Casein Kinase 1 Underlies Temperature Compensation of Circadian Rhythms in Human Red Blood Cells

Andrew D. Beale,* Emily Kruchek,* Stephen J. Kitcatt,* Erin A. Henslee,* Jack S.W. Parry,* Gabriella Braun,* Rita Jabr,* Malcolm von Schantz,* John S. O’Neill†,‡ and Fatima H. Labeed*,†

*Faculty of Engineering and Physical Sciences, University of Surrey, Guildford, Surrey, UK, †Faculty of Health and Medical Sciences, University of Surrey, Guildford, Surrey, UK, and ‡Medical Research Council Laboratory of Molecular Biology, Francis Crick Avenue, Cambridge, UK

Abstract Temperature compensation and period determination by casein kinase 1 (CK1) are conserved features of eukaryotic circadian rhythms, whereas the clock gene transcription factors that facilitate daily gene expression rhythms differ between phylogenetic kingdoms. Human red blood cells (RBCs) exhibit temperature-compensated circadian rhythms, which, because RBCs lack nuclei, must occur in the absence of a circadian transcription-translation feedback loop. We tested whether period determination and temperature compensation are dependent on CKs in RBCs. As with nucleated cell types, broad-spectrum kinase inhibition with staurosporine lengthened the period of the RBC clock at 37°C, with more specific inhibition of CK1 and CK2 also eliciting robust changes in circadian period. Strikingly, inhibition of CK1 abolished temperature compensation and increased the Q10 for the period of oscillation in RBCs, similar to observations in nucleated cells. This indicates that CK1 activity is essential for circadian rhythms irrespective of the presence or absence of clock gene expression cycles.

Keywords casein kinase, temperature compensation, erythrocyte, dielectrophoresis, electrophysiology

Most organisms use endogenous timekeeping mechanisms to coordinate their physiology with the external cycle of day and night. These circadian rhythms have a cell-autonomous basis (Balsalobre et al., 1998; Welsh et al., 2004) orchestrating many cellular functions, such as DNA repair, protein synthesis, and cell division, to an approximately 24-h beat (Dekens et al., 2003; Kang et al., 2009; Lipton et al., 2015; Matsuo et al., 2003). Circadian rhythms bestow a selective advantage upon organisms in rhythmic and arrhythmic settings by synchronising them with predictable daily environmental cycles and facilitating the temporal segregation of mutually antagonistic processes (Beale et al., 2016; Vaze and Sharma, 2013). In most cell types, these temporal programs are thought to be dependent on the rhythmic regulation of gene expression by oscillating “clock gene” activity, in turn facilitated by transcriptional feedback repression of clock gene transcription by their own protein products (Takahashi, 2017).

1. To whom all correspondence should be addressed: Fatima Labeed, Faculty of Engineering and Physical Sciences, University of Surrey, Guildford, Surrey GU2 7XH, UK; e-mail: f.labeed@surrey.ac.uk.
2. To whom all correspondence should be addressed: John O’Neill, Medical Research Council Laboratory of Molecular Biology, Francis Crick Avenue, Cambridge. CB2 0QH, UK; +44 1223 267037; e-mail: oneillj@mrc-lmb.cam.ac.uk.

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Transcriptional oscillations are not essential for circadian rhythms; bona fide circadian rhythms occur in the complete absence of transcription in the alga *Ostreococcus tauri* (O’Neill et al., 2011), in human red blood cells (RBCs) (O’Neill and Reddy, 2011), and in other organisms (Lakin-Thomas, 2006; Tomita et al., 2005; Woolum, 1991). Importantly, these examples of transcription-independent circadian rhythms satisfy the 3 fundamental properties that define circadian clocks (Pittendrigh, 1960): they are entrained by relevant environmental cues, such as light or temperature; they free run with an approximately 24-h period; and they are temperature compensated, meaning that they do not run faster at higher temperatures, i.e., \( Q_{10} \approx 1 \), unlike most chemical and biochemical reactions, where \( Q_{10} = 2 \).

Although temperature compensation is a defining feature of circadian clocks, it remains poorly understood. Some period-determining processes, and thus candidate mechanisms that facilitate temperature compensation, are known, and these tend to involve the phosphorylation of clock proteins by Ser/Thr kinases (Gallego and Virshup, 2007). One important and evolutionarily conserved post-translational modifier is casein kinase 1 (CK1), which phosphorylates non-orthologous clock protein targets, such as PERIOD (PER) in mammals and FREQUENCY (FRQ) in fungi (He et al., 2006; Meng et al., 2008), and has been shown to regulate period and temperature compensation of circadian rhythms across eukaryotes (Causton et al., 2015; He et al., 2006; Hirota et al., 2010; Meng et al., 2008; van Ooijen et al., 2013).

