Regulated phosphorylation of the K-Cl cotransporter KCC3 is a molecular switch of intracellular potassium content and cell volume homeostasis

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The defense of cell volume against excessive shrinkage or swelling is a requirement for cell function and organismal survival. Cell swelling triggers a coordinated homeostatic response termed regulatory volume decrease (RVD), resulting in K+ and Cl− efflux via activation of K+ channels, volume-regulated anion channels (VRACs), and the K+–Cl− cotransporters, including KCC3. Here, we show genetic alanine (Ala) substitution at threonines (Thr) 991 and 1048 in the KCC3a isoform carboxyl-terminus, preventing inhibitory phosphorylation at these sites, not only significantly up-regulates KCC3a activity up to 25-fold in normally inhibitory isotonic conditions, but is also accompanied by reversal of activity of the related bumetanide-sensitive Na+–K+–2Cl− cotransporter isoform 1 (NKCC1). This results in a rapid (<10 min) and significant (>90%) reduction in intracellular K+ content (Kᵢ) via both Cl-dependent (KCC3a + NKCC1) and Cl-independent [DCPIB (VRAC inhibitor)-sensitive] pathways, which collectively renders cells less prone to acute swelling in hypotonic osmotic stress. Together, these data demonstrate the phosphorylation state of Thr991/Thr1048 in KCC3a encodes a potent switch of transporter activity, Kᵢ homeostasis, and cell volume regulation, and reveal novel observations into the functional interaction among ion transport molecules involved in RVD.

Keywords: K-Cl cotransporters, KCC3, NKCC1, SPAK, cell volume homeostasis, regulatory volume decrease, cerebral edema, neurodegeneration

Introduction

Regulation of cell volume is critical for multiple essential cellular functions and organismal survival. Lacking a rigid cell wall, animal cells effectively combat excessive cell swelling or shrinkage, induced by perturbations in intracellular ion content or extracellular osmolality, by triggering concerted counter-responses termed regulatory volume decrease (RVD) or regulatory volume increase (RVI),
respectively (Kregeow, 1971, 1981; Hoffmann and Dunham, 1995; Lauf and Adragna, 2000, 2012; Hoffmann et al., 2009). These mechanisms are highly regulated by signaling molecules that sense changes in intracellular ionic content and/or cell volume, and transduce these signals to the cell membrane to modulate the transport of ions and/or organic osmolytes via the stimulation or inhibition of ion transporters, pumps, and channels (Hoffmann and Dunham, 1995). The SLC12A family of cation-Cl^-cotransporters [including the Na^+-K^+-2Cl^- cotransporter isoform 1 NKCC1 and the K^+-Cl^-cotransporters (KCCs), such as KCC3], the Na^+/H^+ exchangers (e.g., NH,E1), the Na^+/K^+ pump, and volume-regulated anion channels (VRACs), are important plasmalemmal mediators of ion transport in RVI and RVD (Hoffmann and Dunham, 1995; Lauf and Adragna, 2000, 2012; Hoffmann et al., 2009).

K^+-Cl^-cotransport was first identified as a swelling- and thiol-activated K^+ efflux pathway in low-K^+ sheep red blood cells (Dunham et al., 1980; Lauf and Therg, 1980). The four KCC isoforms (KCC1-4) utilize energetically favorable, outwardly-directed K^+ gradients to drive the extrusion of Cl^- across the plasma membrane. As such, they serve as important determinants of both intracellular K^+ and Cl^- content, which are important for cell volume regulation and other essential functions depending on cell type (e.g., epithelial transport and neuronal excitability) and KCC isoform (Lauf and Adragna, 2012). The physiological importance of the swelling-activated KCCs, and in particular KCC3 (SLC12A6), is clearly illustrated by the phenotypes that result from the knockout of the genes encoding these molecules in mouse (Boettger et al., 2002; Delpierre and Mount, 2002), and their mutations in humans (Howard et al., 2002).

