days before tamoxifen) and induced *Bmal1* deletion via tamoxifen administration for 5 days. Both administrations of NAC were continued for the duration of the experiment. Cortex was harvested 9 days after the start of tamoxifen treatment and analyzed by qPCR. Tamoxifen treatment induced an 87% loss in *Bmal1* and a 79% loss in expression of the *Bmal1* target *Nr1d1* (Figure S3H), as well as a 64% increase if *Gfap* expression (Figure 3G). Administration of NAC was able to provide a significant but partial rescue of the *Gfap* increase seen in *CAG-Cre+;Bmal1^{frf}* animals (Figure 3G). These results suggest that *Bmal1* regulates *Gfap* expression at least in part through modulation of glutathione signaling *in vivo*.

Finally, we examined potential pathogenic consequences of Bmal1 deletion in astrocytes. We generated primary astrocyteenriched cultures from CAG-Cre+:Bmal1^{f/f} mice and Cre- littermates and treated them with tamoxifen in vitro to delete Bmal1. These cultures are primarily astrocytes, but this methods yields cultures which contain ~2% microglia and/or oligodendrocyte precursor cells (Schildge et al., 2013). Primary WT cortical neurons were then co-cultured on these astrocytes for up to 12 days, and a subset was subjected to oxidative stress by hydrogen peroxide exposure (100 μ M H₂O₂) for 24 hr (Figure 4A). We found no difference in the number of neurons surviving on Cre- and Cre+ astrocyte cultures 1 day after plating (as quantified by staining for neuronal nuclei with NeuN, as these cells do not yet have many MAP2 positive neurites) (Figure 4B). At day 7 after plating, there was a 45% decrease in NeuN+ nuclei (Figures 4B and 4C), a trend toward a decrease in MAP2 staining, and a trend toward an increase in cleaved-caspase-3 staining (Figures S4B and S4C) in the Cre+ astrocyte condition. The decrease in MAP2 staining in cultures containing Cre+ astrocytes became significant after 12 days (Figures 4D and 4E). H₂O₂ exposure at day 12 decreased MAP2 levels in Cre- cultures, and the combination of Bmal1 deletion and H2O2 killed nearly all neurons (Figures 4D and 4E). This finding shows that Bmal1-deficient astrocytes are less able to sustain neuronal viability, particularly in the face of oxidative stress.

DISCUSSION

Despite a large body of evidence describing circadian disruptions in both aging and neurodegenerative disease, little is known about the implications of circadian dysfunction on the brain. Here, we provide evidence that behavioral circadian disruption or disruption of the astrocytic molecular clock via manipulations of the master clock gene *Bmal1* induce astrogliosis and astrocyte dysfunction. The critical but complex role of astrocytes in brain pathologies and neuroinflammation underscores the importance of elucidating mechanisms of astrocyte activation, and our data illuminate BMAL1 as a potent regulator of astrocyte activation by both cell-autonomous and non-autonomous mechanisms.

The existence of the astrocyte circadian clock has been well documented (Prolo et al., 2005), and the critical role of the astrocyte clock in maintaining circadian rhythms in the SCN has recently been described (Barca-Mayo et al., 2017; Brancaccio et al., 2017; Tso et al., 2017). Clock genes have been shown to



regulate astrocytic glutamate uptake and ATP release (Beaulé et al., 2009; Marpegan et al., 2011), but the function of the core clock in astrocytes is otherwise unexplored. We have shown here, using several in vitro and in vivo methods, that loss of Bmal1 function in astrocytes causes cell-autonomous astrocyte activation. However, deletion of Bmal1 specifically in neurons also induces a partial astrocyte activation phenotype, suggesting a coexistent cell-non-autonomous mechanism. Bmal1 deletion in cultured neurons causes toxicity (Musiek et al., 2013), implicating a possible damage signal from Bmal1-deficient neurons that activates nearby astrocytes. Alternatively, the neuronspecific Bmal1 KO mice used in this study are behaviorally arrhythmic (Izumo et al., 2014), and this loss of rhythms could induce astrocyte activation, as seen in 10 hr:10 hr L:D-exposed animals (Figure 1). Thus, cell-non-autonomous influences on glial activation in the brain must be examined in the future.

One important question is whether the loss of rhythmic function of the circadian clock plays a key role in this phenotype or if this is a "non-circadian" function of BMAL1. Our data suggest that arrhythmicity in the setting of increased BMAL1 expression, as in *Per1/2^{mut}* mice, does not induce astrocyte activation, as these mice express increased levels of GSTs. Thus, astrocyte activation appears to be dependent on suppression of BMAL1mediated transcription. However, BMAL1 DNA binding is highly rhythmic and regulated by the clock (Koike et al., 2012), making it

Figure 4. Loss of *Bmal1* Impairs Astrocytic Support of Neurons

(A) Diagram depicting co-culture experimental design. DIV, days *in vitro*; TAM, tamoxifen.

(B) Representative images showing NeuN+ WT neuronal nuclei at DIV 1 and 7, grown on primary astrocytes from Cre – control mice, or inducible Bmal1 KO mice (iCAG-Cre+) treated with tamoxifen in vitro. Scale bar, 100 μm.

(C) Quantification of NeuN+ nuclei from (B) normalized to NeuN counts from Cre– condition. n = 8 replicates from 2 independent experiments. (D) Representative images of MAP2+ WT neurons grown as in (B) at DIV 12. Lower panels show neurons from adjacent wells treated for 24 hr with hydrogen peroxide at DIV 11 (H₂O₂, 100 μ M). Scale bar, 100 μ m.

(E) Quantification of MAP2 immunoreactivity (percent area) from (D). n = 9 replicates from 3 independent experiments.

