BK channel inactivation gates daytime excitability in the circadian clock

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Inactivation is an intrinsic property of several voltage-dependent ion channels, closing the conduction pathway during membrane depolarization and dynamically regulating neuronal activity. BK K⁺ channels undergo N-type inactivation via their β2 subunit, but the physiological significance is not clear. Here, we report that inactivating BK currents predominate during the day in the suprachiasmatic nucleus, the brain’s intrinsic clock circuit, reducing steady-state current levels. At night inactivation is diminished, resulting in larger BK currents. Loss of β2 eliminates inactivation, abolishing the diurnal variation in both BK current magnitude and SCN firing, and disrupting behavioural rhythmicity. Selective restoration of inactivation via the β2 N-terminal ‘ball-and-chain’ domain rescues BK current levels and firing rate, unexpectedly contributing to the subthreshold membrane properties that shift SCN neurons into the daytime ‘upstate’. Our study reveals the clock employs inactivation gating as a biophysical switch to set the diurnal variation in suprachiasmatic nucleus excitability that underlies circadian rhythm.
Inactivation gating of ion channels is essential to electrical signalling in most neurons. First characterized in Hodgkin and Huxley's classic studies of the action potential, since then, several distinct inactivation mechanisms have been identified in Na⁺, Ca²⁺ and K⁺ channels. Among voltage-gated K⁺ channels, inactivation can occur via the classic 'ball-and-chain' mechanism involving pore occlusion by an intrinsic N-terminal 'ball' located on the cytosolic side of the α pore-forming subunit. In contrast, large conductance Ca²⁺- and voltage-activated BK K⁺ channels (α2,1,1, KCNMA1) are not intrinsically inactivating. However, inactivating BK currents have been observed in some central neurons, sensory neurons and neuroendocrine cells. In adrenal chromaffin cells, BK inactivation is mediated by the β2 subunit and shares several common features with Shaker N-type inactivation, including a rapid time course and contribution of intracellular domains sensitive to proteolytic cleavage. Chromaffin cells express both inactivating and non-inactivating BK channels, associated with distinct action potential behaviour. Yet despite the well-described mechanistic basis, understanding the physiological relevance for BK inactivation has been complicated by other properties conferred by the β2 subunit, including a shift in the voltage dependence of activation and slowing of activation and deactivation kinetics.

To understand the role of BK channel inactivation in neuronal excitability, we identified a circuit where β2 is expressed and where dynamic regulation of the BK current is critical for neural coding, the suprachiasmatic nucleus (SCN) of the hypothalamus. The SCN circuit undergoes synchronized daily oscillations in action potential firing frequency, and circadian behavioural and physiological characteristics are established by the parameters of the SCN circuit rhythm. BK channel expression varies over the circadian cycle in both mouse SCN and fly clock neurons. The daily rhythm in BK protein abundance is linked to Per2, a component of the canonical transcription–translation feedback loop that encodes circadian time. Correlated with expression levels, SCN neurons exhibit a diurnal difference in steady-state BK current magnitude, with smaller currents recorded during the day and larger currents at night. Both loss of BK current at night and aberrant increase in BK current during the day result in disrupted circuit rhythmicity, establishing the diurnal variation in BK current as essential for expressing behavioural rhythms. Loss of this diurnal rhythm in BK current has been recently linked to degradation of circadian rhythm in aged animals, underscoring the significance of understanding the mechanisms that drive rhythms in BK channel activity in SCN.

Here we report that the β2 subunit produces inactivation of SCN BK channels. Using patch-clamp, long-term action potential, and behavioural recordings, we show that loss of β2 (β2 KO) abolishes the diurnal variation in both BK current magnitude and SCN firing, and disrupts circadian circuit and locomotor rhythmicity. We link these changes in excitability specifically to BK channel inactivation by taking advantage of the modular nature of β2, which allows selective rescue of inactivation via delivery of the isolated β2 N-terminal ‘ball-and-chain’ domain. We further reveal that neurons with inactivating BK currents fire faster, have depolarized membrane potentials and increased input resistance, demonstrating that inactivation unexpectedly contributes to the daytime ‘upstate’ in SCN neurons by controlling the amount of subthreshold BK current during the inter-spike interval. Thus, inactivation underlies the shift in BK’s influence on excitability between day and night, and regulation of this biophysical mechanism is a central node for the circadian regulation of firing in the SCN clock circuit.

**Results**

β2 regulates SCN circuit rhythmicity and circadian behaviour. The SCN expresses two subunits with the potential to modify BK channel properties, β2 and β4 (ref. 22), but of these, only β2 can cause inactivation of BK currents. Since β subunits are proposed to tailor BK channel properties and contribute to distinct patterns of excitability across tissues, to establish the importance of the β2 subunit in the SCN circuit, we made long-term recordings of spontaneous action potential activity from wild-type (WT) and β2 KO SCNs. Individual SCN neurons function as coupled, autonomous oscillators, generating a robust synchronized rhythm in firing in *ex vivo* cultures (Fig. 1a) and β2 KO SCNs had significantly reduced rhythmicity overall (Fig. 1b). This was due to both a decrease in the number of rhythmic recordings, as well as a reduction in the circadian amplitude of the rhythmic recordings (Fig. 1c,d). The circadian period of the rhythmic activity was not different (WT, 24.2 ± 1.3 h and β2 KO, 23.4 ± 1.2 h, P = 0.8), instead, the reduced circadian amplitude stemmed from decreased action potential activity at the peak (Fig. 1e).

Alterations in the SCN circuit were correlated with disrupted circadian behaviour (Fig. 1f–g). Like SCN circuit activity, circadian behavioural amplitudes were reduced in β2 KO mice (Fig. 1h; Supplementary Table 1). Decreased circadian behavioural amplitudes are associated with a more labile pacemaker, and consistent with this, β2 KO mice re-entrained about a day faster to a 6-h phase advance of the light-dark cycle and had augmented responses to a phase-shifting light pulse (Fig. 1i,j; Supplementary Fig. 1; Supplementary Table 1). Taken together, these results suggest that β2 is important for the circadian patterning of action potential activity in the SCN circuit, and that this mechanism is required for proper circadian rhythmicity in the animal as a whole. Furthermore, the circadian alterations were specific to loss of β2, as deletion of the β4 subunit, which is also expressed in SCN, did not result in significant changes in neuronal firing or behavioural activity (Fig. 1c–e; Supplementary Table 1; Supplementary Discussion).