Loss-of-function mutations in human KCC3 cause the autosomal recessive disease Andermann syndrome (OMIM # 218000), characterized by agenesis of the corpus callosum, peripheral neuropathy (ACCPN), and seizures (Delpierre and Mount, 2002; Howard et al., 2002; Uyanik et al., 2006). KCC3 KO mice recapitulate the human ACCPN phenotype (Howard et al., 2002; Boettger et al., 2003) and studies in these mice have shown that disease pathogenesis is associated with neuroglial swelling and subsequent neurodegeneration resulting from impaired RVD (Byun and Delpierre, 2007). Dysfunction of the volume-regulated KCCs has been implicated in other cell volume-related pathologies (Adragna et al., 2004, 2006; Adragna and Lauf, 2007), such as RBC dehydration in sickle cell disease (Brugnara et al., 1986; Brugnara, 1993, 1995, 2001; Joiner, 1993; Lew and Bookchin, 2005) and inner ear cell degeneration in deafness (Boettger et al., 2003).

Despite their physiological importance, the regulation of the KCCs is not fully understood, though serine-threonine kinases/phosphatases have long been known to play an essential role (Jennings and Al-Rohil, 1990). Phosphorylation inhibits the KCCs, while dephosphorylation has the opposite effect (Dunham et al., 1980; Altamirano et al., 1988; Jennings and Schulz, 1991; Lytle and Forbush, 1992; Haas and Forbush, 2000; Adragna et al., 2004). KCC1, KCC3, and KCC4 are inactive in isotonic conditions but briskly activated in response to hypotonic, low Cl^- cell-swelling conditions by a regulatory mechanism that is evolutionarily-conserved (Haas and Forbush, 2000; Adragna et al., 2004; Strange et al., 2006). KCC activation by hypotonic cell swelling is prevented by calyculin A, an inhibitor of protein phosphatase 1A (PP1) and PP2, further demonstrating the importance of phosphorylation for KCC regulation (Lauf and Adragna, 2000).

Two threonine (Thr) residues in human KCC3a, Thr991, and Thr1048, were recently shown to undergo rapid dephosphorylation in response to hypotonic low Cl^- (swelling/activating) conditions in HEK293 cells and human RBCs (Rinehart et al., 2009; De Los Heros et al., 2014); physiologically, these sites are phosphorylated in isotonic (inhibitory) conditions. Homologous sites (i.e., “site 1” and “site 2”) are phosphorylated in all human KCCs, including KCC2, and double alanine substitution at these residues, preventing phosphorylation, result in constitutively-active KCC2 and KCC3 (Rinehart et al., 2009; De Los Heros et al., 2014). The with no lysine [K] (WNK)-regulated STEP20/SPS1-related proline/alanine-rich kinase (SPAK) and oxidative stress responsive kinase-1 (OSR1) kinases, known to regulate the activity of the KCCs in oocytes (e.g., Kahle et al., 2005; Gagnon et al., 2006), were recently shown to directly phosphorylate site 2 in the KCCs, but not site 1 (De Los Heros et al., 2014). The kinase that directly phosphorylates site 1 in the KCCs has not yet been identified (Rinehart et al., 2009).

Despite the detailed biochemical characterization of the swelling-induced KCC3 Thr991/Thr1048 dephosphorylation mechanism, the cellular physiology of this event has not been systematically explored. Here, we utilized unidirectional net ion flux uptake/loss assays under zero-trans conditions, to measure intracellular K^+ (K_i) content and uptake of ^85Rb, and cell volume analysis in two isogenic pairs of human epithelial cell lines (HEK-293) engineered with doxycycline-inducible expression of wild type KCC3 (KCC3 WT) or KCC3 Thr991Ala/Thr1048Ala (i.e., “KCC3 AA,” preventing inhibitory phosphorylation), on (1) KCC3 transport activity; (2) the activity of other key molecules involved in cell volume homeostasis [e.g., NKCC1 and the Na^+/K^+ pump (herein termed “NKP”)]; (3) K_i; and (4) cell volume and RVD in conditions of hypotonic stress.