Whilst mutation of individual PERIOD homologs can alter circadian period by several hours, these perturbations have only modest effects on temperature compensation (Dibner et al., 2009). Recently, it was shown that CK18/ε-dependent phosphorylation is intrinsically temperature compensated, with a CK1-specific domain around K224 sufficient to confer temperature compensation on other kinases (Isojima et al., 2009; Shinohara et al., 2017). Since temperature compensation and the circadian function of CK1 exhibit far greater evolutionary conservation than any known clock protein substrate or clock gene transcriptional circuit, we hypothesised that the timekeeping and temperature compensation function of CK1 might extend beyond its known interaction with clock gene transcriptional circuits to include transcription-independent circadian rhythms. Here, we employed a reduced model of non-transcriptional circadian rhythms—isolated human RBCs—to test whether period determination and temperature compensation remain dependent on CK1 activity in the absence of clock protein expression and cycling transcription.

**MATERIALS AND METHODS**

**Experimental Model and Subject Details**

Studies were conducted in accordance with the principles of the Declaration of Helsinki, with approval/favourable opinion from our local Research Ethics Committee. Participants in the study were screened for relevant self-reported health issues, including sleep disorders or excessive daytime sleepiness. All participants provided written, informed consent after having received a detailed explanation of the study procedures. RBCs were isolated from male participants, with RBCs from 4 separate donors for each treatment or experimental condition.

**Human RBC Isolation**

RBCs were isolated following the protocol of Henslee et al. (2017). RBCs were isolated from anticoagulant-treated whole blood (lithium heparin Vacutainer, BD Biosciences; Franklin Lakes, NJ) by density gradient centrifugation, layering a mixture (1:2.67) of whole blood and Dulbecco’s phosphate-buffered saline (DPBS; Sigma-Aldrich, St Louis, MO) on top of Histopaque-1077 (Sigma-Aldrich) at 50 \( \times g \) for 30 min. RBCs were washed once in DPBS (250 \( \times g \) for 15 min) and once in modified Krebs-Henseleit buffer (KHB; Sigma-Aldrich) (250 \( \times g \) for 10 min). KHB was modified by supplementation with calcium chloride dihydrate (to a final concentration of 2.54 mM), sodium bicarbonate (25 mM), bovine serum albumin (1 g/l), penicillin (100 U/l) and streptomycin (100 µg/l), and adjusted to pH 7.4 and 290-299 mOsm. After washing, 330 µl of packed RBCs were added to 40 ml of KHB.

**Circadian Entrainment**

RBCs were subjected to daily temperature cycles (12 h, 37°C; 12 h, 32°C) for 2 days using a programmed thermal cycler before transfer into constant temperature conditions of either 37°C or 32°C (as specified). Samples were taken at 3- or 4-hourly intervals upon transfer into constant conditions, with a single 500-µl PCR tube of RBCs at each time point for each donor. Following convention, the transition to constant conditions was taken as \( t=0 \).

**Dielectrophoresis**

After the 48-h entrainment, at each timepoint, a single PCR tube was removed from the thermal cycler to assess cell electrophysiology by dielectrophoresis.
(DEP). The modified KHB medium was removed and the pellet was suspended in iso-osmotic DEP medium (248 mM sucrose, 16.7 mM dextrose, 250 mM MgCl₂, 100 mM CaCl₂, adjusted to 290-299 mOsm and a conductivity of 0.043 S/m using DPBS). Cells in DEP medium were then centrifuged for 2 min at 500 × g, and the medium was replaced with fresh DEP medium to reduce ionic carryover from the high ionic strength of the KHB medium. The cells were subsequently adjusted to a final concentration of 10⁶ per ml and pipetted into 3DEP chips (Labtech; Heathfield, East Sussex, UK). Chips were analysed by a 3DEP 3D Dielectrophoresis Cell Analysis System (3DEP reader, www.labtech.com/3dep-3d-dielectrophoresis-cell-analysis-system) to extract the electrophysiological parameters; models that produced an R value (Pearson correlation coefficient) of 0.95 or lower were excluded.