β2 mediates BK current inactivation in SCN neurons. To determine the functional relevance of β2 regulation of SCN circuit rhythmicity, we recorded BK currents from SCN neurons using whole-cell voltage clamp. We observed two types of macroscopic BK currents which varied by the amount of current decay observed during the voltage step (Fig. 2a). Currents that exhibited a pronounced decay within 30 ms were termed BK and had a fractional current ratio (I_{BK}/I_{BKpeak}) < 0.7 at +90 mV (Fig. 1b; Supplementary Fig. 2a,b). These BK currents decayed with time constants of 44 ± 4 ms (at +90 mV, n = 13) (Fig. 2c; Supplementary Fig. 2c,d), similar to inactivating BK currents described in other neuronal types. Other currents were relatively sustained with little decay, termed BK (I_{BK}/I_{BKpeak} > 0.7). The much slower time course of decay for BK currents (172 ± 13 ms, n = 17; Fig. 2c; Supplementary Fig. 2c) is consistent with non-inactivating BK currents that show a limited attenuation of current due to Ca²⁺ clearance. Overall, during the day, most currents were BK, while at night, most currents were BK (Fig. 2d). This suggested that the number of neurons exhibiting BK and BK currents between day and night SCNs was responsible for the diurnal difference observed in the steady-state BK current magnitude within SCN (Fig. 2e).

Supporting this idea, we did not find that daytime BK currents decayed more rapidly, and thus to a lower fractional current value, compared with night time BK currents. Instead, the average BK current density (Fig. 2f) and fractional current values (Supplementary Fig. 2b) were similar between day and night.
These observations support a model where the simple summation of BK\textsubscript{i} to BK\textsubscript{s} currents is a central factor in the day versus night BK current levels observed across the SCN.

Since the \( b_2 \) subunit produces inactivation of BK channels expressed in heterologous cells and native tissues \(^7,^8,^{18,34,39} \), we examined \( b_2 \) levels to determine whether expression of the \( b_2 \) subunit was correlated with the increased number of BKi currents observed in daytime SCNs. BK complexes were immunoprecipitated from SCNs harvested across the circadian cycle. Although the expression of \( b_2 \) did not differ across time points (Fig. 3a, b), the relative ratio of \( b_2 \) associated with the BK \( \alpha \) subunit was higher during the day (Fig. 3c), correlated with the higher proportion of BK\textsubscript{i} currents (Fig. 2d). At night as \( \alpha \) expression increased, the ratio of \( b_2:\alpha \) was lower and consistent with the increased number of BK\textsubscript{s} currents.

To determine whether \( b_2 \) was required for the macroscopic BK\textsubscript{s} current decay in SCN, we recorded BK currents from \( b_2 \) KO neurons. Only BK\textsubscript{s} currents were observed in \( b_2 \) KO neurons (Fig. 4a, b; Supplementary Fig. 2a), implicating inactivation as the mechanism for the decay of BK\textsubscript{s} currents. As a consequence of converting all currents to BK\textsubscript{s} during the day, the diurnal difference in BK current magnitude was abolished in \( b_2 \) KO neurons.
SCNs (Fig. 4c). Although the small number of BK currents observed in WT neurons at night were also eliminated, the overall night-time β2 KO BK current levels remained unchanged. This was expected since the vast majority of currents were already identified as BK, in WT neurons at night. Further consistent with loss of inactivation, the fractional current values and macroscopic current decay time constants were increased in β2 KO during the day, with no change at night (Fig. 4d,e; Supplementary Fig. 2a–c). These results demonstrate that β2 underlies the reduction in steady-state BK current levels during the day and is thereby the central regulator of the diurnal difference in SCN BK current. As a control, we also assayed the expression level and diurnal pattern of the BK channel α2 subunit7,11,40,41. If inactivation is required for the diurnal difference in BK current (Supplementary Fig. 6). The restoration of daytime BK current levels comparable to WT neurons (Fig. 6b), without affecting the voltage dependence of BK current activation (Supplementary Fig. 5a–d). Furthermore, β2N decreased β2 KO BK current levels back to levels comparable to WT neurons (Fig. 6b), without affecting the voltage dependence of BK current activation (Supplementary Fig. 5) or other voltage-activated outward currents (Supplementary Fig. 6). The restoration of daytime BK current levels by β2N support the conclusion that inactivation is the critical function of the β2 subunit that underlies its role in driving the diurnal variation in BK current properties in SCN.

Inactivation controls daytime membrane properties and firing. If inactivation is required for the diurnal difference in BK currents, then loss of the β2 subunit or selective rescue of inactivation with β2N should alter neuronal activity. To test this, we recorded spontaneous action potentials from WT and β2 KO SCN neurons. WT neurons fire at higher frequencies during the day and decrease firing at night (Fig. 7a,b). Neurons with BK and BKs currents were isolated by subtraction with the antagonist paxilline (Methods). The β2 subunit or selective rescue of inactivation with β2N had no effect (β2NAFW). The voltage dependence of inactivation with β2N was similar to the intact β2 subunit (Fig. 5e), and τ inact values at β2N concentrations above 50 μM were similar to WT BK currents recorded during the day from SCN neurons (Fig. 5d). We next applied the β2N peptide to SCN neurons via the patch pipette.

In β2 KO SCN neurons, application of β2N, but not β2NAFW, restored the macroscopic decay and reduced the fractional current (Fig. 6a; Supplementary Fig. 2a). The voltage dependence of inactivation and τ inact values with β2N in β2 KO were similar to WT BK currents recorded during the day (β2 KO + β2N at +90 mV; 40 ± 4 ms, n = 20, Fig. 6a; Supplementary Fig. 2c,d). Furthermore, β2N decreased β2 KO BK current levels compared to WT neurons (Fig. 6b), without affecting the voltage dependence of BK current activation (Supplementary Fig. 5) or other voltage-activated outward currents (Supplementary Fig. 6). The restoration of daytime BK current levels by β2N support the conclusion that inactivation is the critical function of the β2 subunit that underlies its role in driving the diurnal variation in BK current properties in SCN.

