Regulation of persistent sodium currents by glycogen synthase kinase 3 encodes daily rhythms of neuronal excitability

Jodi R. Paul1, Daniel DeWoskin2, Laura J. McMeekin1, Rita M. Cowell1, Daniel B. Forger2 & Karen L. Gamble1

How neurons encode intracellular biochemical signalling cascades into electrical signals is not fully understood. Neurons in the central circadian clock in mammals provide a model system to investigate electrical encoding of biochemical timing signals. Here, using experimental and modelling approaches, we show how the activation of glycogen synthase kinase 3 (GSK3) contributes to neuronal excitability through regulation of the persistent sodium current ($I_{NaP}$). $I_{NaP}$ exhibits a day/night difference in peak magnitude and is regulated by GSK3. Using mathematical modelling, we predict and confirm that GSK3 activation of $I_{NaP}$ affects the action potential afterhyperpolarization, which increases the spontaneous firing rate without affecting the resting membrane potential. Together, these results demonstrate a crucial link between the molecular circadian clock and electrical activity, providing examples of kinase regulation of electrical activity and the propagation of intracellular signals in neuronal networks.
Much information processing in the brain occurs through electrical signalling, where the opening and closing of ion channels controls the properly timed firing of action potentials (APs). At the same time, biochemical networks within neurons separately regulate transcription, translation and post-translational modifications to determine cell fate, time cellular events and control cell size. How neurons integrate information between the two levels has remained a key open problem in neuroscience.

An important model system for studying this phenomenon is the neuronal network within the suprachiasmatic nuclei (SCN) of the hypothalamus, which forms the central circadian (24-h) pacemaker. Within the neurons of the SCN, an intracellular clock generates daily rhythms through regulation of transcription factors, kinases, and other signalling molecules. Timing information is relayed to the rest of the body through a variety of electrical behaviours and signals produced by SCN neurons. Understanding how cellular timekeeping and electrical activity are integrated is an essential ongoing question in the field. Since many of the molecular and electrical components of the clock are well characterized, the SCN is a perfect system for investigating the greater question in neuroscience of how intracellular biochemical information is electrically encoded.

One potential link between cellular biochemistry and electrical activity is glycogen synthase kinase 3 (GSK3). GSK3 activation enhances the spontaneous firing rate (SFR), is regulated in a circadian manner, and modulates molecular timekeeping within SCN neurons, providing a potential link between the molecular and electrical activities of the SCN. Here, we test the hypothesis that GSK3 regulates specific ionic currents that affect the spontaneous firing of APs. As GSK3 is implicated in many other intracellular processes (for example, inflammatory pathways, neurotrophic signalling, Wnt and mTOR cascades), GSK3 could provide a universal link between the biochemical state of the cell and its electrical activity.

It has been previously reported that several specific ionic currents are activated in a circadian manner, and thus, may provide a primary mechanism for controlling circadian variation in electrical activity. The presumed mechanism of circadian regulation of many currents, however, is through transcriptional regulation of their channels, which is energetically inefficient. Another challenge to this hypothesis is that experimental and computational studies implicate sodium currents as the most important regulators of firing rate in SCN neurons, but no evidence to support a day–night difference in these currents has been presented to date. Here, we propose another mechanism for the regulation of the electrical activity of a neuron by molecular signalling through GSK3. Using a combination of mathematical modelling and experimental work, we show that GSK3 efficiently regulates the firing rate of the cell by modulation of a persistent sodium current without causing the significant depolarization that could signal back to the molecular circadian clock.

**Results**

**GSK3 inhibition reduces spiking in a time-specific manner.** We initially determined whether acute regulation of GSK3 changes the electrical activity of SCN neurons during the day and night using the minimally invasive loose-patch technique. Compared to vehicle, inhibition of GSK3 with a pharmacological inhibitor (CHIR-99021, abbreviated CHIR throughout the manuscript; 1 μM) significantly decreased the SFR of SCN neurons from hypothalamic slices in a phase-specific manner (P = 0.018; Fig. 1a,b). GSK3 inhibition significantly suppressed SFR during the day between Zeitgeber time (ZT) 4 and 8 (P < 0.001). Conversely, at night (ZT 14–18), both treatment groups were similarly quiescent (P = 0.608). The observed day-time suppression was replicated with another GSK3 inhibitor (SB415286; 1 μM), which significantly decreased SFR by 66% compared with vehicle (mean ± s.e.m., vehicle: 5.05 ± 0.44; SB415286, 1.71 ± 0.28; n = 34–38 cells/group). These results suggest that inhibition of GSK3 suppresses SCN neuronal excitability during the day.

The firing rate of neurons can be regulated by numerous mechanisms, including depolarization of the membrane. We conducted whole-cell current clamp recordings to determine whether CHIR-induced suppression of excitability was due to membrane hyperpolarization. Consistent with recent reports, day-phase SCN neurons exhibited a wide range of resting membrane potentials (RMPs) which was not affected by GSK3 inhibition (P = 0.133; Fig. 1c). There was also no significant difference in input resistance (Supplementary Fig. 1a,b), suggesting that the decreased excitability in CHIR-treated neurons was not due to changed membrane properties.
neurons was not due to an increase in the potassium leak conductance\textsuperscript{17}.