Bioluminescence Imaging

Human U2OS cells (HTB-96, ATCC; Manassas, VA) were stably transfected with BMAL1:Luc and cultured in Dulbecco’s Modified Essential Medium (DMEM) supplemented with 10% HyClone FetalClone II Serum (Thermo Fisher Scientific; Waltham, MA), 1 mM luciferin (Biosynth AG; Staad, Switzerland), 100 U/mL penicillin, 100 µg/mL streptomycin, and 2 µg/mL puromycin (to maintain selection). Four days before bioluminescence imaging, cells were trypsinized, resuspended in culture medium and were maintained at a constant temperature and 5% CO₂ (Crosby et al., 2017). The recording media (± drug or vehicle) and the plates were placed into an ALLIGATOR for bioluminescence imaging, and were maintained at a constant temperature and 5% CO₂ (Crosby et al., 2017). The recording media was identical to the culture media except that the serum concentration was reduced to 1%; B27 supplement (Thermo Fisher) was added to a final concentration of 2%, and puromycin was omitted. For constant time-course experiments. The drugs used in this study—staurosporine, PF-670462, D4476 (all Sigma-Aldrich), and TTP22 (Tocris Biosciences; Bristol, UK)—were prepared in DMSO according to the manufacturer’s instructions. Blots were blocked in 5% (w/v) non-fat dried milk (Marvel) dissolved in Tris-buffered saline containing 0.05% Tween-20 (TBST) for 1 h at room temperature, and then incubated overnight at 4°C on a rotary shaker with primary antibodies diluted in blocking buffer. Primary antibodies were as follows: rabbit anti-CK1α (1:1000, ab108296; Abcam, Cambridge, UK), mouse anti-CK1ε (1:1000, ab85320; Abcam), mouse anti-CK1ε (1:2000, ab82426; Abcam), rabbit anti-CK2α’ (1:1000, ab10474; Abcam) and rabbit anti-CK2β (1:1000, ab133576; Abcam). Blots were washed for 3 × 5 min in TBST then incubated for 1 h with HRP-conjugated anti-rabbit (1:5000) or antimouse (1:10,000) secondary antibodies. Blots were washed again for 3 × 15 min with TBST before chemiluminescence detection using Immobilon Western Chemiluminescent HRP Substrate (Millipore; Billerica, MA), according to manufacturer’s instructions. RIPA extract lysates of U2OS cells transfected with mammalian expression constructs of V5-CK1α (Addgene plasmid #92014, pCAGImC V5-CSNK1A1, a gift from Joshua Mendell), CK1ε-V5His, HA-CK1ε (Addgene plasmid #13724, V405 4HA-CK1ε, a gift from David Virshup), HA-CK2α’ and Myc-CK2β (Addgene plasmids #27087 and #27088, pZW16 [CK2alpha’] and pZW12 [CK2beta] respectively; gifts from David Litchfield) served as positive controls.

Pharmacological Treatments

For each drug, cell viability and morphology were initially screened over a period of 2 to 4 days using a range of concentrations. Concentrations of the drug that did not result in hemolysis or altered viability or morphology under our experimental conditions (following Henslee et al., 2017) were chosen for subsequent time-course experiments. The drugs used in this study—staurosporine, PF-670462, D4476 (all Sigma-Aldrich), and TTP22 (Tocris Biosciences; Bristol, UK)—were prepared in DMSO according to
the manufacturer’s instructions. Drugs were diluted in KHB and delivered as a 10X concentration bolus to RBC suspensions to achieve the final concentration. In each case, an equivalent % of DMSO was used as a vehicle control. Drugs were added to RBC suspensions at ZT0 on day 1 (staurosporine) or circadian time (CT)-0 on day 3 (PF-670462, D4476, TTP22), as indicated in the figure legend. For human U2OS cells, PF-670462 was diluted from a 10 mM DMSO stock directly into recording medium, and was delivered to cells through a media change at the beginning of the recording. This was compared with vehicle control (between 0.03% to 0.1% DMSO).