HEK293 cells (Fig. 5). β2N produced a dose-dependent inactivation, while a peptide harbouring the FIW motif produced no effect (β2NAFW; Fig. 5a–d). The voltage dependence of inactivation with β2N was similar to the intact β2 subunit (Fig. 5e), and τ inact values at β2N concentrations above 50 μM were similar to WT BK currents recorded during the day from SCN neurons (Fig. 5d). We next applied the β2N peptide to SCN neurons via the patch pipette.
diurnal difference in firing across the SCN (Fig. 7b), suggesting BK inactivation may be fundamental to the diurnal regulation of firing rate. Consistent with this, β2 KO neurons did not exhibit a day–night difference in firing rate, stemming from a marked reduction in daytime firing (Fig. 7b,c). To verify that the reduced firing, which was correlated with larger BK currents in β2 KO neurons (Fig. 6b), was due to BK, the BK channel blocker paxilline was applied. The reduced firing in β2 KO neurons was reversed by paxilline (Fig. 7b), demonstrating that BK channels were responsible for the altered excitability in β2 KO neurons.

These results corroborate the reduced peak firing observed in SCN circuit recordings (Fig. 1e). In addition, these data may also explain why loss of BK currents during the day has apparently little effect on WT firing rates. A detectable change in firing after BK block in WT neurons may be precluded under standard daytime conditions when two-thirds of SCN neurons undergo BK inactivation. Following this idea, application of β2N, but not β2NAFIW, increased daytime β2 KO firing rates back to WT levels (Fig. 7b,d). β2N had no further consequences on firing beyond that of blocking the BK current with paxilline (Fig. 7b), demonstrating that the restoration of firing with β2N acts selectively through BK channels. These results suggest that inactivation normally removes BK currents from influencing action potential firing rate during the day. Illustrating this in another way, β2N could also aberrantly suppress BK current (Fig. 8a–c) and elevate firing if applied to WT neurons at night (Fig. 8d), underscoring the basis for the normal reduction in the number of BK currents found in the SCN at night.

What membrane properties does BK inactivation regulate to facilitate daytime firing? A central node for circadian regulation of action potential frequency is the balance of subthreshold excitatory and inhibitory currents that bring the resting membrane potential closer to the firing threshold during the day, or further from threshold at night. K⁺ currents are proposed to be central regulators of the day–night difference in firing by regulating the daily transition between the daytime...

**Figure 3 | Expression of BK α and β2 in WT SCNs.** (a) BK channel complexes were immunoprecipitated with an anti-BK α subunit antibody from WT SCNs at the indicated time points. Western blot analysis was performed for BK α (top panel), β2 (middle) or α-tubulin (bottom). α-Tubulin westerns (loading control) were obtained by running an equivalent volume of supernatant as was used for the immunoprecipitation. Protein was also harvested from α + β2 subunits co-expressed in HEK293 cells (positive control) or from β2 KO SCNs (negative control). ZT, zeitgeber time. Images have been cropped for presentation. Full size images are presented in Supplementary Fig. 3. (b) BK α and β2 band intensities normalized to α-tubulin. α expression increases at night, while β2 expression does not change. Data are the average from four independent timed SCN collections (four SCNs at each timepoint). (c) Ratio of β2:α expression in WT SCNs across the circadian cycle. All values are mean ± s.e.m. *P < 0.05, Bonferroni post hoc.

**Figure 4 | The β2 subunit is required for BK current decay and the diurnal difference in BK current magnitude in SCN neurons.** (a) Representative BK current traces and macroscopic decay time constants (t_inact) for BKs currents from β2 KO neurons. Voltage protocol same as in Figure 2. (b) All currents are BKs from β2 KO SCNs, day or night. (c) β2 KO neurons do not show a day–night difference in BK current density. The β2 KO daytime BK current magnitude is larger than WT, comparable to WT levels at night. (d) The fractional BK current (I_BK/peak) was similar during the day compared with night in β2 KO neurons. (e) t_inact values from β2 KO BKs currents are all >100 ms. All values are mean ± s.e.m. n values: β2 KO, day (20) and night (20).
expressed the macroscopic current decay with the macroscopic current traces from HEK293 cells expressing BKSRKR channels, a daytime BK variant previously cloned from SCN (ref. 60). Figure 5 | The \( \beta 2 N \) terminus (\( \beta 2 N \)) causes inactivation of BK currents from \( \alpha \)-only channels expressed in heterologous cells. (a) Representative macroscopic current traces from HEK293 cells expressing BK\text{SRKR} channels, a daytime BK variant previously cloned from SCN (ref. 60). \( \beta 2 N \) (0–100 μM), corresponding to the first 45 amino acids of the N terminus, was dissolved in recording solution and applied to the intracellular side via the patch pipette. (b) Representative currents in the presence of 50 μM \( \beta 2 N^{\text{FIW}} \), a peptide mutating the three residues required for inactivation\(^{42} \). There was no macroscopic current decay with \( \beta 2 N^{\text{AFW}} \). (c) Representative traces from the +90 mV voltage step at each concentration of \( \beta 2 N \). The current peaks were scaled to illustrate the dose-dependent speeding of inactivation. (d) \( t_{\text{inact}} \) values were calculated from the current elicited at +90 mV and plotted as a function of \( \beta 2 N \) concentration. \( t_{\text{inact}} \) was plotted for BK, and \( \beta 2 KO \) SCN BK currents for cross-comparison. (e) \( t_{\text{inact}} \) versus voltage for BK channels co-expressed the \( \beta 2 \) subunit, or with 50 μM \( \beta 2 N \). There is no difference in the voltage dependence of activation using the isolated \( \beta 2 N \) compared with the intact \( \beta 2 \) subunit. All values are mean ± s.e.m. For currents from HEK293 patches, \( n = 4 \) at each concentration of \( \beta 2 N \) or \( \beta 2 N^{\text{FIW}} \) peptide. For currents from daytime SCN neurons, \( BK_i \) (\( n = 18 \)) and \( \beta 2 KO + \beta 2 N \) (\( n = 10 \)).

**Figure 6 |** \( \beta 2 N \) rescues inactivation and restores daytime BK current levels in \( \beta 2 \) KO SCN neurons. (a) Representative traces showing rescue of the macroscopic BK current decay in \( \beta 2 \) KO neurons with 50 μM \( \beta 2 N \) applied intracellularly. 50 μM \( \beta 2 N^{\text{AFW}} \) did not rescue the current decay. Voltage protocol same as in Fig. 2. (b) Application of \( \beta 2 N \) reduced the BK current density in \( \beta 2 \) KO neurons to levels comparable to WT. \( \beta 2 N^{\text{AFW}} \) had no effect on BK current levels. WT and \( \beta 2 \) KO I-Vs re-plotted from Figs 2 and 4 for cross-comparison. \( n \) values: WT (27); \( \beta 2 \) KO (20); \( \beta 2 \) KO + \( \beta 2 N^{\text{AFW}} \) (18); \( \beta 2 \) KO + \( \beta 2 N \) (20). All values are mean ± s.e.m. *\( P < 0.05 \), Bonferroni post hoc.