As seen in the vehicle-treated controls during the mid-day (Fig. 1c), SCN neurons are capable of exhibiting multiple phenotypes in spontaneous electrical activity\textsuperscript{18}, and these characteristics (for example, spiking versus silent) can impact neurophysiological response\textsuperscript{19}. Using a k-means cluster analysis based on RMP and event amplitude, we classified each neuron into one of four categories while blind to treatment condition. Based on the results, SCN neurons were classified as either (i) silent, (ii) spiking, (iii) having depolarized electrical states with low-amplitude membrane oscillations or (iv) exhibiting depolarization block, in which spiking could be reinstated by injecting hyperpolarizing current (Supplementary Fig. 2) as reported in other brain areas\textsuperscript{20,21} as well as in the SCN\textsuperscript{16}. In the remaining spontaneously spiking neurons, APs were separated (or generated) by smooth depolarizing ramps to threshold (Fig. 1e). When GSK3 was inhibited, however, cells within this same voltage range seldom exhibited spontaneous APs (Fig. 1d). Interestingly, some cells displayed low-amplitude membrane potential oscillations. These oscillations were very similar to the effects of the voltage-gated sodium channel blocker, tetrodotoxin (TTX), when applied to SCN neurons\textsuperscript{12,22}. Given that the frequency of these oscillations was not different from vehicle-treated cells (Supplementary Fig. 1c), we hypothesized that, like TTX, GSK3 inhibition may modulate SCN excitability through suppression of a TTX-sensitive, voltage-gated sodium current. The voltage-gated sodium channels that are responsible for different components of Na\textsuperscript{+} current\textsuperscript{23} are expressed in SCN neurons (Supplementary Fig. 3), and at least Na\textsubscript{V}1.6 has been shown to be regulated by GSK3 (refs 26,27). Although small in magnitude, \(I_{\text{NaP}}\) typically contributes to excitation of repetitively firing neurons by augmenting small depolarizations during the interspike interval\textsuperscript{12}. In SCN cells, \(I_{\text{NaP}}\) is proposed to provide the excitatory drive towards threshold\textsuperscript{12}, and silencing \(I_{\text{NaP}}\) suppresses AP firing in SCN cells\textsuperscript{28}. Therefore, we examined the effect of GSK3 inhibition on \(I_{\text{NaP}}\) by measuring the current response to a slow (59 mV s\textsuperscript{-1}) depolarizing voltage ramp from –100 mV to 10 mV (ref. 29) before and after treatment with the selective persistent sodium channel blocker riluzole (20 \(\mu\)M, 3 min). The voltage ramp elicited an inward current in vehicle-treated cells at the range of \(I_{\text{NaP}}\) activation, starting at \(-52.4 \pm 1.1\) mV and peaking at \(-26.8 \pm 0.7\) mV (as in refs 30–32; means ± s.e.m.; \(n = 20\) cells). During the day, the magnitude of peak \(I_{\text{NaP}}\) was significantly reduced in CHIR-treated cells compared with vehicle-treated controls at baseline (\(P < 0.001\); Fig. 2a,b). Riluzole significantly suppressed the amount of inward current (\(P < 0.001\) and eliminated the

Figure 2 \(\text{GSK3 inhibition suppresses} I_{\text{NaP}} \text{ during the day but not the night. (a,b,d,e) Average normalized response to slow depolarizing voltage ramp (–100 to +10 mV; 59 mV s}^{-1}\) from SCN cells treated with (a,d) vehicle (DMSO, 0.002%; black) or (b,e) GSK3 inhibitor (CHIR, 1 \(\mu\)M, blue) at baseline and after 3-min treatment with persistent sodium current blocker, riluzole (20 \(\mu\)M, grey). (c,f) Peak inward current (mean ± s.e.m.) at baseline and after riluzole treatment. During the day (c) CHIR-treated cells (blue) had significantly reduced inward current at baseline (* \(P < 0.042\)) that was lost after blockade of \(I_{\text{NaP}}\) with riluzole (\(P = 0.350\)). Two-way, mixed-design ANOVA; Treatment X Riluzole interaction, \(F_{(1,33)} = 5.455, P = 0.026; n = 13\text{–}20\) cells, four animals per group. At night (f) peak inward current did not differ between CHIR- and vehicle-treated cells at baseline or after treatment with riluzole (non-significant main effect of CHIR, \(F_{(1,33)} = 0.150, P = 0.748; n = 13\text{–}22\) cells, four animals per group). Recordings made between ZT 4–8 (a-c) or ZT 14–18 (d-f).
difference between CHIR- and vehicle-treated cells (Fig. 2c). However, at night, CHIR did not suppress baseline $I_{\text{NaP}}$ compared with vehicle-treated controls (Fig. 2d–f). These results suggest that GSK3 inhibition decreases $I_{\text{NaP}}$ in day-phased SCN neurons and that this current is driven at least in part by GSK3 activation. Because circadian regulation of GSK3 inactivation increases throughout the night when SCN cells are mostly quiescent, GSK3 regulation of $I_{\text{NaP}}$ provides a previously unexplored mechanism underlying the daily rhythms in SCN excitability.

To investigate whether GSK3 regulates overall sodium current, we next measured the effect of GSK3 inhibition on $I_{\text{NaT}}$. In the presence of blockers (0.1 mM CdCl$_2$, 10 mM TEA, 3 mM 4-AP), the average peak $I_{\text{NaT}}$ was similar with and without the GSK3 inhibitor (peak $I_{\text{NaT}}$ in pA: vehicle, $-883.6 \pm 109.8$; CHIR: $-960.4 \pm 145.6$; $I_{(21)} = 0.43$, $P > 0.05$). While there was no change in the inactivation of $I_{\text{NaT}}$ (slope factor $V_{\text{C0}}$ vehicle: $5.8 \pm 1.2$; CHIR: $5.7 \pm 1.1$; $I_{(21)} = 0.03$, $P > 0.05$), GSK3 inhibition positively shifted the activation of this current by $-8$ mV (vehicle: $V_{\text{mid}} = -29.3 \pm 1.7$ mV; CHIR: $V_{\text{mid}} = -20.7 \pm 2.0$ mV; $I_{(21)} = 3.3$, $P < 0.05$). Thus, GSK3 inhibition did not appear to affect $I_{\text{NaT}}$ magnitude but may impact regulatory sodium channel subunits, such as one of the four $\beta$ subunits.$^{33}$

$I_{\text{NaP}}$ is regulated by time-of-day and activation of GSK3. Complementary to the present results that GSK3 inhibition suppressed SCN activity during the day but not the night, we have recently reported that chronic GSK3 activation increases neuronal firing at night, but not the day.$^{45}$ To determine if chronic GSK3 activation enhances $I_{\text{NaP}}$ in SCN neurons, we measured $I_{\text{NaP}}$ currents from wild-type (WT) mice and transgenic mice with constitutive GSK3 activation (that is, GSK3-KI mice) due to serine–alanine substitutions at the two inhibitory phosphorylation sites in subunits GSK3α and GSK3β (S21 and S9, respectively).$^{34}$ Although previous reports have suggested that $I_{\text{NaP}}$ does not exhibit a day/night difference in the SCN based on transcriptional gene expression,$^{35,36}$ until now, no study has examined $I_{\text{NaP}}$ isolated current at both times of the day.