Sequence Comparisons

The following sequences were compared: the catalytic domain (8-282) and C-terminus (283-end) of human CK1 delta isoform 1 (NP_001884), isoform 2 (NP_620693) and isoform 3 (NP_001350678); the catalytic domain (isoform 1: 16-309; isoforms 2 and 3: 16-281) and C-terminus (isoform 1: 309-end; isoforms 2 and 3: 282-end) of human CK1 alpha isoform 1 (NP_00120276), isoform 2 (NP_001883.4) and isoform 3 (NP_001258670.1). Orthologous sequences in Arabidopsis thaliana (NP_193170), Neurospora crassa (ESA44278), Ostreococcus tauri (OUS42226) and Drosophila melanogaster (CK1 alpha: CAA64358; dou-

Quantification and Statistical Analysis

All data are presented as the mean ± SEM, where n refers to the number of biological replicates. To determine whether a circadian rhythm was present in time-course experiments, and subsequently the circadian parameters period, amplitude and phase, a damped cosine was fit to the data, as described by Hirota et al., (2008), using least squares non-linear regression curve fitting in GraphPad Prism (v7.03, Hirota et al., (2008), using least squares non-linear damped cosine was fit to the data, as described by dian parameters period, amplitude and phase, a time-course experiments, and subsequently the circa-

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Statistically significant differences were defined as those with a p value < 0.05. The significance level is indicated as *p < 0.05, **p < 0.01 and ***p < 0.001.

RESULTS

We previously reported the circadian regulation of membrane electrophysiology and potassium transport in isolated human RBCs under constant conditions using an assay based on DEP (Henslee et al., 2017). Kinases fulfill essential timing functions in the circadian clocks of other mammalian cells (i.e., those with nuclei) and we were therefore interested in understanding whether this extended to encompass the transcription-independent circadian clock of RBCs. We used staurosporine, a broad-spectrum kinase inhibitor first shown to dramatically alter the free-running period in the dinoflagellate Gonyaulax polyedra (Comolli and Hastings, 1999), as a proof-of-principle test of the hypothesis that kinases regulate the RBC circadian clock. Staurosporine treatment dramatically increased the period of rhythms in membrane conductance (G_{eff}) and cytoplasmic conductivity (\sigma_{cvl}) (Fig. 1A-C), similar to its activity in nucleated cells (Comolli and Hastings, 1999). RBCs are therefore an amenable platform with which to test kinase function in the RBC clockwork using small-molecule inhibitors.

Casein kinases play functionally conserved roles in timekeeping across eukaryotes, with roles attributed to CK1 homologs (CK1\alpha, CK1\beta, CK1\epsilon) and CK2 (He et al., 2006; Hirota et al., 2010; Meng et al., 2008; van Ooijen et al., 2013). RBCs express one CK1 paralogue, CK1\alpha, as well as catalytic and regulatory subunits of CK2: CK2\alpha\gamma, and CK2\beta (Fig. 2A and Supplementary Figure S1) (Bryk and Wiśniewski, 2017). Therefore, we explored whether their timekeeping role extended to non-transcriptional oscillations in RBCs by applying casein kinase inhibitors that are known to affect period

For the calculation of Q_{10} value, the period values were entered into the following equation:

$$Q_{10} = \left(\frac{\tau_1}{\tau_2}\right)^{10}$$

where \tau_1 and \tau_2 are the period at temperature T_1 and T_2, respectively. T_1 = 32\degree C and T_2 = 37\degree C. Rhythm parameters (i.e., period), derived from the best fit \cosine, were compared using the extra sum-of-squares F comparison test and 2-way ANOVA. If ANOVA revealed significant (p < 0.05) effects, Holm-Sidak’s multiple comparison post hoc tests were used to determine p values for all relevant comparisons. All analyses were performed in Graphpad Prism. 

$$Y(X) = mX + c$$

was compared with a damped cosine + baseline fit:

$$Y(X) = mX + \text{Baseline} + \text{Amplitude} \times e^{-ax} \times \cos \left(\frac{2\pi(X - \text{Phaseshift})}{\text{Period}}\right)$$

where X = time, using the extra sum-of-squares F comparison test in GraphPad Prism, with the simpler model being preferred unless the p value was < 0.05.
**Figure 1.** RBC circadian rhythms are affected by the inhibition of protein kinases. Isolated RBCs were cultured in the presence of 10 μM staurosporine (STS) or vehicle control at constant 37°C after 2 days of temperature entrainment. DEP measurements of membrane conductance ($G_{eff}$) (A) and cytoplasmic conductivity ($\sigma_{cyt}$) (B) are reported (mean ± SEM, $n = 4$). Damped cosine curves were fitted to the data using non-linear regression comparison of fits ($H_1 =$ damped cosine model, $H_0 =$ straight line) extra sum-of-squares F test. Cosine fit was preferred over straight line fit if $p < 0.05$. $G_{eff}$ and $\sigma_{cyt}$ oscillate in antiphase, as previously reported (Henslee et al., 2017). (C) STS treatment significantly increases the period of both $G_{eff}$ and $\sigma_{cyt}$ rhythms relative to vehicle control (period values derived from best fit cosinor ± SEM; $n = 4$. Two-way ANOVA $p$ value ($\beta_{TWA}$) and Holm-Sidak's post hoc tests reported).