Materials and Methods

Chemicals

Chemicals from Thermo Fisher Scientific (Fair Lawn, NJ) were: Tris (hydroxymethyl) aminomethane (Tris) free base, 3-morpholin-4-ylpropane-1-sulfonic acid (MOPS), sodium chloride (NaCl), potassium chloride (KCl), magnesium chloride (MgCl_2), sodium hydroxide (NaOH), sucrose, D-glucose, perchloric acid, 70% (PCA), and bicinchoninic acid (BCA) protein assay reagents. Magnesium gluconate was from Sigma-Aldrich (St. Louis, MO). 4-(2-hydroxyethyl)-1-piperazine ethane sulfonic acid (HEPES) free acid, and anhydrous calcium chloride (CaCl_2), were from J.T.Baker Chemical Co (Center Valley, PA). Rubidium chloride (RbCl), 99.8% (metals basis), and amidosulfonic acid (sulfamic acid, S), 99.99% (metals basis) were purchased from Alfa Aesar (Ward Hill, MA); N-methyl D-glucamine (NMDG) from...
Fluka Biochemika (St. Louis, MO); cesium chloride (CsCl) and calcine-AM from Life technologies (Carlsbad, CA) and calcium gluconate from Acros Organics (NJ). Ouabain octahydrate was purchased from Calbiochem (San Diego, CA), furoside and bumetanide from Sigma-Aldrich (St. Louis, MO), 4,4′-(2-Butyl-6,7-dichloro-2-cyclopentyl-2,3-dihydro-1-oxo-1H-inden-5-yl)oxy]butanoic acid (DCP1B), 1,2-Bis(2-aminophenoxo) ethane-N,N,N′,N′-tetra acetic acid (BAPTA) from Tocris Bioscience (Bristol, UK), tetra ethyl ammonium (TEA) from Abcam (Cambridge, MA), clotrim tolsylate from Enzo life sciences (Farmindale, NY), and 2,4-dichloro-N-isopropyl-N- (2-isopropylamino)benzenesulfonamide (RN-1734) and Ruthenium Red (RR) from Santa Cruz Biotechnology (Santa Cruz, CA).

**Solutions**

The solution compositions for the different steps in the flux protocol are as follows (with all salt concentrations in mM). Initial wash: 300 mMolar balanced salt solution (BSS-NaCl) (20 HEPES-Tris, 5 KCl, 2 CaCl2, 1 MgCl2, 10 glucose, 135 NaCl, pH 7.4, 37 °C) or BSS-NaS (20 HEPES -Tris, 5 K+ sulfamate, 2 Ca gluconate, 1 Mg gluconate, 10 glucose, 135 NaS, pH 7.4, 37 °C). Pre-incubation/equilibration: BSS-NaCl-BSA (bovine serum albumin) (300 mMolar BSS-NaCl + 0.1 % BSA, pH 7.4, 37 °C) or BSS-NaS-BSA (300 mMolar BSS-NaS + 0.1 % BSA, pH 7.4, 37 °C). Flux (in mM): 300 mMolar BSS-RbCl-BSA (20 HEPES -Tris, 10 RbCl, 2 CaCl2, 1 MgCl2, 10 glucose, 0.1 % BSA, 135 NaCl, pH 7.4, 37 °C) or BSS-RbS-BSA (20 HEPES -Tris, 10 Rb+ sulfamate, 2 Ca gluconate, 1 Mg gluconate, 10 glucose, 0.1 % BSA, 135 NaS, pH 7.4, 37 °C). Final wash: 300 mMolar containing 10 MOPS-TrisMgCl2, pH 7.4, 37 °C (Supplementary Table 2).

Ions were extracted for 15 min at 4 °C with 5 % perchloric acid (PCA) and measured by atomic absorption spectrophotometry in a Perkin Elmer 5000, as described elsewhere (Adragna et al., 2002; Zhang et al., 2003). Total protein was determined by protein extraction with 1M NaOH and measured with the BCA protein assay as previously described (Di Fulvio et al., 2001).