All data represent mean \pm SEM. *p < 0.05, **p < 0.01 by 2-tailed t test.

difficult to separate entirely from circadian rhythms. *Bmal1* KD in astrocytes abrogates circadian *Per2*-luc oscillations, which could contribute to increased *Gfap* expression in cell culture. Moreover, we show in Figure 1 that circadian disruption by exposure of mice to 10 hr:10 hr L:D conditions, which blunts *Bmal1* oscillations in the brain, can induce *Gfap* expression. Thus, dampening the rhythmicity of BMAL1 expression through

circadian clock disruption likely plays a role in mediating astrogliosis, perhaps by restricting BMAL1 levels at certain key times of day. BMAL1 levels can be suppressed by several factors such as aging (Wyse and Coogan, 2010), inflammation (Curtis et al., 2015), or amyloid-beta (Song et al., 2015), all of which could potentially influence astrocyte activation and function in a BMAL1-dependent manner.

The activation profile induced in Bmal1-deficient astrocytes encompasses upregulation of a variety of genes across the predefined neurotoxic, neurotrophic, and pan-reactive categories (Liddelow et al., 2017). However, it is important to remember that these astrocyte phenotypes are thought to be induced by cytokine release from microglia, whereas activation due to loss of Bmal1 can be cell autonomous. Our findings contrast with those of Nakazato et al., who found that S100b-Cre+;Bmal1^{f/f} mice did not develop astrocyte activation and attributed the astrocyte activation phenotype of Nestin-Cre+;Bmal1^{f/f} mice to pericyte dysfunction (Nakazato et al., 2017). While our data do not exclude a contribution of blood-brain barrier dysfunction to the Nestin-Bmal1 phenotype, we provide multiple lines of evidence demonstrating cell-autonomous astrocyte activation. While Bmal1-deficient astrocytes do not assume a clear neurotoxic ("A1") phenotype, they are less able to support neuronal survival in a co-culture system and express numerous transcripts associated with inflammation and neurodegeneration. These include *C4b*, a complement component and marker of astrocyte activation and aging (Boisvert et al., 2018; Clarke et al., 2018), *Megf10*, which mediates astrocytic synapse elimination (Chung et al., 2013), and *Pla2g3*, a phospholipase implicated in ROS-induced neuronal damage (Martínez-García et al., 2010). Activation marker *Lcn2* (lipocalin 2), an anti-inflammatory mediator in the brain (Kang et al., 2018), was not increased, again suggesting a pro-inflammatory state.

Our data support a mechanism in which loss of BMAL1 induces astrogliosis through disruption of GST-mediated protein glutathionylation. In addition to its role in quenching oxidative stress, glutathione can form adducts on cysteine residues in a process termed glutathionylation, which is catalyzed by GSTs and can serve a signaling function (Grek et al., 2013; Mailloux and Treberg, 2016). Circadian rhythms of protein S-glutathionylation in the SCN may regulate neuronal excitability (Wang et al., 2012). Because Bmal1-deficient astrocytes have normal total glutathione levels, it is likely that disruption of GST-mediated glutathione signaling or utilization, but not glutathione depletion, mediates BMAL1-controlled astrogliosis. Accordingly, we observed an altered pattern of protein S-glutathionylation in Bmal1-deficient astrocytes, which resembled that seen in BSO-treated cells. Thus, supplementing Bmal1-deficient astrocyte cultures or mice with NAC presumably prevents astrogliosis by promoting nonenzymatic glutathionylation of protein targets, thereby circumventing the loss of GSTs. Further studies are needed to identify specific glutathionylation targets that mediate astrocyte activation and understand regulation of astrocyte glutathione signaling in health and disease.

In summary, our data identify the circadian clock protein BMAL1 as a cell-autonomous regulator of astrocyte activation and neurotrophic function and posit altered GST-mediated protein glutathionylation as a mediator of this effect. Our findings suggest that BMAL1 serves as a link between the core circadian clock and astrogliosis and provide insights into how the circadian clock might influence neurodegeneration.

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SUPPLEMENTAL INFORMATION

Supplemental Information includes four figures and one table and can be found with this article online at https://doi.org/10.1016/j.celrep.2018.09.015.

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AUTHOR CONTRIBUTIONS

B.V.L. and E.S.M. performed most experiments and data analysis. C.J.N. and M.R.C. generated mouse colonies. C.J.N., M.R.C., D.D.X., M.I., J.D., C.A.M., C.F.T., P.G., and P.W.S. performed various assays and imaging. S.A.L. performed microfluidic qPCR assays. I.N.K. performed circadian desynchrony experiment and analysis. Certain mouse lines/tissue and intellectual guidance were provided by M.I., J.A.H., B.A.B., and J.S.T. B.V.L. and E.S.M. wrote the paper.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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STAR***METHODS**