‘upstate’ and night time ‘downstate’\(^{44–47} \). These states are designated by differences in the subthreshold membrane properties, with the most attention focused on the increase in night time \( K^+ \) current associated with hyperpolarization of the membrane and lower input resistance\(^{44,45,47,48} \). BK channels comprise a component of this upregulated \( K^+ \) current, and loss or block of BK currents causes depolarization of the baseline membrane potential and hyperactive firing at night\(^{23–25} \). However, block of BK currents also depolarizes the membrane potential during the day\(^{43,47,49} \), suggesting a role for BK channels in daytime excitability not previously explored. Thus to determine whether BK inactivation makes a converse contribution to the daytime reduction in \( K^+ \) current that defines the ‘upstate’ that is required for higher frequency firing, we compared membrane parameters from neurons during the day and night, and with loss of \( \beta 2 \) or restoration of inactivation.

First, we found that \( BK_i \) neurons had inter-spike membrane potentials ranging from −36 to −51 mV, while \( BK_i \) neurons ranged from −47 to −54 mV during the day. Loss of inactivation in \( \beta 2 \) KO neurons resulted in hyperpolarization of the inter-spike membrane potential compared with WT \( BK_i \) neurons (−57 ± 1.4 mV versus −50 ± 1.1 mV, respectively; \( P < 0.001 \), \( t \)-test). This shift could be rescued by \( \beta 2 N \) (−52 ± 0.8 mV), but not \( \beta 2 N^{\text{AFW}} \) (−56 ± 1.5 mV; one-way ANOVA, \( P = 0.04 \); Bonferroni post hoc \( P < 0.05 \) for \( BK \), versus \( \beta 2 \) KO or \( \beta 2 N^{\text{AFW}} \)), suggesting that BK channel inactivation affects membrane potential during the inter-spike interval. To further address this, resting membrane potential (\( V_m \)) and input
Resistance ($R_i$) recordings were made in the presence of tetrodotoxin to block action potential activity. Daytime neurons had more positive $V_m$ and higher $R_i$, while night time neurons had more hyperpolarized $V_m$ and lower $R_i$ (Fig. 9a,b), consistent with previous studies. The average daytime membrane potential ($-48 \pm 0.8$ mV) was dominated by the increased number of BK$_i$ neurons, which had more positive $V_m$ ($-47 \pm 0.4$ mV) and higher $R_i$ ($1.4 \pm 0.06 \text{ G} \Omega$) than BK$_s$ neurons.

**Figure 7** | Loss of β2 eliminates the diurnal difference in firing rate, and rescue of inactivation with β2N restores daytime firing rates in SCN neurons. (a) Spontaneous action potential activity from representative day (BK$_i$) and night (BK$_s$) WT neurons. Dotted line (a,c,d) denotes $-50$ mV. (b) In WT SCNs, BK$_i$ neurons fired at higher frequencies than BK$_s$, similar to the average day-night difference in firing. β2 KO neurons did not exhibit a diurnal difference in frequency, and during the day, fired at levels similar to WT night. Application of β2N to daytime β2 KO neurons had no effect on frequency, but β2N increased firing rate to WT levels. (c) Day (BK$_i$) and night (BK$_s$) β2 KO neurons. (d) Day β2 KO neurons with 50 μM β2N (BK$_i$) or 50 μM β2N to (BK$_s$). All values are mean ± s.e.m. n values: WT: BK$_i$ (17), BK$_s$ (10), day (17), night (20); β2 KO: day (19), night (19), β2N (20), β2N/pax (8), and pax (8).

**Figure 8** | β2N can confer inactivation to WT neurons at night. (a) Representative macroscopic traces from WT neurons at night showing a typical BK$_s$ current, and a BK$_i$ current resulting from application of 50 μM β2N. Voltage protocol same as in Fig. 2. (b) β2N reduced the night time current density in WT neurons to daytime levels. WT day and night data re-plotted from Figs 2 and 4 for cross-comparison. (c) The proportion of BK$_i$ currents increased significantly with β2N ($P = 0.0001$, Fisher’s exact test). (d) β2N increased night time firing to daytime levels. WT day and night data re-plotted from Fig. 7 for cross-comparison. All values are mean ± s.e.m. n Values: WT, day (27); WT, night (22); and WT + β2N, night (20). *P<0.05, Bonferroni post hoc.
Overall, BK currents were larger in BKi compared with BKs neurons. This suggests less current would be required to elicit an action potential in BKi neurons. We found that the ratio of Vm and Ri values in BKβ2KO neurons back to values comparable to WT, establishing the sufficiency for inactivation of the BK current to convert the daytime membranes to pro-excitable. Although β2 can also affect excitability by shifting the voltage dependence of activation and slowing activation and deactivation gating (Supplementary Discussion), the data presented here affirm that these functions are not directly involved in the diurnal regulation of membrane properties that control firing frequency in SCN.

The effect of inactivation on resting membrane potential and input resistance predicts that inactivation of BK currents would occur during the inter-spike interval. To demonstrate this, we recorded BK currents from traditional voltage step commands across the subthreshold range of membrane potentials, as well as BK currents evoked by SCN action potential commands (Fig. 10; Supplementary Fig. 7). BK current–voltage relationships showed that daytime BK current activated at voltages more positive than −60 mV (Fig. 10a,b). There was 5–12 pA of current evoked in the subthreshold range, between the average resting and threshold potentials of SCN neurons (−55 to −40 mV). Night time subthreshold BK currents (BKb) were also larger than the net daytime BK current (Supplementary Fig. 7a). In contrast, daytime BKβ2KO neurons had significantly reduced current in the subthreshold voltage range (Fig. 10a,b). Consistent with data obtained from voltage steps, using daytime SCN action potential commands delivered at the normal daytime firing frequency, the amount of BK current inactivated during the inter-spike interval was negligible in BKβ2KO neurons compared with BKb neurons (Fig. 10c,d) or at night (Supplementary Fig. 7b,c). BK currents evoked at the peak of the action potential command were also reduced in BKβ2KO neurons compared with BKb neurons during the day (Fig. 10e) or at night (Supplementary Fig. 7d). In parallel experiments using daytime action potential commands applied to channel complexes expressed in HEK293 cells (Supplementary Fig. 8a–j), the subthreshold current was reduced in patches containing BKβ2 or BKβ2N compared with α-only channels (Supplementary Fig. 8f–i).

