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A Role for the Circadian Clock Protein Per1 in the Regulation of the NaCl Co-transporter (NCC) and the with-no-lysine Kinase (WNK) Cascade in Mouse Distal Convoluted Tubule Cells*

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Background: The role of the circadian protein Per1 in the regulation of sodium reabsorption in the distal convoluted tubule (DCT) is unknown.

Results: Per1 transcriptionally regulates the sodium transporter NCC and the WNK kinase cascade.

Conclusion: Per1 regulates sodium reabsorption in the DCT through NCC and the WNK cascade.

Significance: These data demonstrate a role for Per1 in the regulation of renal sodium transporters.

It has been well established that blood pressure and renal function undergo circadian fluctuations. We have demonstrated that the circadian protein Per1 regulates multiple genes involved in sodium transport in the collecting duct of the kidney. However, the role of Per1 in other parts of the nephron has not been investigated. The distal convoluted tubule (DCT) plays a critical role in renal sodium reabsorption. Sodium is reabsorbed in this segment through the actions of the NaCl co-transporter (NCC), which is regulated by the with-no-lysine kinases (WNKs). The goal of this study was to test if Per1 regulates sodium transport in the DCT through modulation of NCC and the WNK kinases, WNK1 and WNK4. Pharmacological blockade of nuclear Per1 entry resulted in decreased mRNA expression of NCC and WNK1 but increased expression of WNK4 in the renal cortex of mice. These findings were confirmed by using Per1 siRNA and pharmacological blockade of Per1 nuclear entry in mDCT15 cells, a model of the mouse distal convoluted tubule. Transcriptional regulation was demonstrated by changes in short lived heterogeneous nuclear RNA. Chromatin immunoprecipitation experiments demonstrated interaction of Per1 and CLOCK with the promoters of NCC, WNK1, and WNK4. This interaction was modulated by blockade of Per1 nuclear entry. Importantly, NCC protein expression and NCC activity, as measured by thiazide-sensitive, chloride-dependent 22Na uptake, were decreased upon pharmacological inhibition of Per1 nuclear entry. Taken together, these data demonstrate a role for Per1 in the transcriptional regulation of NCC, WNK1, and WNK4.

The circadian clock regulates a multitude of physiological functions, such as blood pressure (BP),2 metabolism, the immune response, sleep-wake cycles, and renal function (reviewed in Refs. 1 and 2). In terms of renal function, sodium excretion, glomerular filtration rate, renal blood flow, and plasma aldosterone levels exhibit rhythmic fluctuations (reviewed in Ref. 3). The molecular mechanism of the circadian clock consists of four core proteins, which interact with one another to regulate expression of circadian target genes (4). The four core proteins are CLOCK, Bmal1, Period (Per1–3), and Cryptochrome (Cry1 and -2). CLOCK and Bmal1 heterodimerize and interact with E-box response elements to transcriptionally up-regulate circadian target genes, which include Per and Cry. Per and Cry interact and then repress the transcriptional activity of CLOCK and Bmal1 (5). In addition to transcriptional regulation, the circadian clock also undergoes post-translational modifications through the phosphorylation of the Per proteins by the circadian kinases casein kinase 1 isoforms δ/e (CK1δ/e). Phosphorylation by CK1δ/e allows Per1 nuclear entry (6, 7).

We have shown previously that Per1 regulates the basal and aldosterone-mediated expression of αENaC, the regulated subunit of the renal epithelial sodium channel (ENaC) (7–10). We also recently demonstrated that Per1 coordinately regulates the expression of multiple genes that are involved in the regulation of renal sodium reabsorption in collecting duct cells (8). This includes the positive regulation of Fxyd5, a positive regulator of the Na,K-ATPase (11), and the negative regulation of endothelin-1, a powerful inhibitor of ENaC (12). It was initially presumed that Per1 behaved primarily as a repressor of CLOCK and Bmal1 activity; however, increasing evidence suggests that Per1 can activate gene expression through an unknown mech-