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Gfap – Rabbit polyclonal, 1:5000	Dako/Agilent	Cat# Z0334; RRID:AB_10013482
Gfap- Mouse monoclonal, 1:1000 (Co-stain only)	Novus	Cat# NBP1-05197; RRID:AB_1555288
3mal1- Rabbit Monoclonal, 1:2000 (IF/IHC only)	Novus	Cat# NB100-2288; RRID:AB_10000794
Bmal1- Rabbit polyclonal, 1:1000 (WB only)	Bethyl	Cat# A302-616A; RRID:AB_10555918
MAP2- Mouse monoclonal, 1:1000	EMD/Milipore	Cat# AB5622; RRID:AB_91939
Ki67- Rat monoclonal, SolA15, 1:500	ThermoFisher	Cat# 14-5698-82; RRID:AB_2016711
NeuN- Mouse monoclonal, 1:1000	EMD/Millipore	Cat# MAB377; RRID:AB_2298772
P42/44 MAPK ERK 1/2- Rabbit monoclonal, 1:1000 (WB)	Cell Signaling Technology	Cat# 9102; RRID:AB_330744
Bacterial and Virus Strains		
AAV8-GFAP-eGFP	UNC Viral Vector Core	N/A
AAV8-GFAP-GFP-Cre	UNC Viral Vector Core	N/A
Chemicals, Peptides, and Recombinant Proteins		
Lipofectamine RNAiMAX transfection reagent	ThermoFisher/Life	Cat# 13778075
N-acetyl-L-cysteine	Sigma	Cat# A7250
Tamoxifen	Sigma	Cat# T5648
L-Buthionine sulfoxime	Sigma	Cat# 5082280001
Poly-D-lysine hydrobromide	Sigma	Cat# P6407
Critical Commercial Assays		
BioGEE Glutathionylation assay kit	ThermoFisher	Cat# G36000
High Capacity RNA-to-cDNA Kit	Applied Biosystems	Cat# 4387406
GSH/GSSG Ratio Detection Assay Kit	Abcam	Cat# ab138881
Taqman Universal PCR Master Mix	Applied Biosystems	Cat# 4304437
PureLink RNA Mini Kit	Ambion/Life	Cat# 12183025
Deposited Data		
Agilent 4x44k mouse microarray v2 data from Nestin- Bmal1, Per1;Per2 double mutant, and control cortex at CT 6 and 18.	This paper	ArrayExpress (EMBL-EBI) Accession#: E-MTAB-7151
Experimental Models: Organisms/Strains		
B6.Cg-Tg(CAG-cre/Esr1*)5Amc/J (CAG-Cre ^{ERT2} mice)	Jackson Labs	Cat# 004682
B6;FVB-Tg(Aldh1l1-cre/ERT2)1Khakh/J (Aldh1l1-Cre ^{ERT2} mice)	Jackson Labs	Cat# 029655
B6.Cg-Tg(Nes-cre)1Kln/J (Nestin-Cre mice)	Jackson Labs	Cat# 003771
B6.129S4(Cg)-Arntltm1Weit/J (Bmal1 ^{f/f} mice)	Jackson Labs	Cat# 007668
CamKIIalpha iCre BAC (CamK2a-Cre mice)	Casanova et al., 2001	N/A
Oligonucleotides		
SMARTpool ON-TARGETplus Arntl siRNA	Dharmacon	Cat# L-040483-01-0005
ON-TARGETplus Non-targeting siRNA1	Dharmacon	Cat# D-001810-02-05
Gfap Taqman Gene expression assay	Applied Biosystems	Mm01253033_m1
Actb Taqman Gene expression assay	Applied Biosystems	Mm02619580_g1
Bmal1 (Arntl) Taqman Gene expression assay	Applied Biosystems	Mm00500223_m1
Aqp4 Taqman Gene expression assay	Applied Biosystems	Mm00802131_m1
Nr1d1 Taqman Gene expression assay	Applied Biosystems	Mm00520708_m1
Nr1d1 Taqman Gene expression assay Serpina3n Taqman Gene expression assay	Applied Biosystems Applied Biosystems	Mm00520708_m1 Mm00776439_m1

(Continued on next page)

Continued		
REAGENT or RESOURCE	SOURCE	IDENTIFIER
II6 Taqman Gene expression assay	Applied Biosystems	Mm00446190_m1
1/33 Taqman Gene expression assay	Applied Biosystems	Mm00505403_m1
C4b Taqman Gene expression assay	Applied Biosystems	Mm00437893_g1
Cxcl5 Taqman Gene expression assay	Applied Biosystems	Mm00436451_g1
Mmp14 Taqman Gene expression assay	Applied Biosystems	Mm00485054_m1
Timp1 Taqman Gene expression assay	Applied Biosystems	Mm01341361_m1
Dbp Taqman Gene expression assay	Applied Biosystems	Mm00497539_m1
Gsta2 Taqman Gene expression assay	Applied Biosystems	Mm03019257_g1
Gsta4 Taqman Gene expression assay	Applied Biosystems	Mm00494803_m1
Gstt3 Taqman Gene expression assay	Applied Biosystems	Mm00519528_m1
Software and Algorithms		
Ingenuity Pathway Analysis	QIAGEN	Build Version: 400896M; Content version 28820210
Imaris	Bitplane	Version 9
Prizm	GraphPad	Version 7
DAVID Bioinformatics Resource	https://david.ncifcrf.gov/	Version 6.8
Fiji	https://fiji.sc/	N/A

CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Erik Musiek (musieke@wustl.edu).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Mice

Nestin-Cre+, *CAG-Cre^{ERT2}*+, *Global Bmal1 KO*, *Aldh1L1-Cre^{ERT2}*+, and *Bmal1^{f/f}* mice were obtained from The Jackson Laboratory (Bar Harbor, ME) and were bred at Washington University. *Camk2a-iCre+;Bmal1^{f/f}* mice were bred at University of Texas Southwestern. All cohorts of mice were mixed sex with ages specified in figure legends. All mice were backcrossed at least 10 generations and were maintained on a C57Bl \ 6 background and housed under 12-hour light/12-hour dark conditions. In all cases, a Cre+; *Bmal1^{f/f}* male mouse was bred with *Bmal1^{f/f}* females to generate an equal mix of Cre- and Cre+ mice, all on the *Bmal1^{f/f}* background. All littermates were thus used for experiments. *Aldh1L1-Cre^{ERT2+}* and *CAG-Cre^{ERT2+}* mice (Cre- and Cre+ littermates from several litters) were given tamoxifen (Sigma, dissolved in corn oil, 2mg/mouse/day for 5 days) by oral gavage at 1mo or 2mo, respectively, to induce *Bmal1* deletion in the applicable tissue. Behavioral circadian desynchronization experiments were group housed for gene expression experiments (N = 4-5 per cage), or singly housed for actigraphy, with food and water available *ad libitum*. All mouse experiments were conducted in accordance with protocols approved by the Washington University Department of Comparative Medicine, or by the IACUCs at UTSW or WSU.