To further demonstrate that inactivation could affect BK current levels at these subthreshold voltages in SCNs, BKβ2KO neurons were held at a conditioning voltage for 100 ms, corresponding to the shortest inter-spike intervals encountered in spontaneously firing SCN neurons, followed by a step to +90 mV to elicit a maximal BK current response (Fig. 10f). Inactivation delivered via BKβ2N reduced the maximal BK current elicited following conditioning voltages in the subthreshold range, compared with BKβ2KO cells treated with BKβ2N/β2FIW (Fig. 10f,g). Further depolarization of the conditioning potential resulted in progressive current reduction that was dependent on the presence of inactivation (Fig. 10f,g). Similar results were obtained with BK channels expressed in heterologous cells with the intact β2 subunit or with BKβ2N (Supplementary Fig. 8a,b). These results suggest that inactivation can reduce BK current under conditions similar to the native inter-spike interval and provide a basis for understanding the effect of BK inactivation on the resting membrane potential and input resistance.

The data further propose that because some BK current would be inactivated before an action potential, the current evoked by an action potential would be reduced. This would be expected to occur during the day in BKβ2KO neurons. To test this, we quantified the peak BK current evoked by daytime action potentials applied to SCN neurons (Fig. 10c,e) and to BKβ2/BKβ2FIW channels expressed in heterologous cells (Supplementary Fig. 8e–h,i). The action potential-evoked BK current was reduced in BKβ2KO neurons (Fig. 10e) and in BKβ2 or BKβ2FIW channels in HEK293 cells compared with α-only channels (Supplementary Fig. 8j). Taken together, these data support a model where current inactivation is necessary during the day to regulate the membrane parameters that initiate firing (membrane potential and input resistance), while concurrently reducing the BK current activated during an action potential. This idea further predicts there would be little difference in the action potential waveforms of BKβ2KO versus BKβ2KO neurons, an observation confirmed by analysis of daytime action potential parameters (Supplementary Fig. 9).
β2-mediated inactivation thus comprises an essential switch that shifts membrane properties between day and night states by inhibiting or permitting subthreshold BK current activation.

Discussion
The circadian patterning of neuronal excitability is an evolutionarily conserved feature of time-encoding circuits. In SCN neurons, which fire relatively slow, spontaneous action potentials characteristic of pacemaker neurons, the balance of currents during the inter-spike interval is proposed to be the predominant regulator of firing rate and a major node for circadian regulation. Correlated with the increased number of BK currents in SCN during the day, most neurons have a more depolarized resting membrane potential that is closer to threshold (ΔV ≈ 7 mV, from our data), increasing the likelihood of firing an action potential. At night, and correlated with the shift to BK currents, the hyperpolarizing shift places the membrane potential further from threshold (ΔV ≈ 12 mV), reducing excitability. The amount of current required to produce this shift is small (~5 pA), owing to the high input resistance of SCN neurons, and the balance can be tipped either by altering the

**Figure 10 | Steady-state and action potential-evoked BK currents are reduced compared to BKs in SCN neurons.**

(a) Representative macroscopic BK current traces from daytime WT neurons in response to a subthreshold voltage protocol (−60 to −30 mV in 5 mV steps). BK currents were identified as BK or BKs from a maximally activating step to +90 mV (as in Fig. 2). (b) Current–voltage relationship showing significantly more activation of BKs current above −60 mV compared to BKs. (c) Pre-recorded action potential commands (top) were used to elicit BK currents from BK or BKs neurons. From a holding potential of −150 mV to remove inactivation completely, cells were stepped to the inter-spike potential (−48 mV) with a sequence of three action potential commands as depicted (Peak, 8 mV; t1/2, 5.5 ms; and AHP/antipeak, −54 mV). Arrow, the subthreshold current level in d was taken just before the second action potential command. (d,e) Average subthreshold BK current density from the inter-spike interval (d) or at the peak of the action potential (e). n values for (a–d): WT: day/BKs (9), day/BKi (5). (f) BK current as a function of conditioning potential in SCN neurons. Representative macroscopic BK current traces from β2 KO neurons during the day with 50 μM β2NΔFIW or β2N. Neurons were held at −90 mV for 150 ms to remove inactivation, and then stepped to a conditioning potential for 100 ms to allow channels to transition into the inactivated state, followed by a maximally activating step to +90 mV. (g) The peak BK current elicited from the +90 mV step was plotted as a function of the conditioning potential. β2N causes a reduction in current, but no reduction is observed in the absence of inactivation with β2NΔFIW. n = 5 for each condition. All values are mean ± s.e.m. *p<0.05, Bonferroni post hoc.
excitatory currents that bring the membrane potential closer threshold, or the inhibitory currents that take the membrane away from threshold. Recently, the identity of the channel underlying the persistent Na\(^+\) current that provides the depolarizing drive on excitability was identified as the leak Na\(^+\) channel NALCN\(^{51}\). However, the identity of the K\(^+\) channels counterbalancing the excitatory current has been uncertain, particularly during the day when SCN neurons undergo a dramatic reduction in K\(^+\) current\(^{44,45,49}\). Although abrogation of one type of K\(^+\) current during the day, the fast delayed rectifier, has also been shown to decrease firing frequency, the reduced firing stems from slower action potential repolarization and not from depolarization of the resting membrane potential\(^{52}\). Consequently, a direct test of whether the daily shift in submembrane K\(^+\) currents is required to drive the day–night difference in action potential firing has been elusive.