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2 The abbreviations used are: BP, blood pressure; CK1δ/e, casein kinase 1 δ/e; Ckh, casein kinase 1 δ/e inhibitor; ENaC, epithelial sodium channel; DCT, distal convoluted tubule; NCC, NaCl co-transporter; WNK, with-no-lysine kinase; KS-WNK1, kidney-specific WNK1; hnRNA, heterogeneous nuclear RNA; het, heterozygote; qPCR, quantitative real-time PCR; Pol, polymerase.
Per1 in NCC and WNK Cascade Regulation

anism, in a manner that appears to be gene- and tissue-specific (8–10, 13–15). We have recently demonstrated that a putative mechanism for Per1 action may occur through repression of the circadian repressor Cry2 (16). Taken together, these data predict that loss of Per1 should result in decreased renal sodium reabsorption, with subsequent decreased plasma volume and decreased BP. Consequently, we have shown previously that Per1 KO mice exhibit significantly lower BP (8) and increased urinary sodium levels and urine volume (9) compared with WT controls. Consistent with these results, we have shown recently that mice with reduced levels of Per1 have a defect in renal sodium retention and decreased plasma aldosterone (17).

We have shown that Per1 plays an extensive role in the regulation of ENaC and multiple other genes that contribute to renal sodium reabsorption in the collecting duct. However, the potential role of Per1 in other parts of the nephron has not been investigated. The distal convoluted tubule (DCT), which precedes the collecting duct, plays a critical role in sodium reabsorption in the renal cortex. Sodium is reabsorbed in this nephron segment through the actions of the NaCl co-transporter (NCC). NCC is post-translationally regulated through the actions of the with-no-lysine kinases (WNKs), including WNK1 and WNK4 (18). WNK1 increases NCC activity through activation of Ste20/SPS-1-related proline/alanine-rich kinase (19, 20), a downstream kinase of the WNKs; at baseline, WNK4 appears to inhibit NCC activity (21). The goal of this study was to test whether Per1 contributes to the regulation of sodium reabsorption in the DCT through transcriptional modulation of NCC and the WNK kinases. Pharmacological blockade of nuclear Per1 entry resulted in decreased mRNA expression of NCC and WNK1 and increased expression of WNK4 mRNA in the renal cortex of mice. To investigate this potential regulation further, a recently generated cell model of the mouse DCT (mDCT15 cells) was employed (22–24). Using several independent methods, we demonstrate a novel role for Per1 in the transcriptional regulation of NCC, WNK1, and WNK4. Importantly, NCC function, as measured by thiazide-sensitive, chloride-dependent 22Na uptake, was decreased upon pharmacological inhibition of Per1 nuclear entry. Stable knockdown of WNK4 prevented this effect. Taken together, these data demonstrate a role for Per1 in the coordinate regulation of sodium reabsorption in mouse DCT cells.

EXPERIMENTAL PROCEDURES

Animals—All animal use protocols were approved by the University of Florida and North Florida/South Georgia Veterans Affairs Institutional Animal Care and Use Committees in accordance with the National Institutes of Health Guide for the Care and Use of Laboratory Animals. WT and Per1 KO mice (129/sv) mice were originally provided by Dr. David Weaver (University of Massachusetts) (25). WT and Per1 heterozygote (het) mice were bred in house by University of Florida Animal Care Services staff and maintained on a normal 12-h light/dark cycle. For in vivo PF670462 (CK18/e inhibitor) experiments, weight-matched male WT 129/sv mice were given either vehicle (20% hydroxypropyl β-cyclodextrin) or 30 mg/kg PF670462 subcutaneously every 12 h for 2.5 days starting at noon and sacrificed at midnight 12 h after the last injection. Mice were anesthetized by inhalant isoflurane, and tissues were collected and snap-frozen in liquid nitrogen. The dose and time of PF670462 administration have been published previously and described by others (26). Kidneys were later dissected, and cortex was removed for protein or RNA isolation.

Cell Culture—mDCT15 cells were maintained in DMEM/F-12, 5% heat-inactivated FBS, and 1% penicillin-streptomycin-neomycin at 37 °C. For RNA silencing experiments, non-target siRNA (Dharmacon) and a previously characterized individual siRNA for Per1 (9) were used according to the manufacturer’s instructions and as described previously (7, 16). Cells were transfected using Dharmafect 1 (Dharmacon). Casein kinase 18/e inhibitor experiments were performed as described previously (7, 16). For dose and time course, mDCT15 cells were treated with either 0.1, 1, 10, or 100 μM casein kinase 18/e inhibitor PF670462 or vehicle (water) for 24, 48, or 72 h (and as described previously (7)). For nuclear entry experiments, mDCT15 cells were treated with 10 μM PF670462 or vehicle for 48 h.