Primary cell cultures

Primary cells were extracted from mixed sex P1-3 (astrocytes) or E16-17 (neurons) CD1 WT, *Bmal1^{f/f}*, or *iCAG-Cre⁺;Bmal1^{f/f}* mice per experimental designation. Wild-type (WT) CD1 mice for culture experiments were obtained from Charles River Laboratories (Wilmington, MA). Mice were briefly anesthetized by cooling on ice, cleansed with ethanol, then decapitated with scissors, and the brain placed in ice-cold DMEM (Life Technologies). Cortices were dissected, stripped of meninges in ice-cold DMEM, and incubated in 0.05% Trypsin-EDTA (Life Technologies) at 37°C for 15 (neurons) or 20 (astrocytes) minutes. 37°C DMEM plus 10% FBS (GIBCO) was then added to neutralize trypsin, cortices were spun down for 5 minutes at ~300 g and gently triturated in 37°C DMEM plus 10% FBS (GIBCO). For astrocytes cultures, cells were then plated on 10cm or T75 tissue culture plates (coated for 2 hours at 37°C with 50 µg/mL poly-D-lysine (PDL- MP Biosciences, Santa Ana, CA), then rinsed twice with ddH₂O) at a concentration of 2 cortices (1 brain) per plate. Astrocytes were then grown to confluence (~10 days), then shaken at 37°C in a heated shaker at 225 rpm for 2 hours, followed by media change to remove adherent microglia. Cells were later lifted using Trypl-E dissociation reagent (Life Technologies) for 5 minutes at 37°C, washed with DMEM plus 10% FBS, and replated on 12-well (mRNA experiments),

48-well (neuronal co-cultures), or 96-well (GSH experiments) plates (coated with PDL as above) at a 1:2 dilution. Astrocyte-enriched cultures were then grown in DMEM+ 10% FBS + penicillin/streptomycin until used for experiments.

METHOD DETAILS

Behavioral Desynchronization

Activity was monitored by implanted Mini-mitter telemetry device (Star BioSciences). For circadian desynchronization, mice were transferred into a 20h light/dark cycle, with 10h of light (\sim 200 Lux) and 10h of dark for 6 weeks then harvested as described below at CT0, 6, 12, and 18; N = 4-5/group/time.

Intracerebroventricular injections

Bmal1^{f/f} mice were injected on postnatal day 2 (P2) with either AAV8-GFAP-GFP or AAV8-GFAP-Cre^{GFP} viral vectors (2 µL, bihemi-spheric, from UNC viral vector core), using a 30 g Hamilton syringe (10ul, 45 degree bevel) with a cuff to ensure proper depth (1.5mm). The injection site was 1mm lateral and 1mm posterior to bregma. Mice were housed as described above, harvested at 5 months of age and analyzed as described in the immunohistochemisty section.

Immunohistochemistry

Mice were anesthetized by i.p. pentobarbital (150mg/kg), followed by perfusion for 3 minutes with ice-cold Dulbecco's modified PBS (DPBS) containing 3 g/l heparin. One hemisphere was post-fixed in 4% paraformaldehyde for 24 hours (4°C), then cryoprotected with 30% sucrose in PBS (4°C) for 48 hours. 50-micron serial coronal sections were then cut on a freezing sliding microtome and stored in cryoprotectant solution (30% ethylene glycol, 15% sucrose, 15% phosphate buffer in ddH20). Sections were washed in Tris buffered saline (TBS), blocked for 30 minutes in TBSX (TBS with 0.25% Triton X-100 (Sigma-Aldrich, St. Louis, MO)) containing 3% goat serum, then incubated overnight in TBSX containing 1% goat serum plus primary antibody at 4°C. Sections were then washed in TBS, incubated for 1 hour at room temperature in TBSX with 1:1000 fluorescent secondary antibody, and mounted on slides using Fluoromount G. Epiflourescent images were obtained on a Nikon Eclipse 80i fluorescent microscope. A Nikon A1Rsi scanning confocal microscope was used at 20x for Ki67/GFAP co-localization, at 40x for *Aldh1L1-Cre* BMAL1/GFAP quantification, and at 60x for viral injection and representative *Aldh1L1-Cre* co-localization pictures.

Culture experiments

For neuronal cultures, triturated cells were transferred to a second tube to remove debris, then diluted in Neurobasal+B27+glutamine (Life Technologies) and counted prior to plating on astrocyte beds. For co-culture experiments, astrocytes were grown to confluence after replating in 48-well plates and neurons were added immediately following primary dissection at a concentration of 37,500 cells/ well in Neurobasal media plus B-27 supplement and glutamine. After 48 hours, 50% of the media was changed to Neurobasal plus anti-oxidant-free B-27 (AOF B-27, Life Technologies) and glutamine. Subsequently, 50% media changes were performed using Neurobasal plus AOF B-27 and glutamine every 3 days until neuronal DIV 7 or 11. In cultures grown to DIV 11, half of the wells received 100 μ M H₂O₂ for 24 hours. All wells were then fixed with 4% paraformaldehyde for 15 minutes, washed 3x with TBS and stained for MAP2, Cleaved-Caspase-3 or NEUN and quantified using protocol in *immunoreactivity quantification* section.