**Figure 2.** Casein kinase inhibition alters the circadian rhythm of red blood cells (RBCs). (A) Immunoblots showing the expression profiles of human CK1 and CK2 paralogs in human RBCs and in human U2OS cells. Data from 2 donors are shown. U2OS cells transfected with V5-CK1α (Golden et al., 2017), CK1β-V5, HA-CK1ε (River et al., 1998), HA-CK2α' and Myc-CK2β (both Turovec et al., 2010) served as positive controls. (B-D) Dielectrophoresis (DEP) measurements of membrane conductance ($G_{eff}$) in isolated RBCs incubated at constant 37°C over 2 days in the presence of 100 nM TTP22 (CK2 inhibitor) (B), 1 μM D4476 (CK1 inhibitor) (C), or 3 μM PF-670462 (highly potent CK1 inhibitor) (D) versus vehicle control (mean ± SEM, $n = 4$. Extra sum-of-squares F test $p$ values reported from non-linear regression comparison of fits, $H_1 =$ damped cosine model, $H_0 =$ straight line). (E) The period of CPR rhythms are significantly affected by inhibition of CK1 and CK2 (period values derived from best fit cosinor ± SEM, $n = 4$. Extra sum-of-squares F test $p$ values reported following comparison of best fit cosinor. **$p < 0.01$, *$p < 0.05$.**
determination in nucleated cells (Badura et al., 2007; Meng et al., 2010). We found that TTP22, an inhibitor of CK2 (Golub et al., 2011), and D4476, an inhibitor of CK1 (Bain et al., 2007; Rena et al., 2004), elicited significant effects on the period of the circadian rhythm in RBC membrane electrophysiology, lengthening, and shortening period, respectively (Fig. 2B, C, E). This demonstrates a role for these enzymes in the transcription-independent time-keeping mechanism of RBCs.

A lengthening of the circadian period in response to CK2 inhibition has been observed previously for other mammalian and eukaryotic models (Tsuchiya et al., 2009). We were surprised, however, that CK1 inhibition elicited a significantly shorter circadian period in RBCs compared with controls, because, to our knowledge, in all other contexts CK1 inhibition slows down the cellular clock (Badura et al., 2007; Hirota et al., 2010; Isomina et al., 2009; Meng et al., 2010; van Ooijen et al., 2013). We therefore repeated our assays using a pharmacologically distinct and highly potent inhibitor of CK1, PF-670462 (Badura et al., 2007; Bibian et al., 2013). This resulted in a similar reduction in circadian period length as that observed with D4476 (Fig. 2D and E).

Temperature compensation is an evolutionarily conserved property of the circadian clock and a defining feature of bona fide circadian rhythms. CK1 and CK2 have proposed roles in temperature compensation in organisms as diverse as *Neurospora* fungi, algae and mammals (Isomina et al., 2009; Mehra et al., 2009; Tosini and Menaker, 1998; Zhou et al., 2015). We examined the role of these cellular kinases in temperature compensation in the absence of known clock protein targets. If temperature compensation is reliant on the activity of either kinase, their inhibition will result in a $Q_{10}$ for the period of oscillation that lies outside the temperature-compensated range of 0.8 to 1.2; i.e. inhibition of the kinase should have markedly different effects on circadian period at different temperatures. As shown in previous studies in nucleated cells (Isomina et al., 2009), inhibition of CK1 disrupted temperature compensation in (nucleated) human U2OS cells expressing a BMAL1:Luc reporter, with $Q_{10}^{\text{PF670}} = 1.60 \pm 0.14$ (Fig. 3A and C). Consistent with daily rhythms becoming temperature-dependent under CK1 inhibition, in a separate experiment, the magnitude of period lengthening due to CK1 inhibition was further reduced in U2OS cells when cultured at the higher temperature of 40°C (Supplementary Figure S2). In contrast, CK2 inhibition had no effect on temperature compensation, i.e. $Q_{10}^{\text{TTP22}} = 1.15 \pm 0.13$ was not significantly different from the vehicle $Q_{10}^{\text{control}} = 1.02 \pm 0.11$ (Fig. 3C). In anucleate RBCs, as with U2OS cells, temperature compensation was observed both for vehicle control