Assessment of NCC Function—The determination of NCC activity in mDCT15 cells was performed as described previously (22). Briefly, mDCT15 cells were seeded in 12-well plates and prepared as described. mDCT15 cells were treated with either 0.1, 1, 10, or 100 μM casein kinase 18/e inhibitor PF670462 or vehicle (water) for 48 h prior to uptake. Thirty minutes before uptake, 0.1 mM metolazone was added to the medium. The medium was then changed to a 22Na+-containing medium (140 mM NaCl, 1 mM CaCl, 1 mM MgCl, 5 mM HEPES/Tris, pH 7.4, 1 mM amiloride, 0.1 mM bumetanide, 0.1 mM benzamil, 1 mM ouabain, and 1 μCi/ml 22Na+) with or without thiazide (0.1 mM metolazone) and incubated for 20 min. Tracer uptake was then stopped via washes with ice-cold wash buffer. Cells were subsequently lysed with 0.1% SDS. Radioactivity was measured via liquid scintillation, and protein concentrations of the lysates were determined (BCA protein assay, Pierce). Uptakes were normalized to nmol/mg. Thiazide-sensitive uptake was given by the difference of the uptakes with and without thiazide.

RNA Isolation and Quantitative Real-time PCR—Total RNA was isolated and DNase I-treated using Direct-Zol™ RNA Miniprep (Zymo Research) as per the manufacturer’s instructions. DNase I-treated RNA (2 μg) samples were used as template for reverse transcription with the High Capacity cDNA Reverse Transcription Kit (Applied Biosystems). The resulting cDNAs (20 ng) were then used as a template in quantitative real-time PCR (qPCR) reactions (Applied Biosystems) to evaluate changes in Per1, NCC, WNK1, WNK4, and actin mRNA levels. Cycle threshold (Ct) values were normalized against β-actin, and relative quantification was performed using the ΔΔCt method (27). -Fold change values were calculated as the change in mRNA expression levels relative to the control. TaqMan primer/probe sets were purchased from Applied Biosystems.

Membrane Protein Analysis, Nuclear Protein Isolation, and Western Blot Analysis—Membrane protein was isolated using the Cell Surface Protein Isolation Kit (Pierce) according to the manufacturer’s instructions. Nuclear extracts were isolated using the NE-PER kit (Pierce) according to the manufacturer’s instructions. Protein concentrations were then quantified by
Per1 in NCC and WNK Cascade Regulation

Briefer et al.

Table 1

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Table 3

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decreased levels of NCC (Fig. 1A) and WNK1 (Fig. 1B). Conversely, CK1β/ε inhibition resulted in significantly increased levels of WNK4 (Fig. 1C). To assess if PF670462 treatment in vivo resulted in decreased nuclear Per1 protein levels, Western blot analysis was performed on nuclear fractions obtained from the mice described above. As shown, nuclear Per1 levels were significantly decreased by more than 50% (Fig. 1D; quantified in Fig. 1E).

Per1 Knockdown in Vitro or in Vivo or Pharmacological Inhibition of Per1 Nuclear Entry in mDCT15 Cells Results in Decreased mRNA Expression of NCC and WNK1 and Increased Expression of WNK4—To further investigate our in vivo findings, Per1 was knocked down using siRNA in a model of the distal convoluted tubule, mDCT15 cells. Total mRNA levels of NCC, WNK1, and WNK4 were measured by qPCR. As expected, Per1 knockdown resulted in significantly decreased mRNA expression of Per1 (Fig. 2A), as we have shown previously (7, 8, 10, 16). Consistent with our in vivo findings, Per1 knockdown resulted in significantly decreased expression of NCC (Fig. 2B) and WNK1 (Fig. 2C) but increased expression of WNK4 (Fig. 2D).

To further assess the role of Per1 in the regulation of NCC, WNK1, and WNK4 in vivo, mRNA levels of NCC, WNK1, and WNK4 were assessed by qPCR in WT and Per1 het mice euthanized at midnight (Fig. 3). We have previously demonstrated that Per1 het mice have a ~50% reduction of Per1 in the renal cortex (16). Consistent with our in vitro results and our in vivo CK1β/ε inhibitor results, mRNA expression of NCC and WNK1 were both significantly decreased in Per1 het mice when compared with WT (Fig. 3, A and B). Conversely, WNK4 mRNA expression was significantly increased (Fig. 3C).