**Cell Line Construction and Culture**

WT or AA (Thr991Ala/Thr1048Ala) human KCC3a cDNAs harboring an N-terminal myc epitope (gifts of P. Gallagher, Yale University), under the expression control of tetracycline, were introduced as single copies into HEK-293 (Tet-On) cells using the Flp-In™ T-REX™ System as described elsewhere (Rinehart et al., 2009) (Supplementary Figure 2). Doxycycline, a tetracycline derivative, was used to induce gene expression per manufacturer’s directions (Life Technologies-Invitrogen, Grand Island, NY, 14072). Cells were cultured in Dulbecco’s Modified Eagle Medium (DMEM) with high glucose (Life Technologies, Catalog No. 11965-092), supplemented with 10 % tetracycline-free fetal bovine serum (FBS) (Clontech, Catalog No. 631106), 100 units/mL penicillin, 100 µg/mL streptomycin (HyClone, Catalog No. SV30010) and 10 µg/mL blasticidin S hydrochloride (Santa Cruz, Catalog No. sc-204655), and maintained in a humidified incubator with 5 % CO2 at 37 °C. For live cell imaging experiments, viable cells (0.2 × 106 cells/well) were plated on poly-D-Lysine coated glass coverslips (22 × 22 mm) in 6-well plates. To induce KCC3 (WT or AA) expression, cultures were treated with 1 µg/mL doxycycline for up to 16 h.

**Ion Flux Studies**

Ion fluxes were measured according to previously published protocols (Adragna et al., 2002; Zhang et al., 2003) with some modifications. Cells were grown to 60 % confluence in 12-well plates; KCC3 was expressed by overnight doxycycline (0–1 µg/mL) induction. 85Rb+ uptake or influx and intracellular 39K+ (Ki) content were determined under zero-trans conditions as follows. WT- or AA-transfected cells were removed from the incubator; culture medium was aspirated, washed three times with 1 ml 300 mMolar BSS-NaCl or BSS-NaS and then equilibrated in BSS-NaCl-BSA or BSS-NaS-BSA for 10 min. Thereafter, supernatants were removed and then exposed to pre-warmed flux media 300 mMolar BSS-RbCl-BSA or BSS-RbS-BSA during pre-determined time points (0–15 min). 85Rb+ was used as a 39K+ congener (Lauf, 1983; Delpire and Lauf, 1991). Thereafter, cells were washed with washing solution to block Rb+ influx and K+ loss. Cellular Rb+ and K+ were extracted as described above. Intracellular K+ was simultaneously measured with a Na+ -K+ lamp by absorption and Rb+ uptake with 85Rb+ by flame emission spectrophotometry in a Perkin-Elmer 5000 Atomic Absorption spectrophotometer (Perkin-Elmer, Boston MA). Total protein was determined by the BCA assay as described above. Rb+ uptake as a function of time and Ki, (nanomoles per milligram protein) as a function of time were calculated for each well as described elsewhere (Adragna, 1991). Ouabain and bumetanide at final concentrations of 0.1 mM and 10 µM, respectively, were added to the flux media to block the Na+/K+ pump (NKP) and Na+/K+ -2Cl− cotransporter (NKCC), respectively. NKP-mediated Rb+ influx was calculated as the ouabain-sensitive flux (Cl− - ClO) and NKCC as the ouabain-insensitive but bumetanide-sensitive (ClO− - ClOBr) Rb+ influx in Cl− media, The Cl−-dependent Rb+ influx (KCC) is the calculated difference between the Rb+ influx in Cl− and S media, both containing ouabain and bumetanide (ClO− - SOB) (Supplementary Tables 1, 2).