We found that WT neurons exhibited a significant increase in $I_{\text{NaP}}$ during the day compared with the night ($P = 0.02$; Fig. 3a). Further, $I_{\text{NaP}}$ was enhanced in GSK3-KI cells compared with WT cells ($P = 0.008$; Fig. 3b,c). Surprisingly, the effect of chronic GSK3 activation was independent of day as both genotypes exhibited a day–night difference in $I_{\text{NaP}}$ magnitude ($P = 0.01$; Fig. 3b–d). Follow-up recordings from WT and GSK3-KI cells at night in the presence of CdCl$_2$ (0.1 mM) and TEA (10 mM) yielded similar results, yet this current could be blocked by a different Na$^+$ channel blocker, TTX (Supplementary Fig. 4). These results suggested that this TTX-sensitive, inward current was not driven by calcium or potassium currents and could be due to shunting or sequestration of constitutively active GSK3.

Mathematical model predicts GSK3 effect on $I_{\text{NaP}}$. To investigate how GSK3 affects neuronal excitability through $I_{\text{NaP}}$, we incorporated the $I_{\text{NaP}}$ current into a conductance-based single-cell model of SCN electrophysiology that simulates electrical activity with millisecond resolution based on sodium, potassium, calcium (L-type and non L-type), calcium-activated potassium and leak (Na-leak and K-leak) currents.$^{37}$ We modified the published model with the addition of $I_{\text{NaP}}$ with dynamics fit using the voltage ramp data (Figs 2 and 3) as described in the Methods section. We found that the dynamics of $I_{\text{NaP}}$ are consistent with a conductance-based model of the form $I_{\text{NaP}} = -g_{\text{NaP}}(V - E_{\text{NaP}})P$, with the activating gating variable $p$ ranging between 0 and 1. Currents for all experimental conditions could be fit to this form by varying the maximal conductance $g_{\text{NaP}}$ (Supplementary Fig. 5; $g_{\text{NaP}}$ values are GSK3-KI, day: 2.27; GSK3-KI, night: 2.13; WT, day: 2.09; WT, night: 1.59; CHIR, day: 1.97; CHIR, night: 1.46). This finding suggests that the action of GSK3 on $I_{\text{NaP}}$ is primarily through changing the maximal conductance of $I_{\text{NaP}}$ channels rather than the channel kinetics.

We validated the model after the addition of the $I_{\text{NaP}}$ current by performing voltage ramp simulations mimicking the analogous experiments presented in Figs 2 and 3 (Fig. 4). Simulated voltage ramps with and without $I_{\text{NaP}}$ matched the magnitudes of the inward currents seen experimentally in each condition: with simulated application of vehicle (Fig. 4a), CHIR (Fig. 4b), and with simulated constitutively active GSK3 (Fig. 4c), where each condition is reproduced simply by tuning the maximal conductance of the $I_{\text{NaP}}$ current ($g_{\text{NaP}}$).

$I_{\text{NaP}}$ mediates GSK3-induced excitability at night. Since the effects of $I_{\text{NaP}}$ have been shown to be different in isolated cells or in a network, we incorporated this ionic single-cell model into a previously validated SCN neuronal network model.$^{18,36}$ This model also included a validated model of the intracellular timekeeping system within individual cells. A total of 1,024 γ-aminobutyric acid-ergic (GABAergic) and vasoactive intestinal polypeptide-coupled SCN neurons were studied. Neurons within the network were excited or inhibited by GABA depending on cellular GABA reversal potentials, which had a distribution based on previously published experimental measurements.$^{38}$ This led to a distribution of excitation across the SCN causing WT cells to exhibit the varied firing activities reported here and in previous
WT and WT g
Fig. 2b and shows the effect of inhibiting GSK3 in our model. (P WT levels (X2 = 32.428, P < 0.001). Blocking hNaP with riluzole increased the proportion of silent cells in GSK3-KI slices (X2 = 44.735, P < 0.001) up to that of WT levels (X2 = 0.430, P = 0.313), whereas riluzole had no effect on the proportion of silent cells in WT slices (X2 = 0.034, P = 0.5). For d and f, WT and WT + riluzole, n = 60 cells, 2 animals; GSK3-KI, n = 84 cells, three animals; GSK3-KI + riluzole, n = 86 cells, three animals.

Figure 5 | Riluzole blocks GSK3-induced excitability. (a, b) Representative model recordings (1 s) from nighttime cell with hNaP at WT (a) or GSK3-KI (b) level shows that increasing hNaP alone is sufficient to induce spontaneous activity in a silent cell. Scale bar, 20 mV, 200 ms. Grey line indicates RMP (–63.4 mV). (c) Percentage of silent versus spiking neurons seen in SCN network model at night with WT or GSK3-KI levels of hNaP before and after hNaP blockade with riluzole. (d) Box plot of early-night spontaneous AP frequencies of SCN neurons from WT (grey) or GSK3-KI (red) mice treated with vehicle (DMSO, 0.01%) or riluzole (10 µM), showing 10th and 90th percentiles (whiskers), 25th and 75th percentiles (box borders), median (centre line) and outliers (symbols). Scheirer–Ray–Hare Kruskal–Wallis test, main effect of genotype, H(3) = 7.009, P < 0.001; main effect of treatment, H(3) = 8.089, P < 0.001 and interaction, H(3, 11) = 4.834, P < 0.001, post hoc asymptotic significance, P < 0.001. (e) Representative cell-attached loose-patch traces (5 s) from each group. Scale bar, 20 mV, 1 s. (f) Quantification of silent versus non-silent cells for each group in d and e. Three-way loglinear analysis, three-way interaction, X2(10) = 25.852, P < 0.001. Follow-up χ2 tests revealed that GSK3-KI cells were significantly more likely to be spiking than WT vehicle-treated cells (X2(10) = 32.428, P < 0.001). Blocking hNaP with riluzole increased the proportion of silent cells in GSK3-KI slices (X2(10) = 44.735, P < 0.001) up to that of WT levels (X2(10) = 0.430, P = 0.313), whereas riluzole had no effect on the proportion of silent cells in WT slices (X2(10) = 0.034, P = 0.5). For d and f, WT and WT + riluzole, n = 60 cells, 2 animals; GSK3-KI, n = 84 cells, three animals; GSK3-KI + riluzole, n = 86 cells, three animals.