Figure 3. Temperature compensation of RBC circadian rhythms is abolished by inhibition of casein kinase 1 (CK1). (A) Nucleated human U2OS cells (BMAL1:Luc) were cultured at constant 32°C (top, light tone) and 37°C (bottom, full tone) following 4 days of temperature entrainment in the presence of 0.31 μM PF-670462 (CK1 inhibitor, red) or 2.5 μM TTP22 (CK2 inhibitor, blue) vs. vehicle control (black). In vehicle control (black) (mean ± SEM, n = 6), detrended using a moving average of 24-h for vehicle and TTP22 or 30-h for PF-670462. (B) Dielectrophoretic (DEP) measurements of membrane conductance ($G_{\text{mem}}$) in isolated RBCs over 2 days incubated at 32°C in the presence of 3 μM PF-670462 (light red) or 100 nM TTP22 (light blue) vs. vehicle control (mean ± SEM, n = 4). P values for DEP assays report non-linear regression comparison of fits ($H_1$ = damped cosine model, $H_2$ = straight line). (C and D) Inhibition of CK1 abolishes temperature compensation in U2OS (top) and RBCs (bottom). In vehicle control (black) and TTP22-treated (blue) cells, period does not change with increased temperature, reflected by a $Q_{10}$ of approximately 1. Conversely, treatment with PF-670462 results in a significantly increased $Q_{10}$ vs. control, reflected by the shorter period at 37°C relative to 32°C. Period values derived from best fit $T_{\text{min}}$ vs. SEM (nU2OS = 6, nRBC = 4). Holm-Sidak’s multiple comparisons $p$-value reported, following one-way ANOVA (panova $= 0.015$, pANOVA $= 0.017$), for $Q_{10}$ vs. vehicle. *p < 0.05.
with period shortening at 37°C, and a Q₁₀ of 2.60 ± 0.46 (Fig. 3B and D). Therefore, inhibition of CK1, but not CK2, activity abolished temperature compensation of circadian rhythms in the presence and absence of transcription-translation feedback loops (Fig. 3C and D).

**DISCUSSION**

Both CK1 and CK2 play roles in circadian period determination in many eukaryotic cellular contexts (O’Neill et al., 2013). CK1 and CK2 homologs are present in RBCs (this study and Bryk and Wiśniewski, 2017), whereas the capacity for nascent transcription is not. Here, we have extended the timekeeping function for these ancient and ubiquitously expressed kinases to encompass regulation of circadian period in the non-transcriptional RBC clock. Whilst it is clear that PER (and other clock gene encoded) proteins are functionally relevant substrates for CK1 and CK2 in cells where cycling clock gene transcription is present, our RBC assays suggest clock protein transcription factors (TFs) are unlikely to be their only relevant substrates with respect to cellular timekeeping. Separately, this is also implied by the complete conservation of CK1 and CK2 in period determination across eukaryotes (Causton et al., 2015; He et al., 2006; Lin et al., 2002; Meng et al., 2008; Tsuchiya et al., 2009; van Ooijen et al., 2013; Yang et al., 2002; Zhang et al., 2013), whereas the clock protein TFs, upon which they act, are not. A parsimonious interpretation of this observation is that there exist one or more CK substrates, evolutionarily conserved but as yet unidentified, whose activity determines periodicity.

In the absence of cycling transcription and clock gene activity, we also found that the catalytic activity of CK1, but not CK2, appears to be relevant to the mechanisms that facilitate temperature compensation of RBC circadian rhythms, similar to the results of nucleated mammalian cells. In two very different human cell types, primary RBCs and the U2OS osteosarcoma cell line, inhibition of CK1 abolished temperature compensation and increased the Q₁₀ for the period of oscillation. Inhibition of CK1 with PF-670462 had a greater effect on the Q₁₀ of rhythms in RBCs than in U2OS cells (Q₁₀ of 2.6 compared with 1.6, respectively), thus explaining the period shortening elicited by CK1 inhibition in RBCs at 37°C (Fig. 3 and Supplemental Figure S2). The differential sensitivity of Q₁₀ to CK1 inhibition between the two cell types may be attributable to cell type-dependent variation in the expression of CK1 isoforms; i.e., in U2OS cells, the α, δ, and ε isoforms were all readily detectable, whereas only CK1α was detected in RBCs, both in our study and in an independent mass spectrometry-based investigation (Bryk and Wiśniewski, 2017).