To further assess the regulation of NCC, WNK1, and WNK4 mRNA expression by blockade of Per1 nuclear entry, mDCT15 cells were treated with CK1β/ε inhibitor, and NCC, WNK1, and WNK4 mRNA expression was determined by qPCR. After 48 h of CK1β/ε inhibition, treatment with 10 μM PF670462 resulted in significant reduction of NCC and WNK1 mRNA (Fig. 4, A and B), whereas WNK4 was significantly up-regulated after 24, 48, or 72 h of 10 μM CK1β/ε inhibition (Fig. 4C). The goal of this experiment was to determine the lowest effective dose, as described previously (7). Although the data points for WNK4 at the 100 μM dose were difficult to interpret, it was clear that 10
M was the lowest effective dose. Thus, the 10 μM dose was used for subsequent experiments.

To confirm that CK1/ε inhibition decreased Per1 nuclear entry, mDCT15 cells were treated with vehicle or 10 μM PF670462 for 48 h, and nuclear extracts were collected with Western blotting performed to assess nuclear Per1 levels. As we have shown previously in other cell types (7, 16), nuclear Per1 levels were significantly decreased with PF670462 treatment (Fig. 5A; quantified in Fig. 5B). It should be noted that nuclear Per1 was detected at a molecular mass of just less than 50 kDa, which has been consistently demonstrated in Western blots and pull-down experiments in vivo in the liver and the kidney and in vitro in many different cell types (7, 8, 10, 16).

Pharmacological Inhibition of Per1 Nuclear Entry Results in Decreased Transcription of NCC and WNK1 and Increased Transcription of WNK4 and KS-WNK1—Measurement of the short lived hnRNA is a measure of transcriptional activity (31, 32). Therefore, to assess if the effect of CK1/ε inhibition and Per1 knockdown on the inhibition on NCC, WNK1, KS-WNK1, and WNK4 was transcriptional, hnRNA levels were assessed by PCR amplification of intron-exon junctions using cDNA templates from mDCT15 cells treated with

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vehicle or 10 μM PF670462 for 48 h or non-target and Per1 siRNA for 48 h. KS-WNK1 is a kinase-dead splice variant, unique to the kidney, that inhibits the activity of WNK1 (33). Both Per1 siRNA-mediated knockdown and CK1 inhibition led to significantly decreased hnRNA expression of both NCC (Fig. 6A) and WNK1 (Fig. 6B) but increased expression of KS-WNK1 (Fig. 6C) and WNK4 (Fig. 6D).

**Pharmacological Inhibition of Per1 Nuclear Entry Results in Increased Transcription of KS-WNK1 in Vivo**—To assess if KS-WNK1 is up-regulated in vivo, hnRNA levels were assessed in WT versus Per1 het mice and in vehicle-treated versus CK1δ/ε-treated mice (Fig. 7). Significantly increased hnRNA expression of KS-WNK1 was observed in Per1 het mice compared with

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**FIGURE 4.** Pharmacological blockade of Per1 nuclear entry results in decreased NCC and WNK1 expression and increased WNK4 expression in vitro. mDCT15 cells were treated with the casein kinase 1δ/ε inhibitor PF670462 with either 0.1, 1, 10, or 100 μM for 24, 48, or 72 h. qPCR was used to evaluate changes in NCC (A), WNK1 (B), or WNK4 (C) expression in PF670462-treated cells versus vehicle (water)-treated cells (n = 3). *, p < 0.05; **, p < 0.01. Values are represented as the mean ± S.E. (error bars).

**FIGURE 5.** Pharmacological blockade of Per1 nuclear entry results in decreased nuclear Per1 in vitro. A, nuclear extracts were collected from mDCT15 cells treated with 10 μM casein kinase 1δ/ε inhibitor PF670462 or water for 48 h. Western blot analysis was performed using anti-Per1 or anti-β-actin antibodies as a loading control. Data are representative of three independent experiments. B, densitometry analysis was used to quantitate the level of Per1 in A (n = 3). *, p < 0.05. Values are represented as the mean ± S.E. (error bars).
WT and in CK1δ/ε-treated mice compared with vehicle-treated controls, confirming our in vitro findings.