**Cell Volume Measurements**

Cell volume change was determined using calcine as a marker of intracellular water volume, as established previously (Lenart et al., 2004). Briefly, cells on coverslips were incubated with 0.5 µM calcine-AM for 30 min at 37 °C. The cells were placed in a heated (37 °C) imaging chamber (Warner Instruments, Hamden, CT) on a Nikon Ti Eclipse inverted epifluorescence microscope equipped with perfect focus, a 40X Super Fluor oil immersion objective lens, and a Princeton Instruments MicroMax CCD camera. Calcine fluorescence was monitored using a FITC filter set (excitation 480 nm, emission 535 nm, Chroma Technology, Rockingham, VT). Images were collected every 60 s with MetaFluor image-acquisition software ( Molecular Devices, Sunnyvale, CA) and regions of interest (~20–30 cells) were selected. Baseline drift resulting from photo bleaching and dye leakage was corrected as described previously (Lenart et al., 2004). The fluorescence change was plotted as a function of the reciprocal of the relative osmotic pressure and the
resulting calibration curve applied to all subsequent experiments as described previously (Lenart et al., 2004). The HEPES-buffered isotonic solution contained (in mM) 100 NaCl, 5.4 KCl, 1.3 CaCl$_2$, 0.8 MgSO$_4$, 20 HEPES, 5.5 glucose, 0.4 NaHCO$_3$, and 70 sucrose (pH 7.4) with 310 mOsm determined using an osmometer (Advanced Instruments, Norwood, MA). Anisosmotic solutions (150, 280, and 312 mOsm) were prepared by removal or addition of sucrose to the above solution.

Statistical Analysis

The analysis of two samples in each or for independent experiments was done by paired or unpaired t-test. Multiple intergroup differences in each or for independent experiments was conducted by One-Way analysis of variance (ANOVA) or Kruskal–Wallis One-Way-ANOVA test followed by either, paired or unpaired t-test, Wilcoxon signed-rank test, two-sample t-test or Wilcoxon rank-sum test. A $p < 0.05$ was used as the criterion of statistical significance. Except where indicated, all values were obtained from three independent experiments in which at least triplicate samples were assayed.

Results

An Isogenic Cell Line System with Inducible Expression of Wild Type or Constitutively-Dephosphorylated KCC3 (Thr991Ala/Thr1048Ala)

We constructed HEK-293 epithelial cell lines with doxycycline-regulated expression of either N-terminal myc-tagged wild type human KCC3a (“KCC3 WT”) or KCC3a Thr991A/Thr1048A (“KCC3 AA”) inserted as a single copy, as described in Materials and Methods. In the absence of doxycycline, KCC3 WT or AA is not expressed (Supplementary Figure 1). However, in the presence of doxycycline (0.1 µM), anti-myc antibody revealed robust expression of myc-tagged KCC3 WT or KCC3 AA in 1.5 h (Supplementary Figure 1). KCC3 expression and localization was similar across multiple clonal cell lines of both KCC3 WT and KCC3 AA (data not shown). Consistent with previous reports (Rinehart et al., 2009; De Los Heros et al., 2014), KCC3 in isotonic conditions was phosphorylated at both Thr991 and Thr1048 (Supplementary Figure 1), as revealed using phospho-specific antibodies raised against these residues (De Los Heros et al., 2014). Hypotonic low Cl$^-$ conditions that stimulate KCC3 activity elicited dephosphorylation at Thr991 and Thr1048 in KCC3 WT (data not shown), and mutation of these residues to alanine (A) in KCC3 AA prevented phosphorylation at each site (Supplementary Figure 1).

Constitutive KCC3 Thr991/Thr1048 Dephosphorylation Potently Stimulates KCC3 and is Associated with a Reversal of NKCC1 Activity in Flux Conditions

As a reminder $^{85}$Rb$^+$ (a K$^+$ congener) was used in uptake assays and $^{39}$K$^+$ under zero-trans conditions for K$_i$ content determinations to measure the activities of KCC, NKCC, and NKP (see Materials and Methods for details).

It is also important to stress that only one technique is used to determine simultaneously (in the same cells and at the same time) both Rb$^+$ influx and K$_i$ content, which are measured by atomic absorption (K$^+$) and emission (Rb$^+$) spectrophotometry. Briefly, the technique consists of the following main steps: (1) Cells plated on 12-well plates; (2) Culture medium removal from wells at confluence; (3) Overnight incubation with doxycycline (Induction); (4) Washing of cells 3×; (5) Equilibration with solutions of similar composition as the flux solutions containing K$^+$ but not Rb$^+$ (Preincubation); (6) Flux with solutions containing Rb$^+$ but not K$^+$ (Flux); (7) Washing of cells 3 times; (8) Further processing for cations and protein determinations (See Materials and Methods for more details and Supplementary Tables 1, 2).