In this study we provide surprising evidence linking alterations in BK current with the daytime increase in input resistance, depolarization of the membrane and higher frequency firing. BK currents were previously thought to mainly play a night-time-restricted role as part of the surge of K\(^+\) current that results in the firing nadir\(^{24,44,45}\). The less significant role for BK channels in setting daytime frequency was attributed to a decrease in expression of the pore-forming \(\alpha\) subunit. However, in the \(\beta2\) KO, the intact circadian pattern of \(\alpha\) expression was insufficient to support normal diurnal variation in BK current or SCN firing, suggesting there must be other mechanisms that limit BK channels from influencing daytime membrane properties. We demonstrate that inactivation, achieved through \(\beta2\)-dependent regulation of BK channel biophysical properties between day and night, is the central mechanism that changes the influence of BK channels on membrane properties between day and night. Inactivation facilitates the daytime membrane ‘upstate’ by reducing the BK current between spikes that would normally inhibit action potential initiation. Inactivation also reduces action potential-evoked current, and the sum of these two mechanisms effectively removes BK channels from influencing daytime excitability. In \(\beta2\) KO SCN neurons, the absence of BK inactivation during the day results in passive membrane properties that look much like night time membranes. Correspondingly, with \(\beta2\)N, restoration of inactivation is both necessary and sufficient to recapitulate the \(V_\text{m}\) and \(R_\text{p}\) properties that define the daytime membrane state. Importantly, the identification of this biophysical mechanism to functionally reduce a K\(^+\) current suggests that the balance of excitatory and inhibitory currents is more actively regulated during the day than previously thought\(^{45}\). Moreover, the disruption of circadian behaviour in the absence of the \(\beta2\) subunit underscores the importance of both BK inactivation, and the diurnal variation in SCN submembrane properties, in the regulation of circadian rhythmicity. As such, BK\(_i\) currents comprise an electrophysiological signature for a previously undetected class of essential oscillator in SCN that makes a vital contribution to circadian behaviour. Even so, the residual rhythmicity in \(\beta2\) KOs hints that neuronal heterogeneity contributes redundant mechanisms to circadian behavioural rhythmicity (Supplementary Discussion), and it remains to be determined how SCN neurons without inactivating BK channels balance their daytime currents to control firing, and what separable contribution these cells make to circadian behaviour.

Given that the BK current must be functionally reduced by inactivation, what is the relevance of the daytime clock-linked decrease in BK \(\alpha\) subunit expression\(^{24,30}\)? Such differences in ion channel expression across the circadian cycle are broadly proposed to be central to rhythmicity\(^{26}\), but an explicit mechanism that translates channel expression differences into action potential activity has not been established. In this study, we showed \(\beta2\) KO SCNs retained the normal day–night difference in BK \(\alpha\) expression, yet still displayed disrupted BK steady-state current levels and action potential firing, demonstrating that circadian differences in the BK \(\alpha\) subunit alone are insufficient to drive rhythmicity. Our data instead suggest a model in which the clock-linked changes in \(\alpha\) expression control the ratio of \(\beta2\alpha\). Since expression of \(\beta2\) does not vary appreciably between day and night, the higher daytime \(\beta2\alpha\) ratio facilitates inactivation in the majority of SCN neurons, permitting transition to the ‘upstate’. At night, inactivation is overcome by increased \(\alpha\) subunit expression, reducing the \(\beta2\alpha\) ratio and facilitating the ‘downstate’. In support of this hypothesis, aberrantly increasing daytime BK expression reduces inactivation in the SCN (Supplementary Discussion).

The data in SCN also provide a framework for understanding the physiological relevance of BK channel inactivation, which is too slow to produce substantial BK current decay during a typical central nervous system action potential\(^{53}\). In other neurons where higher frequency firing prevails, BK inactivation is thought to contribute to spike frequency adaptation, which also integrates over timeframes longer than a single action potential\(^{10,15,57}\). In SCN neurons, which exhibit depolarized membrane potentials and slow spontaneous firing, the results here suggest that enough channels undergo inactivation, or remain cumulatively inactivated, during the inter-spike interval (100–500 ms) to impact the input resistance and resting membrane potential. Given \(t_\text{inact}\) values of 35–100 ms in BK\(_i\) neurons, steady-state inactivation may be most relevant for setting the subthreshold BK current level during the inter-spike interval. SCN neurons have been shown to undergo subthreshold Ca\(^{2+}\) oscillations\(^{49,54}\), raising the possibility that Ca\(^{2+}\) influx independent of the action potential could contribute to BK channel activation, and consequently BK inactivation, during the inter-spike interval. In addition, the restoration of firing rate with \(\beta2\)N reveals that inactivation mediated by the N terminus is sufficient for this role. N-type inactivation typically occurs from the open state\(^7,8\), but BK channels may also inactivate from closed states, affecting the amount of steady-state current\(^55\). Thus, the data in this study reveal a surprising twist to how cells use BK channels to regulate excitability. While BK channels perform diverse roles spanning excitable and non-excitable cells, in the CNS they are mostly known for regulating firing frequency by underlying the fast AHP\(^56–58\). Here we show in SCN, BK current activated outside of the action potential is potentially the major contributor to frequency regulation, making dynamic regulation of subthreshold BK current via inactivation a novel role for BK channels.

In summary, although the combination of BK\(_i\) and BK\(_i\) expressing cells has been observed within other tissues\(^{18}\), the significance for how these currents work together to set the ensemble properties underpinning physiological function is not yet apparent\(^{59}\). In SCN, the cycling between predominantly BK\(_i\) currents during the day and predominantly BK\(_i\) currents at night suggests that partnering with the \(\beta2\) subunit generates a hardwired mechanism linked to the intrinsic clock to dynamically disengage BK channels from regulating action potentials during one phase of the circadian cycle.

**Methods**

**Mice.** WT, \(\beta2\) KO (KCNMB2\(^{-/-}\)), gift from C. Lingle), \(\beta4\) KO (KCNMB4\(^{-/-}\), gift from R. Brenner), and Kenmali\(^{-/-}\) mice were maintained on inbred C57BL/6 background and genotyped as previously described\(^{18,24,59}\). KCNMB2 floxed mice are available at The Jackson Laboratory (Stock no. 028416). Male and female mice were used (3 weeks to 5 months old) and were group housed separated by sex until experimental procedures as indicated. Day time points were collected from mice housed on a standard 12 h light, 12 h dark cycle (12:12 LD), and night time points...
were collected from mice housed on a reverse 12:12 LD cycle. All procedures involving mice were conducted in accordance with the University of Maryland School of Medicine Animal Care and Use Guidelines and approved by the IACUC Committee.

Organotypic slice culture. Brains were harvested from postnatal day 4 mice, and the hypothalamus was blocked and sectioned into 300 μm coronal slices using a manual chopper (Stoelting, Wood Dale, IL, USA). Slices containing SCN nuclei were transferred onto Millicell filters (Millipore, Billerica, MA, USA) and cultured as interface explants in a CO₂ incubator as described. After 2 days, 20 μg/ml cytosine β-arabinofuranoside (ara-C, no. C6645; Sigma) was added to inhibit glial proliferation, and the media was changed every 3 days thereafter.