**Pharmacological Inhibition of Per1 Nuclear Entry Results in Decreased Binding of Per1 and CLOCK to the Promoters of Slc12a3 and Wnk1 but Increased Binding to the Wnk4 Promoter**—As discussed above, circadian regulation of target genes is mediated through interaction of the core circadian proteins with E-box response elements in target gene promoters. To assess if Per1 and CLOCK interact with the endogenous promoters of Slc12a3, Wnk1, and Wnk4 and if CK1δ/ε inhibition could modulate this interaction, ChIP was performed in mDCT15 cells treated with either vehicle or 10 μM PF670462 for 48 h. Three primer sets per gene were designed to amplify short regions containing putative E-box sites within approximately the first 10 kb upstream of the transcription start site (Figs. 8A, 9A, and 10A). Loading was demonstrated by performing PCR using 10% of the input as template (Figs. 8B, 9B, and 10B). IgG was used as a negative control (Figs. 8F, 9F, and 10F).
CK1δ/e inhibition led to significantly decreased binding of RNA Pol II at all three sites in the Slc12a3 promoter (Fig. 8C), indicative of decreased transcription, further corroborating our mRNA and hnRNA data. CLOCK and Per1 were detected at all three NCC promoter regions, and these interactions were significantly decreased with CK1δ/e inhibition (Fig. 8, D and E). On the Wnk1 promoter, CK1δ/e inhibition led to significantly decreased RNA Pol II binding at all three promoter regions (Fig. 9C). As with Slc12a3, CLOCK and Per1 were detected on all three promoter regions, and binding was significantly decreased upon CK1δ/e inhibition (Fig. 9, D and E). On the Wnk4 promoter, RNA Pol II has increased binding upon CK1δ/e inhibition in all three promoter regions, indicative of increased transcription, in agreement with the results of the hnRNA assay (Fig. 10C). CK1δ/e inhibition led to increased CLOCK and Per1 interaction with the second and third promoter regions but no apparent interaction with the promoter region amplified by the first primer set (Fig. 10, D and E).

**Pharmacological Blockade of Per1 Nuclear Entry in Vitro and in Vivo Results in Increased mRNA Levels of Cry1/2 and Per2 in Kidney Cortex and mDCT15 Cells**—Recent evidence from our laboratory and others has demonstrated that Per1 may act on target gene expression through modulation of the circadian repressors Cry1/2 and Per2 (15, 16, 34). We have demonstrated that inhibition of Per1 results in up-regulation of Cry1/2 mRNA and protein levels and that Per1 and Cry2 mediate opposing actions on target gene expression in the kidney, liver, and adrenal glands in vitro and in vivo (16, 17). However, this has not been investigated in the context of CK1δ/e inhibition in vitro in mDCT15 cells or in vivo in kidney cortex. Therefore, mRNA levels were assessed by qPCR for Clock, Bmal1, Cry1, Cry2, and Per2, in either mDCT15 cells (Fig. 11) or WT animals treated with CKInh (Fig. 12). Pharmacological blockade of Per1 nuclear entry in vitro did not affect Clock or Bmal1 mRNA (Fig. 11, A and B). Consistent with our previous observations, however, CKInh treatment resulted in significantly increased mRNA levels of Cry1, Cry2, and Per2 (Fig. 11, C–E). These results were corroborated in vivo with CKInh treatment (Fig. 12). These confirm our previously reported results that we have shown for Cry2 both in vitro and in vivo in kidney, liver, and the adrenal gland (17) and that others have shown previously for Per2 and Cry1 (34).

**Pharmacological Inhibition of Per1 Nuclear Entry Results in Decreased Membrane and Intracellular Protein Expression of NCC**—To test whether the effects of CK1δ/e inhibition on NCC expression extended to the level of protein, plasma membrane and intracellular fractions were isolated from mDCT15 cells treated with vehicle or 10 μM PF670462 for 48 h. Western blot analysis was performed to determine changes in NCC protein levels from both intracellular and membrane fractions. CKInh treatment led to a significant decrease of NCC at the intracellular level and a ~40% decrease at the membrane level (Fig. 13). Decreases at both the intracellular and membrane level are consistent with our hypothesis that regulation of NCC by Per1 occurs at the translational level.