Figure 4 | Modelling of voltage-ramp experiments. (a) Simulations of the voltage-ramp protocol in WT model cells with and without riluzole (R = 3.1, vehicle: gNaP = 2.08, riluzole: gNaP = 0). Our model matched the data collected in Fig. 2a and shows the effects of the persistent sodium current in the model. (b) Simulations of the voltage-ramp protocol under CHR with and without riluzole (R = 3.1, vehicle: gNaP = 1.46, riluzole: gNaP = 0). This matches Fig. 2b and shows the effect of inhibiting GSK3 in our model. (c) Simulations of the voltage-ramp protocol in the GSK3-KI compared with WT (R = 3.1, WT: gNaP = 2.08, GSK3-KI: gNaP = 2.85). This matches the data in Fig. 3b, and shows how the overactive kinase increases the persistent sodium current.

studies with circadian variation36. Using the model, we tuned the strength of INaP to mimic the transition from low to normal to high GSK3 activity (for example, CHIR to WT to GSK3-KI) and then examined the resulting changes in neuronal excitability. As gNaP is increased to the level of GSK3-KI mice, the model predicted that many of the cells which were normally quiescent at night became spontaneously active. Voltage for a sample quiescent WT neuron at night varied little from the RMP (Fig. 5a), whereas the identical simulated neuron with gNaP increased to GSK3-KI levels fired APs spontaneously (Fig. 5b). Furthermore, approximately half of cells in the WT network were quiescent at night, while in the GSK3-KI network, the majority of neurons were spiking (Fig. 5c). Since all parameters other than gNaP were identical between the simulations, the change in activity must have been due to the persistent sodium current. To test this model prediction, we examined the SFR of WT and GSK3-KI SCN neurons following application of riluzole (10 µM) and compared it to vehicle controls. Consistent with our previous
work, vehicle-treated GSK3-KI cells had a significantly increased SFR compared with WT neurons ($P < 0.001$; Fig. 5d,e). This hyper-excitability was rescued in riluzole-treated GSK3-KI cells ($P < 0.001$), which did not differ from either WT group. Since the majority of SCN neurons are relatively quiescent during the night, we also quantified the number of spiking versus non-spiking cells from WT and GSK3-KI mice in riluzole- and vehicle-treated conditions and found significant differences between groups ($P < 0.001$; Fig. 5f). While $\sim 40\%$ of WT neurons were silent, almost all of the GSK3-KI neurons were spiking ($P < 0.001$), as predicted by the model. Moreover, the ratio of spiking to non-spiking cells was restored to WT levels upon riluzole treatment, while riluzole had no effect on WT neurons at night ($P = 0.5$; Fig. 5g). Taken together, these results suggest that active GSK3 promotes neuronal excitability in an $I_{\text{NaP}}$-dependent manner.

**GSK3 promotes excitability through AHP but not RMP.** We next investigated how $I_{\text{NaP}}$ affects intrinsic excitability. Often, increases in excitability are due to depolarization of the RMP; however, the model unexpectedly predicted that increasing $I_{\text{NaP}}$ in firing neurons causes an increase in firing rate with minimal change to the RMP of the cells (Fig. 6a,b). This model prediction is consistent with our experimental results that GSK3 inhibition suppresses SCN activity without affecting RMP (Fig. 1).

Further analysis of our model simulations revealed that the increased excitation through $I_{\text{NaP}}$ is instead caused by suppression of the AP afterhyperpolarization (AHP). In both spiking and initially silent WT neurons at night, increasing the magnitude of $I_{\text{NaP}}$ by increasing $g_{\text{NaP}}$ from CHIR to GSK3-KI levels caused the minimum voltage attained after an AP to become more depolarized without changing the RMP (Fig. 6a,b). Thus the magnitude of the AHP diminished, allowing cells to fire more rapidly (Fig. 6a,b). Note that in the WT-silent neuron (Fig. 6b), increasing $g_{\text{NaP}}$ beyond a threshold level caused a bifurcation in the dynamics of the cell, and the onset of oscillations$^{37}$, as seen in Fig. 5. According to the model, the degree of AHP following an AP depends on the balance of $I_{\text{NaP}}$ (positive-inward) and $I_{\text{KCa}}$ (positive-outward) currents. By plotting each of the currents that contribute to changes in membrane voltage, it is apparent that most of the currents involved in generating the AP shut off rapidly (Fig. 6c). Only $I_{\text{NaP}}$ and $I_{\text{KCa}}$ persist through the interspike interval, setting the AHP magnitude and the depolarizing ramp back towards threshold. Increasing $I_{\text{NaP}}$ or decreasing $I_{\text{KCa}}$ causes a decrease in the AHP magnitude and consequently speeds up the firing rate. Conversely, decreasing $I_{\text{NaP}}$ or increasing $I_{\text{KCa}}$ increases the AHP magnitude and slows firing (Fig. 6d).