Immunoreactivity quantification

GFAP and MAP2 staining were quantified from 10x images using ImageJ software (NIH). GFAP 8-bit images were thresholded by hand to encompass all GFAP immunoreactivity, and a threshold value was set and maintained for all sections. GFAP percent area coverage was then calculated for each image. In general, 1 image from each section and 2-3 sections/mouse were averaged, and the average from each mouse used as a data point. For Ki67/GFAP co-localization, puncta were hand-counted from maximal intensity projections of z stacks in each region of interest. GFAP, BMAL1, and DAPI co-localizations were hand counted by 2 independent, blinded experimenters in 6 images/mouse (3 hippocampus, 3 cortex) with the average for each mouse reported. Cleaved-Caspase-3 puncta co-localizing with MAP2 were hand counted by 2 independent, blinded experimenters in 3-5 images/well over 11 wells from 3 independent experiments with averages/well reported.

qPCR

Flash-frozen brain tissue was homogenized with a mechanical homogenizer for 20 s in RNA kit lysis buffer (PureLinkTM RNA Mini Kit, Life Technologies, Carlsbad CA) plus 1% β -mercaptoethanol. RNA was then purified using the kit protocol. Cell lysates were homogenized in TRIzol (Life Technologies). The aqueous layer was then utilized, after chloroform extraction (added 1:5, sample then spun at 13000xg for 15 minutes), in RNA isolation using the kit protocol. RNA concentration was then measured by Nanodrop spectrophotometer and cDNA was made using a high capacity RNA-cDNA reverse transcription kit (Applied Biosystems/Life Technologies) with 1 μ g RNA used per 20 μ L reaction. Real-time quantitative PCR was performed with ABI TaqMan primers and ABI PCR Master Mix buffer on either the ABI StepOnePLus or QuantStudio 12k Real-Time PCR thermocyclers. During analysis, mRNA measurements were normalized to β -actin (*Actb*) mRNA levels. All Taqman qPCR primers were obtained from Thermo/Fisher.

Microfluidic qPCR was performed on flash frozen hippocampal tissue from 6mo *Global Bmal1 KO* and WT littermates mice at Stanford University using a BioMark Real-Time PCR system (Fluidigm, South San Francisco, CA) and analyzed as previously described (Liddelow et al., 2017).

Western blotting

Tissue samples were homogenized by sonication radioimmunoprecipitation (RIPA) buffer (Cell Signaling Technologies, Danvers MA) containing cOmplete protease inhibitors (Roche Lifesceinces USA, Indianapolis IN). Invitrogen nuPAGE Novex gels, SDS running buffer, and an iBlot transfer device were utilized per the manufacturer's instructions (Life Technologies). Bands were visualized using Lumigen TMA-6 chemiluminescence reagents (Lumigen, Southfield MI) on a Syngene GBOX imaging system. Band density was calculated using ImageJ software and normalized to ERK as a loading control.

Microarray

11mo NBmal1 KO, Cre- littermate, and *Per1/2^{mut}* mice were placed in constant darkness for 36 hours, then anesthetized with i.p. pentobarbital in the dark, followed by perfusion. Mice were harvested at CT 6 and 18 (6am and 6pm), 2-4 mice/genotype/time point. RNA was isolated from flash frozen cortex samples as above and submitted to the Genome Technology Access Center at Washington University for quality control, MessageAmp RNA library preparation, and Agilent 4x44k mouse v2 microarray. Raw data was normalized and analyzed using Partek Genomics Suite v6.6. For DAVID analysis, version 6.8 was used. The gene list from Table S1 was uploaded using Official Gene Symbol as the identifier, species = *Mus musculus*, and 692/842 genes were recognized. Two analyses were run: Functional Annotation Clustering (stringency: high), and KEGG pathway analysis. For Ingenuity Pathway Analysis (QIAGEN, Venlo, Netherlands), content version 28820210, build version 400896 was used, and the same gene list was employed (with ratios and p values). Cannonical Pathway analysis was performed. Dataset will be uploaded to GEO Datasets website (https://www.ncbi.nlm.nih.gov/gds/).

siRNA transfections

Astrocytes were transfected with siRNA using lipofectamine RNAiMAX (Life Technologies) in Optimem (LifeTechnologies) according to manufacturer's instructions 24 hours after replating. After 48 hours, media was changed to DMEM plus 10% FBS. SiRNAs targeting mouse *Bmal1* or non-targeting (designed to target no known mouse gene) were obtained from Dharmacon (Lafayette, CO). ON-TARGETplus SmartPool siRNA was used, which contains 5 separate siRNAs targeting the gene of interest which are pooled together (Bmal1: product L040483-01-005, non-targeting: D-001210-01-05). An siRNA:RNAiMAX ratio of 1:1.25) was used, and 40 pmol of siRNA (2uL of 20uM stock) was added to each well, using 12-well plates. Media was changed 24-48 hours later.

BSO and NAC experiments

For NAC experiments, astrocytes were incubated in growth media supplement with 1mM N-acetylcysteine (Sigma) starting 24 hours after transfection with siRNA, and for the remainder of the experiment. 72 hours before harvest, media was changed to DMEM plus 2% FBS and, where indicated, containing 250 µM buthionine sulfoxime (BSO, Sigma) and cells were harvested as described above at day 8 after siRNA treatment.