This technique, which has been developed several decades ago, allows to measure simultaneously unidirectional fluxes under 0-trans conditions (Rb$^+$ influx and K$^+$ efflux) without the use of radioisotopes (Lauf, 1983; Delpire and Lauf, 1991), and was applied when measuring simultaneously K$^+$ loss and Rb$^+$ influx through the IK channel of lens epithelial cells (Lauf et al., 2008). This technique also allows measuring K$_i$ in a simplified manner, especially in KCC3 AA cells, because K$_i$ loss is of such magnitude that the signal to noise ratio is large enough to produce accurate data.

Rb$^+$ influx through KCC is defined as the difference in Rb$^+$ influx between Cl-containing and Cl-free medium with sulfamate (S) as Cl$^-$ replacement, in the presence of both ouabain (a NKP inhibitor) and bumetanide (a NKCC inhibitor); Rb$^+$ influx through NKCC is defined as the ouabain-insensitive and bumetanide-sensitive Rb$^+$ influx in Cl-containing media; and Rb$^+$ influx through NKP is defined as the ouabain-sensitive Rb$^+$ influx in Cl-containing medium (Supplementary Tables 1, 2).

Induction of KCC3 WT in isotonic conditions did not elicit a significant KCC-mediated Rb$^+$ influx. For instance, in control cells (i.e., cells not induced with doxycycline) the Rb$^+$ influx was 10.6 ± 1.8 nmol/mg protein × 5 min, whereas in cells induced with doxycycline (shown in Figure 1B), it was 12.4 ± 0.9 nmol/mg protein × 5 min, $n = 12$ individual determinations from 4 independent experiments, consistent with the presence of inhibitory phosphorylation of KCC3 at Thr991 and Thr1048 (Supplementary Figure 1). Likewise, in KCC3 WT cells, NKCC was 30.2 ± 7.4 in non-induced cells and 29.7 ± 6.0 nmol/mg protein × 5 min in induced cells (shown in Figure 1B), $n = 12$ individual determinations from 4 independent experiments.

In contrast, induction of KCC3 AA caused a significant increase in Rb$^+$ influx in Cl$^-$ and ouabain +/− bumetanide media (CIO and CIoB), which was almost completely inhibited in Cl-free medium (SOB) (Figure 1A), resulting in a greater than ~10-fold elevation of KCC activity [non-induced cells, 19.1 ± 0.42 nmol/mg protein × 5 min and in induced cells (shown in Figure 1B), 266.1 ± 2.2 nmol/mg protein × 5 min, $n = 12$ individual determinations from 4 independent experiments, $p < 0.00001$]. This effect was associated with a ~5-fold decrease in NKCC activity; note the negative NKCC value in Figure 1B that was calculated as the difference between CIO - CIoB Rb$^+$ content.
in flux (plotted in Figure 1A) [non-induced, 108.5 ± 3.2 and in induced (shown in Figure 1B), −40.1 ± 8.5 nmoles/mg protein × 5 min, n = 12 individual determinations from 4 independent experiments, p < 0.0005].

Analysis of the ouabain-sensitive Rb+ uptake component revealed induction of KCC3 AA triggered a ~3–4 fold increase in NKP activation in 5 min (Figure 2B) when compared with KNP activity upon induction of KCC3 WT (Figure 2A, KCC3 WT 23.2 ± 1.0 and KCC3 AA 87.7 ± 12.9 nmol/mg protein − 5 min), consistent with the known 3–4 fold stimulation of NKP flux to its Vmax by intracellular Na+ and extracellular K+ site saturation (reviewed in Lauf and Adragna, 2012). This became further evident when extracellular Na+ replacement by NMDG eliminated NKP activity (Figure 3). No changes were observed in Ki content between non-induced and induced KCC3 WT cells under the same conditions for KCC3 AA cells (data not shown). These data show that prevention of KCC3 Thr991/Thr1408 inhibitory phosphorylation results in constitutive KCC3a activity that is accompanied by concurrent NKCC1 reversal (i.e., outward Rb+ efflux through NKCC1) and NKP stimulation.