Multielectrode array recordings. After 7 days in culture, organotypic slices were cut from the surrounding membrane, inverted over the 64-electrode grid, and adhered to 0.1% polyethyleneimine and collagen-treated P210A probes (Alpha MED Scientific, Osaka, Japan). Probes were placed on a MED CO2 connector headstage, sealed with a vacuum-greased coverslip, and maintained in a humidified 5% CO₂ incubator at 37 °C. Membrane potentials were amplified (VT1000S vibratome (Leica Microsystems, Wetzlar, Germany) at 3–4 °C). Signals from all electrodes on the probe were collected simultaneously with the 64-channel integrated amplifier (Alpha MED Scientific and ActiLife (Alpha MED Scientific) and analysed as described previously. Second-order Butterworth filters were used with a low frequency cut-off of 0.1 Hz. Spike and extracellular action potentials from visually identified electrodes within the SCN were discriminated offline using threshold-based event counting. Circuit analysis was performed on each slice from three cycles of activity at all electrodes within the SCN. The multiunit spontaneous action potential activity from each electrode located in the SCN area was determined as the highest peak above the 99% confidence interval of a periodogram. The daily firing rate peak, and the ‘day’ firing rate value was the average of activity for the duration of the recordings. Media changes were maintained every 3 days.

For locomotor rhythms, mice (3–5 months old) were housed individually in cages containing a running wheel for 10 days in LD 16:8 in constant darkness (DD). Activity was sampled every 10 s in ClockLab software (Actimetrics). Actograms were constructed by double-plotting consecutive days of activity over the recording period. Circadian period and amplitude were determined from 10 days of wheel running activity in DD in Clocklab software as described previously. For re-entrainment experiments, after 7 days of stable entrainment, the LD cycle was phase advanced by 6 h. The response time was calculated as the number of days to stable re-entrainment. After keeping mice in DD for 8 days, phase shifts in response to light pulses were calculated as the number of hours between the activity onset regression fits before and after a 30-min light pulse delivered in early subjective night (CT16). Alpha was determined as the length of time an animal had consolidated activity using default settings with manual adjustment, with rho defined as the portion of the cycle outside of alpha. Bouts were calculated using default settings. Data was excluded from mice that failed to run on wheels for two consecutive days.

Western blots. A single block of hypothalamus containing the SCN (∼2 mm) was harvested at the indicated time points from mice (2- to 3-months old). Four micrograms solubilized protein from individual SCNs were loaded per lane. Densitometry of BK doublet band (1 μg/ml L6/60 mice monoclonal α-BK antibody, Neurona; University of California, Davis, CA, USA) to DMI2 against α-tubulin (1:75,000; Sigma T-9026) was performed as described previously and presented as a proportion of ZT20. Three independent circadian cycle tissue harvests were performed for each condition. For immunoprecipitation, individual SCNs (∼50 μg of protein) were pre-cleared by incubation with 20 μl TrueBlot anti-rabbit lg IP beads (Rockland Immunomolecular, Limerick, PA, USA) at 2 °C for 4 h (ref. 18). Samples were centrifuged at 13,200 r.p.m. for 30 min to precipitate the beads. Supernatant was collected and mixed with 1 μg of anti-BK α subunit antibody (anti-KCa1.1, #1184–1200, Alamone Labs; Jerusalem, Israel) and rotated for 2 h at 4 °C. 50 μl TrueBlot anti-rabbit lg IP beads were added, and the mixture was rotated overnight at 4 °C, centrifuged at 13,200 r.p.m. for 30 min to precipitate the beads, and the supernatant was discarded. Beads were washed three times and resuspended in 20 μl lysis buffer. Protein was run on a western blot as detailed previously and probed with 1 μg/ml α anti-BK (L6/60; Antibodies Incorporated; Davis, CA, USA); 1:75,000 anti-α-Tubulin (DM1A Sigma; St Louis, MO, USA) or 1 μg/ml α anti-f2 (N53/32; Antibodies Incorporated; Davis, CA, USA). Full size images for western blots are presented in Supplementary Figs 3 and 4.

Acute SCN slice preparation. Mice were sacrificed in the light, at zeitgeber time (ZT) ZTD–2 (day time points) or ZT11–12 (night) from 3- to 6-week-old mice. Brains were immediately removed and placed into cold saline. Brains were cut at the following (in mM): 1.2 MgSO₄, 26 NaHCO₃, 1.25 Na₂HPO₄, 3.5 KCl, 3.8 MgCl₂, 10 glucose and 200 sucrose. Coronal slices were cut at 300 μm on a VT1000S vibratome (Leica Microsystems, Wetzlar, Germany) at 3–4 °C. Slices containing SCN were recovered between 3–4 h (day) and 10–11 h (night) after ZT11. Slices were superfused in oxygenated artificial cerebrospinal fluid (in mM: 125 NaCl, 1.2 MgSO₄, 26 NaHCO₃, 1.25 Na₂HPO₄, 3.5 KCl, 2.5 CaCl₂, and 10 glucose).

Slices containing SCN were transferred to the recording chamber (RC26GLP/PM-1; Warner Instruments, Hamden, CT, USA) with gravity flow bath perfused with artificial cerebrospinal fluid (ACSF). Neurons were visualized with a Luca-R DL-600 EMCCD camera (Andor, Belfast, UK) under IR-dic illumination on an FN1 upright microscope (Nikon, Melville, NY, USA). Recordings were made from cells in the centre of the SCN. Current- and voltage-clamp recordings were made with a Multiclamp 2000B or 700B amplifier and pClAMP v10 software (Molecular Devices, Sunnyvale, CA, USA). Data were acquired at a 20 or 100 kHz sampling rate. Drugs were delivered to the bath by a computer-controlled pressurized perfusion system (ValveLink 8.2; Automatic Scientific, Berkeley, CA, USA) at the concentrations indicated for 1,000 × stocks. Recording windows were at the peak (ZT4–8) and nadir (ZT17–21) of the circadian rhythm in spontaneous action potential firing, corresponding to the ‘day’ and ‘night’ time points, respectively. Although the action potential frequencies in this study were lower than our previous studies performed at around 35 °C (refs 22,23), the day–night difference in firing frequency in this study is preserved (Fig. 4ac). All recordings were made with synaptic transmission intact, unless indicated (Tetrodotoxin was applied for all Vh and R, measurements, as well as the Western blot). BK currents in Fig. 5). Reported N’s are the number of neurons, and data for each condition was derived from 5 to 15 slices (1–6 neuronal recordings per slice).