CAG-Cre+;Bmal1^{fif} mice and Cre- littermates (age 2-3 mo) used for *in vivo* NAC experiments were given either control drinking water or water containing 40mM NAC starting day 0 until sacrifice. Beginning on day 3 and continuing until sacrifice, mice were also given daily intraperitoneal injections of NAC (150mg/kg) or sterile DPBS for controls. Tamoxifen (Sigma, dissolved in corn oil, 2mg/mouse/day for 5 days) was administered by oral gavage to all mice beginning day 5 after water treatment. Water was refreshed once at day 7. Mice were perfused at day 14 (4 days after completion of tamoxifen) and tissue was prepared as described above.

Per2:luciferase Bioluminescence Recording

Primary astrocyte cultures were prepared from P1-2 PER2::LUC transgenic mice, and transfected with siRNA as above. 48 hours later, cells were synchronized by media change, and PER2::LUC activity was continuously monitored from a photomultiplier tube (HC135-11MOD, Hamamatsu) from transfected cortical astrocytes plated on a sealed 35mm Primaria dish (BD Biosystems, San Diego, CA) with 1 mL of serum-free DMEM supplemented with 2% B27 (Invitrogen), pen/strep, 10 mM HEPES, 0.35 g/L NaHCO3, 4.5 g/L D-glucose, 2% L-glutamine, and 0.1 mM beetle luciferin (Promega,Madison,WI). Acquired *in vitro* bioluminescence traces were detrended by division over a 24h running average and fitted with a damped sine function (Chronostar 2.0, PMID: 19299560). Only traces with a coefficient of correlation > 0.80 and period between 18-30h were considered circadian and rhythmic.

BioGEE glutathionylation assay

Primary astrocyte cultures were prepared and transfected with siBmal1 and siSCR siRNA as described above. At day 5 post-transfection media was changed to DMEM, +2% FBS, +penicillin/streptomycin. Also at day 5, 250uM BSO (Sigma) was added to a subset of wells to block glutathione synthesis. At day 7 post-transfection all media was refreshed (DMEM, +2% FBS, +penicillin/ streptomycin, +250uM BSO where appropriate) and 100uM BioGEE Glutathiolation Detection Reagent (Invitrogen, G36000) was added to each well except for one (non-treated control). After 3 hours, cells were lysed in RIPA buffer and processed as in western blotting section. As the entire size range of membrane was needed to blot for glutathione ethyl ester linked biotin, a parallel gel was run using the same sample mix in order to blot for loading control (p42/44 ERK). After transfer, the loading control membrane was blotted and imaged as above in the western blotting section. The membrane for detection of glutathionylation was washed and blocked as above except that TBS-T (tris buffered saline +0.125 tween) was replaced with PBS-T (phosphate buffered saline +0.125% tween) per method recommendation. After blocking, membrane was placed in 5% milk in PBS-T +streptavidin poly-HRP40 conjugate (Fitzgerald, Boston, MA, 65R-S104PHRP, 1:2000) for 1 hour, washed 5 times in PBS-T and imaged as in western blotting section.

Glutathione Measurements

GSH and GSSG levels were quantified in mouse cortex samples using the Glutathione Assay Kit (Cayman Chemical, Ann Arbor, MI). In cell culture experiments, glutathione measurements were obtained at day 7 using the Intracellular Glutathione Detection Assay Kit (Abcam, Cambridge, MA) by kinetic read (beginning immediately upon adding reagent and then every 5 minutes for 60 minutes) on a Tecan Infinite M1000 Pro fluorescent plate reader (Tecan, Switzerland).

QUANTIFICATION AND STATISTICAL ANALYSIS

Statistical analyses were performed using GraphPad Prizm v7.02. In general, data are represented by mean \pm SEM. In mouse experiments, "n" represents the number of mice and in cell culture experiments "n" represents the number of biological replicates. Specific statistical tests used, p value level definitions, and additional details are listed in figure legends. When multiple t test were performed, Holm-Sidak correction test for multiple comparisons was applied unless otherwise noted.

DATA AND SOFTWARE AVAILABILITY

The accession number for the microarray data reported in this paper is ArrayExpress: E-MTAB-7151. Array available at https://www.ebi.ac.uk/arrayexpress/experiments/E-MTAB-7151/

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Supplemental Information

Cell-Autonomous Regulation of Astrocyte

Activation by the Circadian Clock Protein BMAL1

Brian V. Lananna, Collin J. Nadarajah, Mariko Izumo, Michelle R. Cedeño, David D. Xiong, Julie Dimitry, Chak Foon Tso, Celia A. McKee, Percy Griffin, Patrick W. Sheehan, Jeffery A. Haspel, Ben A. Barres, Shane A. Liddelow, Joseph S. Takahashi, Ilia N. Karatsoreos, and Erik S. Musiek

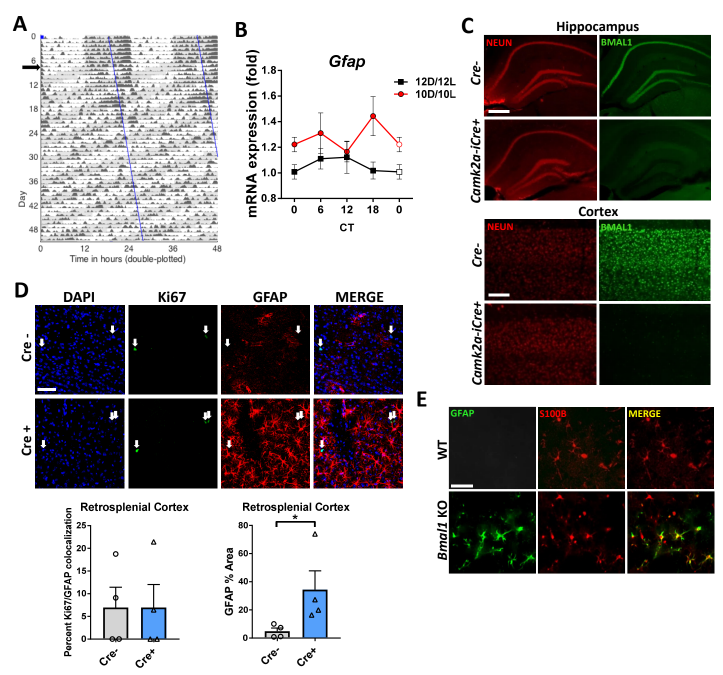


Figure S1. Related to Figure 1; Characterization of mouse models with astrogliosis induced by loss of *Bmal1*.