**Pharmacological Inhibition of Per1 Nuclear Entry Results in Decreased NCC Activity**—Having demonstrated that pharmacological inhibition of Per1 nuclear entry affects expression of NCC, WNK1, and WNK4, NCC activity was measured with and without CK1δ/e inhibition to assess for an effect on NCC function. The mDCT15 cells were treated with vehicle or 1, 10, or 100 μM PF670462 for 24 or 48 h prior to measurement of NCC activity by thiazide-sensitive 22Na+ uptake. At 24 h, no significant change was seen in NCC activity in the CK1δ/e inhibition groups (Fig. 14A). However, at 48 h, NCC activity declined by 32 ± 7% in the 10 μM group and by 39 ± 6% in the 100 μM group (Fig. 14B, n = 4, p < 0.05 compared with vehicle for both). This significant decrease parallels the decrease in NCC membrane protein levels (Fig. 13). Interestingly, a similar treatment in mDCT15 cells with stable knockdown of WNK4 expression (generated as described previously (22)) showed a sharply attenuated decline in NCC activity with CK1δ/e inhibition (non-significant at 10 μM, 8 ± 2% for 100 μM, n = 4, p < 0.05, Fig. 14C).

**DISCUSSION**

The results of the present study demonstrate a potential role for Per1 in the transcriptional regulation of NCC and the WNK cascade. We demonstrated that pharmacological blockade of Per1 nuclear entry or decreased Per1 in vivo and in vitro results in decreased mRNA expression of NCC and WNK1 but increased expression of WNK4 and KS-WNK1. Furthermore, ChIP and hnRNA assays performed in the mouse distal convoluted tubule cell line, mDCT15, consistently demonstrated that Per1 acts on the NCC/WNK pathway via a transcriptional mechanism and that this regulation extends to the level of protein. Importantly, we show for the first time that NCC activity was decreased upon pharmacological inhibition of NCC expression by thiazide-sensitive 22Na+ uptake in vitro.
Pharmacological blockade of Per1 nuclear entry leads to decreased occupancy of CLOCK and Per1 on the Slc12a3 promoter in mDCT15 cells. Chromatin immunoprecipitation experiments were performed using mDCT15 cells treated with either vehicle (water) or 10 μM PF670462 for 48 h. End point PCR was performed using primer sets flanking three putative E-boxes in the Slc12a3 promoter (A, not drawn to scale). B, input control. Chromatin immunoprecipitations were performed using anti-Pol II (Santa Cruz Biotechnology) (C), anti-CLOCK (Pierce) (D), anti-Per1 (Pierce) (E), or rabbit IgG (Bethyl) (negative control) (F) antibodies. Bands were quantitated using densitometry, which was performed using ImageJ. Signal strength was normalized to the relevant vehicle- or CKinh-treated input control (n = 3; *, p < 0.05; **, p < 0.01). Values are represented as the mean ± S.E. (error bars).
FIGURE 9. Pharmacological blockade of Per1 nuclear entry leads to decreased occupancy of CLOCK and Per1 on the Wnk1 promoter in mDCT15 cells. Chromatin immunoprecipitation experiments were performed using mDCT15 cells treated with either vehicle (water) or 10 μM PF670462 for 48 h. End point PCR was performed using primer sets flanking three putative E-boxes in the WNK1 promoter (A, not drawn to scale). B, input control. Chromatin immunoprecipitations were performed using anti-Pol II (Santa Cruz Biotechnology) (C), anti-CLOCK (Pierce) (D), anti-Per1 (Pierce) (E), or rabbit IgG (Bethyl) (negative control) (F) antibodies. Bands were quantitated using densitometry, which was performed using ImageJ. Signal strength was normalized to the relevant vehicle or CKinh treated input control (n = 3; *, p < 0.05; **, p < 0.01). Values are represented as the mean ± S.E. (error bars).
FIGURE 10. Pharmacological blockade of Per1 nuclear entry leads to increased occupancy of CLOCK and Per1 on the Wnk4 promoter in mDCT15 cells. Chromatin immunoprecipitation experiments were performed using mDCT15 cells treated with either vehicle (water) or 10 μM PF670462 for 48 h. End point PCR was performed using primer sets flanking three putative E-boxes in the WNK4 promoter (A, not drawn to scale). Input control. Chromatin immunoprecipitations were performed using anti-Pol II (Santa Cruz Biotechnology) (C), anti-CLOCK (Pierce) (D), anti-Per1 (Pierce) (E), or rabbit IgG (Bethyl) (negative control) (F) antibodies. Bands were quantitated using densitometry, which was performed using ImageJ. Signal strength was normalized to the relevant vehicle- or CKinh-treated input control (n = 3; *, p < 0.05; **, p < 0.01). Values are represented as the mean ± S.E. (error bars).
Per1 nuclear entry, an effect that appeared to be partially dependent on WNK4. Based on these data, we propose a model for Per1 action on the NCC and WNK pathway in which Per1 promotes increased sodium reabsorption in the distal convoluted tubule, through transcriptional regulation of NCC, WNK1, KS-WNK1, and WNK4 (Fig. 15). This apparent coordinate regulation of pathway genes by Per1 suggests a role for Per1 in driving renal epithelial sodium reabsorption in DCT cells; this coordinate regulation is entirely consistent with our previously proposed model for Per1 action on ENaC in CCD cells (8).