To test the prediction that increasing $I_{\text{NaP}}$ decreases AHP without affecting RMP experimentally, we performed whole-cell current clamp recordings in WT and GSK3-KI neurons during the day and night (Fig. 7a,b). As predicted by the model, there was no correlation between peak $I_{\text{NaP}}$ and RMP ($r = -0.094$, $P = 0.463$; Supplementary Fig. 6). Both genotypes exhibited a day/night difference in RMP ($P = 0.008$), and there was no difference in RMP between genotypes for the majority of neurons (two-way analysis of variance (ANOVA), with depolarization block cells excluded; $P = 0.433$; Fig. 7c). Only when the subset of neurons in depolarization block were included in the analysis did constitutive activation of GSK3 result in a more depolarized RMP (two-way ANOVA, with all cells included; $P = 0.016$; Fig. 7c).

**Figure 6** | Model predicts $I_{\text{NaP}}$ promotes excitability through AHP but not RMP. (a,b) Model prediction of RMP (black), minimum voltage attained after an AP (AHP; blue) and SFR (red) at different levels of $I_{\text{NaP}}$ for a WT-spiking (a) or WT-silent (b) SCN neuron at night. The WT-silent neuron does not spike with low to moderate levels of $I_{\text{NaP}}$ and has a SFR of zero and no AHP. When $g_{\text{NaP}}$ increased past a threshold level, the cell undergoes a ‘bifurcation’ in its dynamics and begins to fire. When either neuron fires, increasing $g_{\text{NaP}}$ increases the SFR without changing the RMP by decreasing the magnitude of the AHP. (c) Various currents (colour-coded in legend) associated with a typical SCN AP. During the AHP and ramp back to threshold, all currents other than $I_{\text{NaP}}$ and $I_{\text{KCa}}$ have inactivated. Scale bar, 40 ms. (d) Heat map showing the contributions of $g_{\text{NaP}}$ and $g_{\text{KCa}}$ towards setting the SFR (in Hz) of SCN neurons.
I regulation of kinase GSK3 (ref. 5) promotes neuronal excitability through Here, we present evidence that the rhythmically phosphorylated

Discussion

WAY ANOVA; main effect of time, significantly more depolarized during the day and in GSK3-KI slices. Two-early-night. Lines indicate mean ± s.e.m. for each group. Cells were significantly more depolarized during the day and in GSK3-KI slices. Two-way ANOVA; main effect of time, F(1,64) = 7.474, P = 0.008; main effect of genotype, F(1,64) = 6.099, P = 0.016. From left-to-right, n = 21 cells, four animals; 19 cells, three animals; 11 cells, three animals; 17 cells and three animals. The effect of genotype on RMP was driven by groups of cells in depolarized block (non-significant main effect of genotype when depolarized block cells were excluded from analysis; F(1,64) = 0.623, P = 0.433). Symbol colour indicates subgroup phenotype as indicated in the legend. (d) Model prediction of effect of NaP conductance (gNaP) on RMP shows that increasing gNaP causes a cell in depolarization block (red) to further depolarize but has no effect on the RMP of a hyperpolarized, silent cell (black).

This effect was predicted by the model as well since the voltage-gated sodium channels responsible for INaP are held open at the higher voltages seen in depolarization block-type neurons, leading to an increase in RMP (Fig. 7d). Additionally, analysis of the AP waveforms of the spontaneously active cells from the same recordings revealed a significant decrease in the AHP in the GSK3-KI neurons (P = 0.012), and in the RMP of all the neurons (F(1,64) = 0.302, P = 0.589). Further, consistent with the model prediction, both the AHP amplitude and the SFR of all spontaneously spiking cells are significantly negatively correlated with INaP peak current, such that the larger the INaP magnitude, the smaller the AHP (r = -0.732, P < 0.001; Fig. 8c) and faster the SFR (r = -0.550, P = 0.001; Fig. 8d). To directly test whether INaP magnitude influences AHP, a follow-up current clamp experiment was conducted to measure the AHP of individual SCN neurons before and after acute application of riluzole (10 μM). As shown in Supplementary Fig. 6c, blockade of INaP significantly induced a 70% increase in AHP amplitude (n = 5; mean change ± s.e.m.: 1.66 ± 0.63 mV; P = 0.001), but had no effect on RMP (mean change ± s.e.m.: -0.23 ± 0.42 mV; P = 0.605).

Discussion

Here, we present evidence that the rhythmically phosphorylated kinase GSK3 (ref. 5) promotes neuronal excitability through regulation of INaP. In particular, we find that: (1) GSK3 activity regulates INaP, (2) INaP exhibits a day/night difference in peak magnitude that is regulated by GSK3 and (3) INaP promotes firing and decreases AP AHP without affecting RMP. Our experiments were conducted in the SCN, the site of the central circadian pacemaker. As molecular circadian timekeeping can be found in many parts of the brain38, this mechanism could be widely used in the brain to control many systems. Of particular interest are the regions of the brain that regulate mood. Impaired molecular circadian timekeeping has been implicated in mood disorders39, and differentiated neurons from induced pluripotent stem cells isolated from patients with bipolar disorder show enhanced SFRs and over-excitability40. Given that GSK3 phosphorylation and INaP are altered by mood stabilizing drugs such as lithium41 and riluzole42, respectively, this work may provide a potential basis for the chronotherapeutic control of mood.

Despite the known importance of INaP as a pacemaker potential in many types of spontaneously active neurons43–46, including the SCN12,28,39, INaP has been largely overlooked as a potential contributor to the circadian modulation of rhythmic firing3,28,47. However, the present study is the first to show that INaP is greater in magnitude during the day (~5 pA) than at night. This difference in INaP magnitude is sufficient to have a notable impact on excitability of complex, high RMP neurons such as those in the SCN. Prior work has shown that blockade of INaP with riluzole acutely silences spontaneous firing48 and chronically damps the circadian amplitude of AHP rhythms of individual SCN neurons47. Conversely, our results demonstrate that high, day-like INaP levels seen in GSK3-KI neurons at night are sufficient to induce spiking in neurons that would normally be silent (Fig. 6) as well as increase the SFR of already spiking neurons (Fig. 7a). Moreover, lower INaP level following GSK3 inhibition reduces the SFR (Figs 1 and 2), while transient sodium current was minimally affected.