A. Representative actigraph of a WT mouse housed in 12h:12h light:dark, then switched to 10h:10h light:dark at black arrow, leading to behavioral arrhythmicity.

B. Cortical qPCR showing mRNA levels from 10h:10h light:dark WT mice from Fig. 1A, S1A reveals increased *Gfap* expression at most timepoints throughout the circadian day.

C. Hippocampal (top, scale bars = 150μ m) and cortical (bottom, scale bars = 50μ m) staining of NEUN (red) and BMAL1 (green) from 4mo *Camk2a-iCre+;Bmal^{#/f}* mice and Cre- controls shows complete loss of neuronal BMAL1 in Cre+ animals.

D. Representative images (top) and quantification (bottom) illustrating minimal localization of the cell division marker Ki67 staining with GFAP+ astrocytes in cerebral cortex from 4 mo *Nestin-Cre+;Bmal1^{ff}* (NBmal1 KO) mice (left graph), despite large increases in GFAP+ cells as compared to Cre- controls (right graph). Scale bar= 50µm.

E. NBmal1 KO mice have increased GFAP+ activated astrocytes in the cerebral cortex without major changes in number of total (S100B+) astrocytes. Scale bar = 50µm.

All data represent mean+SEM. *p<0.05 by 2-tailed T-test.

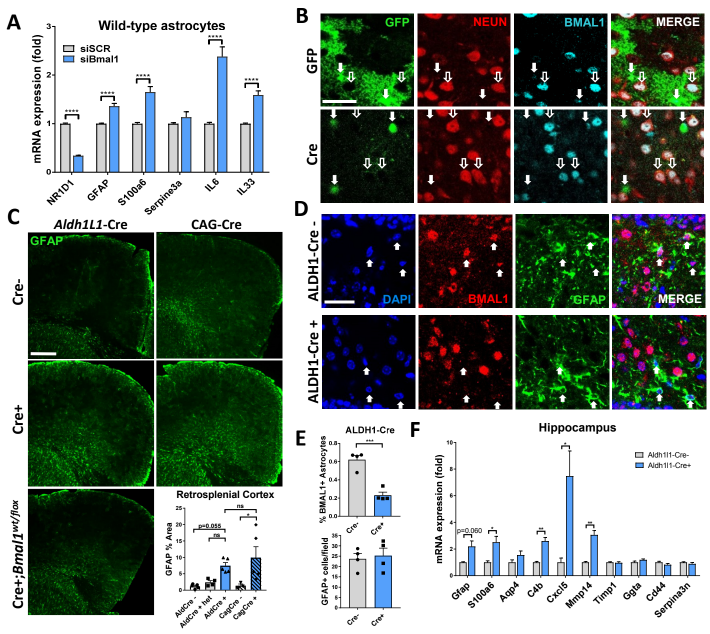


Figure S2. Related to Figure 2; Characterization of astrogliosis seen with loss of *Bmal1* in vitro and in vivo.

A. qPCR showing mRNA levels of several astrocyte activation and inflammatory genes 7 days after treatment of WT astrocytes with non-targeting siRNA (siSCR) or siRNA targeting *Bmal1* (siBmal1). N=12-15 experiments.

B. Representative cortical images from mice in Fig. 2F-H. *Top row*. AAV8-GFAP-GFP-infected cells (solid arrows) show whole-cell GFP expression, persistent colocalized nuclear BMAL1, and fail to colocalize with neuronal nuclei (hollow arrows). *Bottom row*. AAV8-GFAP-Cre infected cells show only nuclear GFP expression (solid arrows, Cre^{eGFP} fusion protein), loss of nuclear BMAL1 and fail to colocalize with neuronal nuclei (hollow arrows). Scale bar= 30µm.

C. Representative images, quantification of %GFAP area in retrosplenial cortex of mice in Fig. 2H. Scale bar = 100µm.

D. Representative confocal images of *Aldh111-Cre+;Bmal^{f/f}* mice and Cre- controls. DAPI+, GFAP+ (solid arrows) astrocytes colocalize with BMAL1 in Cre- mice, but not in Cre+ mice. Scale bar= 30µm.

E. Quantification of DAPI/GFAP/BMAL1+ cells (top) or GFAP+ cells per field of view (bottom) in *Aldh111-Cre+;Bmal^{f/f}* mice vs Cre- controls in hippocampus. Each data point = 1 mouse. 6 fields/mouse with an average of 24.5 GFAP+ cells/field, totaling 1,176 cells counted. Note that there is not an increase in GFAP+ cells in hippocampus, just the size and shape of these cells.

F. mRNA levels of astrocyte activation markers in hippocampus from Aldh111-Cre+;Bmal^{f/f} mice vs Cre- controls.

All data represent mean+SEM. *p<0.05, **p<0.01, ***p<0.001, ****p<0.0001 by 2-tailed T-tests (A), (E), and (F) or 1-way ANOVA (D) with Hom-Sidak correction for multiple comparisons when applicable.