We also demonstrated that CK1δ/ε inhibition led to increased transcription of KS-WNK1, as measured by increased amplification of the intron-exon sequence of the KS-WNK1-specific Exon4A, which is the first exon in KS-WNK1 and replaces the first four exons of WNK1 (33). However, further evidence is required to understand the role of Per1 in the regulation of KS-WNK1. The WNK kinase pathway includes many other factors in addition to WNK1, WNK4, and KS-WNK1. These include WNK3, which promotes sodium reabsorption by activating NCC and inhibiting WNK4. STE20/SPS-1-related proline/alanine-rich kinase and oxidative stress-responsive kinase 1 were determined to be downstream targets of the WNK cascade and are presumably responsible for the phosphorylation of NCC itself (reviewed in Refs. 21 and 35). It was recently demonstrated that the levels of phosphorylated oxidative stress-responsive kinase 1, Ste20/SPS-1-related proline/alanine-rich kinase, and NCC exhibited circadian fluctuations in mice (36), a result that is consistent with our observations demonstrating regulation of this pathway by circadian clock proteins. Interestingly, a microarray study showed that NCC, WNK1, and WNK4 did not appear to oscillate with a circadian pattern (37). More work is needed to determine if the Per1-mediated regulation of NCC and the WNK kinases is independent of time. Consistent with our previous observations in the liver and kidney (16), knockdown or inhibition of Per1 in the present study resulted in increased mRNA expression of Cry2. As well, Cry1 and Per2 mRNA expression was increased following knockdown or inhibition of Per1, but CLOCK and Bmal1 levels did not appear to be affected. Thus, the mechanism of Per1 action probably involves the regulation of other circadian repressors.

Mounting evidence has demonstrated the critical importance of the circadian clock for the regulation of BP control and renal sodium handling (reviewed in Refs. 1, 3, and 38). CLOCK KO mice are relatively hypotensive and display dysregulated urinary sodium excretion and mild diabetes insipidus (37, 39). Cry1/2 KO mice display salt-sensitive hypertension partially due to increased aldosterone levels (40). This result was linked to increased expression of 3β-dehydrogenase isomerase, a rate-
limiting enzyme in the aldosterone synthesis pathway. Per2 KO mice exhibit decreased 24-h diastolic BP, increased heart rate, and a reduction in diurnal dipping (41). Bmal1 KO mice have reduced BP in the active phase, leading to a lack of circadian BP variation (42). We have demonstrated that Per1 KO animals exhibit significantly lower BP than WT controls (8). It is unknown if the BP phenotype of these animals is solely due to modulation of renal sodium retention; however, recently, we have shown that mice with reduced levels of Per1 have a defect in renal sodium retention (17). It is likely that a combination of factors contribute to the BP phenotype of these animals, including the heart (43), sympathetic activity (44, 45), the vasculature (46), and/or nitric oxide (47). However, consistent with our findings that Per1 is important for the regulation of sodium reabsorption in the kidney, we have recently demonstrated that reduced expression of Per1 in mice is associated with decreased renal sodium reabsorption (17). Available data in humans are consistent with a role for Per1 in BP regulation as well; a recent report demonstrated that Per1 was overexpressed in the renal medulla of human hypertensive patients compared with normotensive control patients (48).

Along with our previous data demonstrating the regulation of ENaC and sodium transport genes in collecting duct cells and the kidney (8–10), Per1 appears to regulate gene expression in a cell model of the distal convoluted tubule. It will be interesting to see if Per1 and the circadian clock regulate gene expression in other parts of the kidney. It has been demonstrated that the
sodium hydrogen exchanger NHE3 exhibits circadian oscillations in the mouse renal medulla and is regulated by Bmal1/CLOCK (49, 50) (51, 52). Taken together, the data presented here demonstrate a novel role for Per1 in the regulation of sodium reabsorption in distal convoluted tubule cells via transcriptional regulation of NCC and several kinase genes from the WNK cascade.

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