Although GSK3 may impact neurophysiological properties of neurons through regulation of other ionic conductances yet to be identified (including IkCa), our results show a novel effect of GSK3 activity on excitability through modulation of INaP. This GSK3-mediated increase in INaP may be the result of several different channel protein modifications including channel trafficking, distribution and phosphorylation state49. In fact, recent work in the hippocampus has shown that fibroblast growth factor 14 forms a complex with at least one sodium channel alpha subunit (that is, NaV1.6) in a GSK3-dependent manner26. It is interesting to note that four of the alpha subunits implicated in INaP are expressed in the SCN (Supplementary Fig. 3), with enrichment specifically for NaV1.3. In addition to alpha subunits, GSK3 may also activate other regulatory subunits of voltage-gated sodium channels, such as the NaV β subunits, which can enhance INaP magnitude33. Future research will be necessary to elucidate the contribution of the various TTX-sensitive NaV channel subunits responsible for day–night differences in INaP magnitude in SCN neurons.

Our work also shows how changes in INaP permeate throughout the SCN network. Since the SCN is coupled via mainly inhibitory GABA, increasing the firing rate causes more inhibitory post-synaptic currents, which could slow firing. However, inhibitory post-synaptic currents can both increase and decrease firing in SCN neurons50, so the effects of inhibitory GABA are not always intuitive. Nevertheless, our modelling and experimental data show an increase in firing rate throughout the network, and in particular, a change in the electrical state of neurons that produces an increase in the proportion of neurons that are firing. Together, this work suggests GSK3 as a way of modulating not just firing of a single cell, but also global network properties. Future work can improve our mathematical models by considering the morphology of SCN neurons, additional neurotransmitters and neuropeptides besides GABA and the stochasticity of neuronal firing.
Further clarification of the details of GSK3 regulation is an interesting question for future study. Despite much work, how the circadian clock precisely regulates GSK3 activity is unknown. In addition to rhythmic inhibitory phosphorylation, GSK3 function within the SCN could also be reduced by sequestration, localization or through priming kinases. Although suppression of \( I_{\text{Na}} \) by pharmacological GSK3 inhibition was time-of-day specific (that is, restricted to the day, when GSK3 activity is high), GSK3-KI mice with chronic GSK3 de-phosphorylation had enhanced, but unexpectedly rhythmic \( I_{\text{Na}} \), suggesting that other mechanisms of GSK3 inhibition may still be rhythmic in these mice. Alternatively, we cannot rule out a compensatory mechanism or developmental effect as an explanation of this result; however, it should be noted that no gross developmental differences have been previously reported in this animal model.

By combining modelling and experimental approaches, we present an intriguing method by which the molecular circadian clock controls electrical activity. Unlike Na\(^+\) leak currents which were recently shown to promote excitability of clock neurons by membrane depolarization, our data show a mechanism for how \( I_{\text{Na}} \) regulates the firing rate of APs without affecting the RMP of neurons that are hyperpolarized or spiking. We also show that the balance between \( I_{\text{Na}} \) and \( K_C \) controls both the depolarizing ramp towards the AP threshold and, surprisingly, the magnitude of the AHP following an AP. In fact, pharmacological inhibition of \( I_{\text{Na}} \) increased AHP magnitude without affecting RMP, consistent with a prior study that showed that reduction of \( I_{\text{Na}} \) due to genetic loss of SCN8A significantly increases peak AHP in cerebellar granule cells. Thus, sub-threshold ionic balance could be crucial to rhythmic changes in AP frequency independent of circadian phase, especially in light of a previous modelling/experimental study which showed that changes in RMP can feedback to phase shift the circadian clock. The mechanism we propose, therefore, can faithfully reflect the state of the circadian clock in firing rate without disrupting molecular circadian timekeeping itself.

Intriguingly, aging-induced circadian decline is associated with reduced SCN firing rate amplitude and smaller nocturnal AHP in the absence of altered PER2 expression, leading to the proposal that SCN neuronal activity may be the ‘weak link’ in the circadian system. Given that p-GSK3 levels are decreased in the SCN of aged rodents, GSK3 control of \( I_{\text{Na}} \) may be one source of age-related circadian disruption. While more work is necessary to explore this concept as well as the biophysical mechanism by which GSK3 activates \( I_{\text{Na}} \), this work suggests a promising new motif in which a kinase can directly regulate electrical activity acutely and without transcriptional cost. Because of its efficiency, this is a motif likely to be found in other brain regions as well.

**Methods**

**Animals.** Homozygous Per1::GFP, WT C57-BL6/J, and GSK3\(^{−/−}\) (GSK3-KI) mice (2–5 months old; male and female) backcrossed for at least 10 generations to C57BL/6 mice, in which regulatory serine–alanine substitution on both isoforms of GSK3 rendered GSK3 de-phosphorylated and constitutively active were group-housed (2–5 animals per cage) in a 12:12 light/dark cycle with ad libitum access to food and water, and handled in accordance with the University of Alabama at Birmingham (UAB) Institutional Animal Care and Use Committee (IACUC) and National Institutes of Health (NIH) guidelines. Mice were euthanized with cervical dislocation and rapid decapitation.