Unlike CK1δ/ε, the timekeeping functions of CK1α are less well established, being reported by two separate studies (Hirotö et al., 2010; Lam et al., 2018). Thus, an alternative interpretation of our findings is that temperature-compensated circadian rhythms in RBCs arise by a completely different mechanism from that which operates in other mammalian cell types. In this case, the similar sensitivity to CK inhibition of RBC circadian rhythms compared with other cell types, as well as their common features, such as rhythmic K⁺ transport and over-oxidised peroxiredoxin levels (Cho et al., 2014; Edgar et al., 2012; Feeney et al., 2016; Henslee et al., 2017; O’Neill and Reddy, 2011), is simply coincidental. We note, however, that the CK1α catalytic domain shares very high sequence identity with human CK1δ/ε (78%) and CK1 orthologs in the metazoan, fungal and green lineages (>71%), whereas the C-terminal domains that distinguish mammalian CK1 paralogs and isoforms from each other show much less conservation across the eukaryotic kingdoms. We also note that, whilst CK1ε is dispensable, CK1δ and CK1α both play essential roles in development and proliferation, possess identical substrate specificities that cannot be distinguished pharmacologically, and have many cellular targets in common (Schittek and Sinnberg, 2014). Therefore, we suggest that there is very little evidence that CK1α is not equally as important as CK1δ for the regulation of circadian gene expression rhythms in mammalian cells.

Unlike most enzymes (including CK2), the activity of CK1δ/ε toward certain polypeptides is insensitive to temperature, with a Q₁₀ of around 1.0 (Isojima et al., 2009; Shinohara et al., 2017; Zhou et al., 2015). This temperature insensitivity is inherent to the catalytic domain of CK1, demonstrated by impaired temperature compensation of catalytic mutants against synthetic substrates compared with native sequences (Shinohara et al., 2017). This “temperature-compensated domain” is conserved in other CK18 homologs, such as human CK1α (86% similarity), including the critical lysine residue (K224 in CK1δ; K232 in CK1α). This strongly suggests that temperature compensation is a function of CK1 homologs across the eukaryotic kingdom, independent of the presence or absence of the TTFL, and functions similarly between nucleated and non-nucleated cells.

In this, we note similarities with KaiC phosphorylation in cyanobacteria, whose post-translational
rhythm persists in the absence of nascent gene expression (Nakajima et al., 2005; Tomita et al., 2005). Here, temperature compensation of circadian period is conferred by the intrinsically temperature-insensitive phosphotransferase activity of KaiC (Egli et al., 2012; Terauchi et al., 2007), with circadian period and temperature compensation also being sensitive to amino acid substitution around the ATP-binding site (Murakami et al., 2008; Terauchi et al., 2007). Our present observations support the possibility that the role of CK1 homologs in eukaryotic circadian rhythms may be functionally analogous to that of KaiC in cyanobacteria (Merrow et al., 2006; Wong and O’Neill, 2018).

The circadian clock regulates biological functions, such as metabolism, protein synthesis, and ion transport, in many cell types across phylogenetic kingdoms, both in the presence and absence of transcriptional feedback cycles (Cho et al., 2014; Feeney et al., 2016; Henslee et al., 2017; Lipton et al., 2015; O’Neill et al., 2011; O’Neill and Reddy, 2011; van Ooijen et al., 2011; Woolum, 1991). Though no definitive adaptive advantage has been ascribed to RBC circadian rhythms, daily variation in CO2-carrying potential, adverse cardiovascular events, and osmotic homeostasis are potential consequences of this reduced and highly specialised cellular clockwork (Henslee et al., 2017). Erythrocytes are the most numerous cell type in the human body. Whereas their circadian clock mechanism is not understood, it does involve the same conserved kinases that regulate timekeeping fidelity in other mammalian cell types. Future studies should identify which RBC proteins might be subject to circadian post-translational modification by CK1 and CK2, and whether these also exhibit phosphorylation rhythms in other cell types.

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CONFLICT OF INTEREST STATEMENT

The author(s) have no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

NOTE

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ORCID ID

Andrew D. Beale https://orcid.org/0000-0002-2051-0919

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