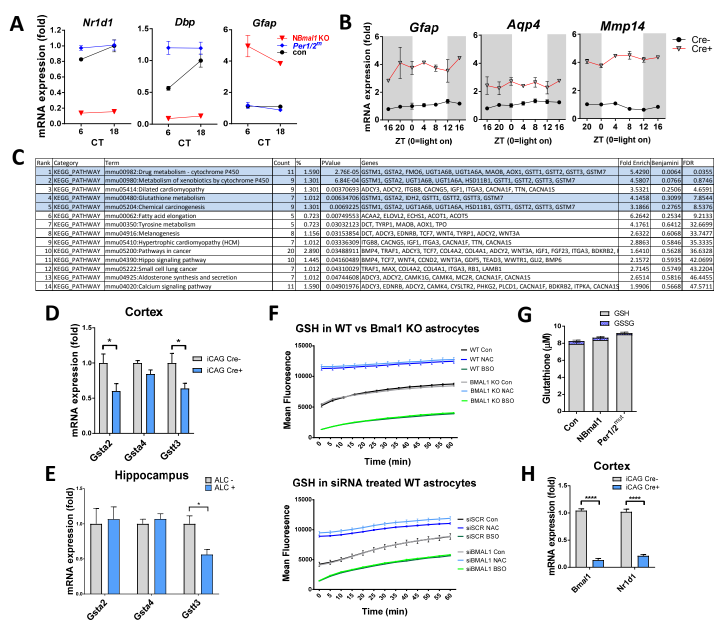


Figure S3. Related to Figure 3; Pathway analysis, GSTx, and GSH with *Bmal1* deficiency.

A. Expression of BMAL1 transcriptional targets and *Gfap* from microarray in Fig. 3A at CT 6 and 18.

B. Expression of astrocyte activation transcripts *Gfap*, *Aqp4*, and *Mmp14* in cortex of NBmal1 KO (Cre+, red) and Crecontrol mice (black) at 4 hour timepoints across the circadian cycle. No clear circadian oscillations were observed in Cre- or Cre+ mice. N=3-4 mice/genotype/timepoint.

C. KEGG pathway analysis of NBmal1 KO microarray data. Blue = pathways related to glutathione homeostasis.

D. qPCR of mRNA expression of glutathione transferases in the cortex of *iCAG-Cre+;Bmal1^{ff}* mice (Fig. 3F), 9 days after tamoxifen treatment, normalized to Cre-, *Bmal^{ff}* controls.

E. qPCR of mRNA expression of glutathione transferases in hippocampus of Aldh111-Cre+; Bmal1^{f/f} mice (Fig. 2H).

F. GSH levels in Bmal1 KO (top) and siBmal1 treated WT (bottom) astrocyte enriched cultures along with appropriate controls as measured by fluorometric intracellular glutathione detection assay.

G. Quantification of GSH and GSSG levels in cortical tissue (CT 10) from control, NBmal1 KO, and *Per1/2^{mut}* mice. None of the averages were significantly different.

H. mRNA expression of *Bmal1* and *Nr1d1* in the cortex of *iCAG-Cre+;Bmal1^{t/f}* mice used in Fig. 3G, 9 days after tamoxifen treatment, normalized to Cre-, *Bmal^{t/f}* controls.

All data represent mean±SEM. *p<0.05, ****p<0.001 by 2-tailed T-test with Holm-Sidak correction for multiple comparisons when applicable.

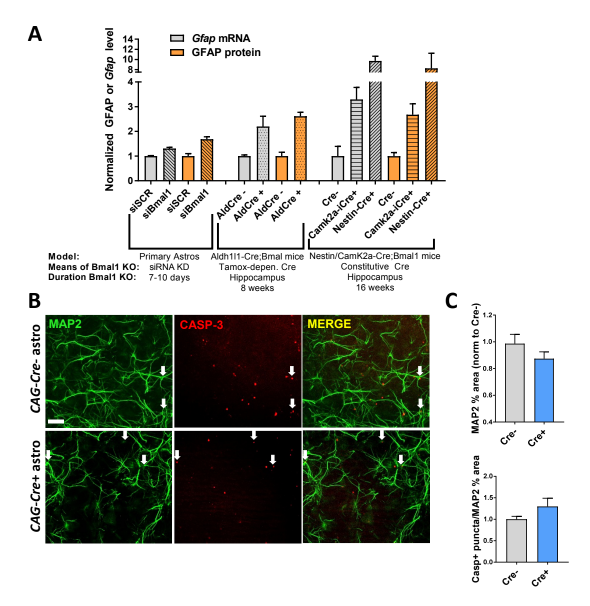


Figure S4. Related to Figures 3 and 4; Comparison of *Gfap* mRNA and GFAP protein across experiments; Further characterization of neuronal support deficits observed with loss of *Bmal1*.

A. Comparison of *Gfap* mRNA (quantified by qPCR) and GFAP protein (quantified by immunofluorescence) across experiments in this manuscript. Note that in both cultures cells and two distinct Cre-Lox mouse models, *Gfap* mRNA and GFAP protein are very closely related. Data is shown separately in Figs. 1 and 2.

B. Representative images showing MAP2+ (green) and cleaved-Caspase-3+ (CASP-3, red, arrows) WT neurons (grown as in Fig. 4A) at DIV 7. Scale bar = 100µm.

C. Quantification of MAP2 percent area (top) and cleaved-Caspase-3+ (bottom) neurons from (C) shows nonsignificant trends toward decreased MAP2 and increased cleaved-Caspase-3+ neurons when neurons are plated on BMAL1-deficient astrocytes (Cre+).

All data represent mean+SEM.