**Electrophysiology.** All chemicals were obtained from Sigma-Aldrich (St Louis, MO) unless otherwise specified. For recordings in which the primary outcome was

**Figure 8 | GSK3 decreases the AHP magnitude through \( I_{\text{Na}} \)** (a) Average AP waveforms from spontaneously active SCN neurons from WT (black) or GSK3-KI (red) mice recorded in mid-day or early-night. For visualization, all waveforms were adjusted to same baseline (−40 mV, dashed line) before averaging. Scale bar, 10 mV, 10 ms (b) Mean ± s.e.m. of AHP magnitude (difference from RMP) from cells represented in a. AHP magnitude was significantly decreased during the day and in GSK3-KI cells. Two-way ANOVA; main effect of time, \( F_{(2,30)} = 10.266, P = 0.003 \); main effect of genotype, \( F_{(1,30)} = 7.085, P = 0.012 \). For a and b: \( n = 4–11 \) cells, three animals per group. (c) Amplitude of AHP and peak \( I_{\text{Na}} \) of spiking cells from all groups are significantly correlated. Pearson correlation, \( R = -0.732, P < 0.001 \). Symbol colour indicates group as indicated in the legend. Lines represent linear fit and 95% confidence intervals for all cells in plot. (d) Spontaneous AP frequency and peak \( I_{\text{Na}} \) of spiking cells from all groups are significantly correlated. Pearson correlation, \( R = -0.550, P = 0.001 \). Lines represent linear fit and 95% confidence intervals for all cells in plot. For c and d: \( n = 4–10 \) cells per group.
AP firing rate, the loose-patch recording technique was chosen, given that it does not affect endogenous membrane properties either by formation of a tight seal or by dialysis of intracellular milieu. Sample sizes were based on previous work.  

Whole-cell electrophysiology. Slices were prepared using the same methods and solutions described previously and above. Riluzole-sensitive current was measured using a slow depolarizing voltage ramp (−100 mV to +10 mV, 59 mV−1) before and after 3 min of riluzole (20 μM) application. This ramp speed was chosen because it was in the middle of the range used by prior studies in SCN neurons. Moreover, this ramp speed produced maximal current without compromising voltage control (similar to ref. 29), and the same ramp speed was used for all experiments. At least 5–10 sweeps of the voltage ramp protocol were averaged for each cell, and baseline subtracted by fitting the linear portion between −85 mV and −65 mV to zero. Peak sodium current was determined as the minimum point of a 4–7th order polynomial fit applied to the baseline subtracted curve between −130 mV and −10 mV. For current clamp recordings, cells were recorded in gap-free current clamp mode. RMP was determined as the average voltage half-way between two APs. Cells were classified by neurophysiological phenotypes based on RMP and spontaneous event amplitude. Using a k-means cluster analysis, each neuron was grouped into one of four categories while blind to experimental group. SCN neurons were classified as either (1) spiking, (2) spiking, (3) having depolarized electrical states with low-amplitude membrane oscillations or (4) exhibiting depolarization block. For cells classified as spiking, the AP AHP was measured using the template search function in Clampfit 10 (Axon Instruments) to determine the average anti-peak amplitude for each cell. In all whole-cell experiments, recordings were made within 5 min of breaking into the cell. Cells with >−50 pA of leak recorded in the seal test, with excessive break through spikes in voltage-ramp protocol (indicative of poor voltage control), or which did not exhibit a rebound spike after terminating a hyperpolarizing current injection were excluded from the analysis. Supplementary Figure 2 depicts an example recording from a depolarized block neuron that was induced to spike with hyperpolarizing current.  

Quantitative, real-time RT-PCR. Slices (600 μm thick) containing the SCN were sectioned in Hanks’ balanced salt solution (14175-103; Gibco, Carlbad, CA) supplemented with NaHCO3 (7.5%; Sigma, St Louis, MO), and 1.0 M HEPES according to previously described methods. The SCN was carefully dissected from the slice under a Zeiss dissection microscope, and tubes containing either the SCN or the remaining brain tissue from the slice were rapidly frozen in liquid nitrogen in RNAase-free tubes and stored at −80 °C until use. RNA isolation and transcript measurements were performed as described in ref. 59. Samples were homogenized using an Omni Bead Ruptor Homogenizer (OMNI International, Kennesaw, GA, USA). Following homogenization, samples from one male and one female were combined to give a total of six samples for each group across the whole brain. RNA was isolated using the Trizol/chloroform-isopropanol method (Invitrogen, Carlbad, CA, USA) per manufacturer’s protocol and purity and concentration were determined using a Thermo Scientific NanoDrop2000 (Fisher Scientific, Pittsburg, PA, USA). Equivalent amounts of RNA (1 μg) were treated with DNAse I (Promega, Madison, WI, USA) at 37 °C for 30 min followed by RNasefree DNase Stop solution at 65 °C for 15 min and reverse-transcribed using the High-Capacity cDNA Archive Kit (Applied Biosystems, Carlbad, CA, USA). Transcription measurement was performed using JumpStart Taq ReadyMix (Sigma) and Applied Biosystems primers with an initial ramp time of 2 min at 50 °C and 10 min at 95 °C followed by 40 subsequent cycles of 15 s at 95 °C and 1 min at 60 °C. Relative concentrations of the genes of interest were calculated in comparison to a standard curve generated from a doubled concentration as well as 1:5, 1:10, 1:20 and 1:40 dilutions of a pooled cDNA of whole-brain samples (that is, the calibrator method). Taqman primer/probe information is included in Supplementary Table 1 for the following transcripts: Scn1a, Scn2a, Scn3b, Scn4b and Scn8a. Values were normalized to β-actin and expressed as fold change in SDS.
with constitutive activation of glycogen synthase kinase 3. Neuroscience 226, 1–9 (2012).

5. Bear, M. F. et al. Circadian rhythmicity of active GSK3 isoforms modulates molecular clock gene rhythms in the suprachiasmatic nucleus. J. Biol. Rhythms 30, 155–160 (2015).

6. Itakura, C., Miyazaki, K., Akaike, T. & Ishida, N. A role for glycogen synthase kinase-3beta in the mammalian circadian clock. J. Biol. Chem. 280, 29397–29402 (2005).

7. Hur, E. M. & Zhang, F. Q. GSK3 signalling in neural development. Nat. Rev. Neurosci. 11, 539–551 (2010).

8. Li, X. & Jope, R. S. Is glycogen synthase kinase-3 a central modulator in mood regulation? Neuropsychopharmacology 35, 2143–2154 (2010).

9. Taelman, V. F. et al. Wnt signaling requires sequestration of glycogen synthase kinase 3 inside multivesicular endosomes. Cell 143, 1136–1148 (2010).