SCN electrophysiology. SCN neurons were recorded in whole-cell configuration, first in current-clamp mode to collect action potential data, then switched into voltage clamp mode to obtain macroscopic voltage-activated outward K⁺ currents. Electrodess (4–7 MD) were filled with internal solution, in mM: 123 K-methanesulfonate, 9 NaCl, 0.9 EGTA, 9 HEPES, 14 Tris-phosphocreatine, 2 Mg-ATP, 0.3 Tris-GTP, and 2 Na₂-ATP, pH adjusted to 7.3 with KOH. The electrode solution was adjusted to 303 mosmol with sucrose and bath osmolarity was 300 mosmol. This internal solution supports BK current activation and spontaneous firingβ25,38,49. After GΩ seal and whole-cell break-in, membrane properties were elicited from a +20 mV step from a holding potential (Vh) of −90 mV. Access resistance was verified to be ≤25 MΩ with ±5% change at the end of the recording (on average 15%). Series resistance was compensated at 80%.

For action potentials, data were acquired in 10 s sweeps and filtered at 10kHz. Frequency was calculated as the average of each sweep. Baseline potential was determined as the average inter-spike potential in the presence of spontaneous activity. Template-based action potential analysis was performed in Clampfit 10 (Molecular Devices) to obtain action potential peak amplitude and inactivate values. After hyperpolarization (AP) peak was determined as the peak negative value of the action potential, and AHF amplitude was the baseline membrane potential minus the AP peak value. Threshold analysis was performed by taking the derivative of each action potential event in the 10s segment to determine the collection in the voltage trajectory (Clampfit). This method gives a set of values per neuron with lower standard deviation than automated analysis using dV/dt of 4–5V s⁻¹ (ref. 49). Action potential analysis was performed with blind with respect to genotype or time point.

In voltage-clamp mode, total voltage-activated K⁺ currents were elicited from a holding potential of −90 mV, stepping from −110 to +110 mV for either 30 or 130 ms in 20 ms increments. BK currents were isolated by focal application of the BK antagonist paxilline (10 μM), and subtracting currents after paxilline from the initial current. The effective concentration of paxilline in the recording chamber was 3.75 μM. Currents were averaged from three to five voltage families. Current–voltage (∆V) relations were constructed from 1 to 110 ms at 30 ms from each potential, unless noted otherwise (peak or steady-state values). Current density values were obtained by normalizing to cell capacitance, which did not differ between day and night (7.0 ± 0.5 and 6.8 ± 0.4 pF, respectively) or between WT and β2 KO (7.7 ± 0.7 and 7.1 ± 0.4 pF, n = 27, 20, respectively). Voltage values were adjusted for the liquid junction potential of 10 mV, tcurrent values were calculated from single exponential fits of the macroscopic current decay from the +90 mV step for 150 ms (Clampfit, Molecular Devices). BK and BK currents were categorized as described using the current ratio at 30 ms from +90 mV step (BK < 0.07, BK > 0.67), corresponding to tcurrent values < 0.110 ms. For passive membrane properties, whole-cell recordings were performed as above using the same internal solution as for current- and voltage-clamp recording, with tetrodotoxin (1 μM) in the bath. After break in, the membrane potential was recorded, followed by current injection steps (0 to −25 pA, in 5 pA increments). Input resistance was calculated as the linear slope of the voltage. For experiments where BK currents were categorized as described using the current ratio at 30 ms from +90 mV step, BK current clamp mode after recording Vh and R, and BK currents were isolated and tcurrent values were obtained as described in the previous section.
For BK, and BK, recordings of $V_{m}$, $R_{in}$, subthreshold SCN BK currents, and BK currents evoked by action potential commands, recordings were performed in 1 mM tetrodotoxin. After recording $V_{m}$ and $R_{in}$ as described above, cells were switched to voltage-clamp mode and stepped from a holding potential of $-90$ to $+90$ mV for 150 ms to determine BK, and BK, by $I_{K, BK}$ as described previously. After holding at $-150$ mV for 100 ms, cells were stepped from $-60$ to $-30$ mV (in 5 mV increments), and then stepped back to $-150$ mV for 100 ms followed by action potential commands. Previously acquired native action potential waveform was obtained from daytime or night time SCN recordings. For day (Fig. 10c) and night (Supplementary Fig. 7), the action potential parameters are indicated in the figure legends. Three sequenced action potentials were applied at the average BK, daytime firing rate (2.25 Hz) or night time BKs firing rate (1 Hz) as depicted in the figures. Current–voltage relationships were plotted, and the action potential-evoked peak and subthreshold currents were normalized to the cell capacitance.

**Drugs and β2N peptides.** Paxilline and tetrodotoxin (Tocris, Bristol, UK) were prepared as 1,000 stocks in DMSO and water, respectively. β2N was synthesized as the first 45 amino acids of the β2 subunit (M1F1WTSGRSSSRYQDEKR-NITYQKIRDLLDKRKTVAKLAGED) (GenScript, Piscataway, NJ, USA)β2N194203 was a 42 amino acid peptide with the underlined FIW deletion. Lyophilized peptides were reconstituted in intracellular solution as 100 stocks and added at the final indicated concentrations.

**HEX cell patch-clamp electrophysiology.** HEK293T cells (ATCC, Manassas, VA, USA) were transfected with the BKα1a α subunit cDNA (Gembark accession #JX46275), with or without β2 (ref. 34), both in pCDNA3.1. DNA was transfected at 1 µg (x only) or 1 µg α and 1.7 µg β2 per 35 mm dish using TransIT (Mirus Bio, Madison, WI, USA). Cells were plated on glass coverslips 3–5 h later and recorded from 20 to 30 h after transfection. Patch-clamp recording was performed using the voltage protocols indicated in the Figure legends. Data were recorded at 50 kHz and filtered at 10 kHz. The bath (intracellular) solution was composed of (mM): 140 KMeSO3, 2 KCl, 2 HEPES and 5 HEDTA. Free [Ca2+] as described previously60. Currents were elicited from patches using the voltage protocols indicated in the Figure legends. Data were recorded at 50 kHz and filtered at 10 kHz. The bath (intracellular) solution was composed of (mM): 140 KMeSO3, 2 KCl, 2 HEPES and 5 HEDTA. Free [Ca2+] was produced by pore block from an associated particle. 

**Statistics.** The number of neurons (BK current and action potential recordings), the number of mice (circadian behaviour) or the number of SCN slice cultures. Whole recordings above. The action potential-evoked peak current, or the subthreshold current, was normalized to the cell capacitance.

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Author contributions
J.P.W. and A.L.M. designed the study. J.P.W. performed the experiments and statistical analysis. J.P.W. and A.L.M. analysed the data. J.R.M. analysed action potential data and performed statistical analysis. A.L.M. wrote the manuscript. All authors approved the final manuscript.

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