10. Beurel, E., Michalek, S. M. & Jope, R. S. Inactive and adaptive immune responses regulated by glycogen synthase kinase-3 (GSK3). Trends Immunol. 31, 24–31 (2010).

11. Sim, C. K. & Forger, D. B. Modeling the electrophysiology of suprachiasmatic nucleus neurons. J. Biol. Rhythms 22, 445–453 (2007).

12. Jackson, A. C., Yao, G. L. & Bean, B. P. Mechanism of spontaneous firing in dorsomedial suprachiasmatic nucleus neurons. J. Neurosci. 24, 7985–7998 (2004).

13. Nunemaker, C. S., DeFazio, R. A. & Moenter, S. M. A targeted extracellular approach for recording long-term firing patterns of excitable cells: a practical guide. Biol. Proced. Online 5, 53–62 (2003).

14. Kuhlman, S. J. & McMahon, D. G. Encoding the ins and outs of circadian clock neurons. PLoS Comput. Biol. 9, e1003196 (2013).

15. Montgomery, J. R. & Meredith, A. L. Genetic activation of BK currents in vivo generates bidirectional effects on neuronal excitability. Proc. Natl Acad. Sci. USA 109, 18997–19002 (2012).

16. Raman, I. M., Gustafson, A. E. & Padgett, D. Ionic currents and spontaneous firing in neurons isolated from the cerebellar nuclei. J. Neurosci. 20, 9004–9016 (2000).

17. Sakhi, K., Belle, M. D., Gossan, N., Delagrange, P. & Piggins, H. D. Daily variation in the electrophysiological activity of mouse medial habenula neurons. J. Physiol. 592, 587–603 (2013).

18. Pennartz, C. M., Bierlaagh, M. A. & Geurtsen, A. M. Biometric modulation of pacemaker potentials and calcium current in the mammalian circadian clock. Nature 416, 286–290 (2002).

19. Lewis, A. H. & Raman, I. M. Persistent current at subthreshold voltages: activation by EPSP waveforms. J. Neurophysiol. 95, 1081–1093 (2012).

20. Yamada-Hanff, J. & Bean, B. P. Persistent sodium current drives conditional pacemaking in CA1 pyramidal neurons under muscarinic stimulation. J. Neurosci. 33, 15011–15021 (2013).

21. Do, M. T. & Bean, B. P. Sodium currents in subthalamic nucleus neurons from Nav1.6-null mice. J. Neurosci. 33, 726–733 (2013).

22. Do, M. T. & Bean, B. P. Subthreshold sodium currents and pacemaking of subthalamic neurons: modulation by slow inactivation. Neuron 39, 109–120 (2003).

23. Kononenko, N. I., Honma, S., Dudek, F. E. & Honma, K. On the role of calcium and potassium currents in circuit modulating of firing rate in suprachiasmatic nucleus neurons: multiteleode disk analysis. Neurosci. Res. 62, 51–57 (2008).

24. Beurel, E., Greco, S. F. & Jope, R. S. Glycogen synthase kinase-3 (GSK3): regulation, actions, and diseases. Pharmacol. Ther. 148, 114–131 (2015).

25. Flourakis, M. et al. A conserved bicycle model for circadian clock control of membrane excitability. Cell 162, 836–848 (2015).

26. Osoiro, N. et al. Persistent Nav1.6 current at axon initial segments tunes spike timing of cerebellar granule cells. J. Physiol. 588, 651–670 (2010).

27. Farajinia, S. et al. Evidence for neuronal desynchrony in the aged suprachiasmatic nucleus clock. J. Neurosci. 32, 5891–5899 (2012).

28. Nakamura, T. J. et al. Age-related decline in circadian output. J. Neurosci. 31, 10201–10205 (2011).

29. LeSauter, J. et al. A short half-life GFP mouse model for analysis of circadian clock organization. Brain Res. 964, 279–287 (2003).

30. Schaap, J. et al. Neurons of the rat suprachiasmatic nucleus show a circadian rhythm in membrane properties that is lost during prolonged whole-cell recording. Brain Res. 815, 154–166 (1999).

31. Meijer, J. H. & Michel, S. Neurophysiological analysis of the suprachiasmatic nucleus: a challenge at multiple levels. Methods Enzymol. 552, 75–102 (2016).

32. Bensing, R. C. et al. Neuropeptide Y-induced phase shifts of PER2: LUC rhythms are mediated by long-term suppression of neuronal excitability in a phase-specific manner. Chronobiol. Int. 29, 91–102 (2012).

33. Habita, L. M., Moltzof, H. E., Paul, J. R., Johnson, R. L. & Gamble, K. L. Suprachiasmatic nucleus function and circadian entrainment are modulated by G protein-coupled inwardly rectifying (GIRK) channels. J. Physiol. 592, 5079–5092 (2014).

34. Gamble, K. L., Kudo, T., Colwell, C. S. & McMahon, D. G. Gastrin-releasing peptide modulates fast delayed rectifier potassium current in peri-expressing SCN2a subtypes. J. Biol. Rhythms 26, 99–106 (2011).

35. Lucas, E. K. et al. PGC-1alpha provides a transcriptional framework for synchronous neurotransmitter release from paraventricular-positive interneurons. J. Neurosci. 34, 14375–14387 (2014).

36. Kim, J. K. et al. Modeling and validating chronic pharmacological manipulation of circadian rhythms. PTP Pharmacometics Syst. Pharmacol. 2, e57 (2013).

37. Kim, J. K. & Forger, D. B. A mechanism for robust circadian timekeeping via stoichiometric balance. Mol. Syst. Biol. 8, 630 (2012).
Author contributions
J.R.P., D.D., D.B.F., R.M.C. and K.L.G. conceived the experimental design; J.R.P., D.D. and L.J.M. performed experiments and compiled the results; J.R.P. and L.J.M. performed statistical analyses; and J.R.P., D.D., D.B.F., L.J.M., R.M.C. and K.L.G. wrote the paper.

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