KCC3 Thr991/Thr1408 Dephosphorylation Triggers Significant Cl-Dependent and Cl-Independent Ki Loss

Because intracellular ions and water are determinants of cell volume, any inward ion flux study would be incomplete without considering the fate of the intracellular counter-ion content, in our case K+. Furthermore, as opposed to the Rb+ influx data shown above, KCC-mediated transport actually results in K+ loss (efflux) and requires Cl−. Figure 4 shows Ki in KCC3 WT (Figure 4A) and KCC3 AA (Figure 4B) cells measured after exposure to Cl− or S media in conditions of NKP inhibition by ouabain, and NKCC inhibition by ouabain + bumetanide, in both non-induced and induced cells. Note that in KCC3 WT cells, there was a 30% loss of Ki in Cl−-free medium (601 ± 5.2 in CIOr vs. 417 ± 10.7 in SOb, i.e., 31.5 ± 1.5% in non-induced and 609 ± 10.6 in CIOr and 423 ± 15.9, i.e., 30.6 ± 1.8% in induced WT cells, n = 9 individual determinations for all the conditions).

Because this K+ loss occurred in S but not in Cl− media, it could not have occurred via KCC3, and instead appeared to involve an unidentified K+ conductive route, such as a selective
K⁺ channel, non-selective cation channel, or transporter. Surprisingly, induction of KCC3 AA triggered a ∼75 % Kᵢ loss in Cl⁻ media, 50 % of which was recovered by incubating the cells in S medium (see Figure 4B, compare ClO in non-induced and induced cells, and ClO with SOB in induced cells, labeled as Cl-dependent K⁺ loss), indicating inhibition of a Cl-dependent Kᵢ loss mechanism presumably through NKCC and KCC3. This Cl-dependent Kᵢ loss through NKCC and KCC3 increased as a function of time, further depletion of the cells K⁺ (data not shown), and was inhibited by replacing extracellular Rb⁺ with K⁺, and 2 mM furosemide, an inhibitor of KCC activity (data not shown). In contrast, the remaining 50 % Kᵢ loss was Cl-independent (see Figure 4B, compare SOB in non-induced and induced cells).

The Cl-dependent and Cl-independent Kᵢ loss triggered by KCC3 AA in isotonic conditions has not been previously reported. The Cl-dependent K⁺ loss that presumably occurs via KCC3 (see above), was matched by a significant increase in Cl-dependent Rb⁺ influx, respectively. KCC3 AA but not KCC3 WT (results not shown) induction causes a potent increase in flux through KCC3, which is accompanied by inhibition of NKCC and stimulation of NKP. The effects of KCC3 AA induction on NKCC and NKP are abolished when extracellular Na⁺ is replaced with NMDG. Use of the Cl-free medium SOB partially reduces the magnitude of this Kᵢ decrease, indicating that there are both Cl-dependent and Cl-independent components of the Kᵢ decrease stimulated by KCC3 AA induction. Cl-dependent Kᵢ loss in induced cells was calculated as the Kᵢ content in SOB—that in ClO, whereas the Cl-independent Kᵢ loss was calculated as the difference in Kᵢ content in SOB between non-induced and induced cells. Flux time, 5 min; n = 9 individual determinations from a single experiment for WT and n = 4 for AA cells. Total number of independent experiments N = 4 for WT and N = 2 for AA cells. There was a statistically significant difference between groups as determined by Kruskal–Wallis One-Way ANOVA for Non-induced vs. Induced (ClO, ClO, and SOB) for WT [F(5, 48) = 21.55] and AA [F(5, 18) = 36.97], p = 0.0005, n = 24, and 24, respectively. These differences were confirmed by paired t-test for (ClO vs. SOB) in Non-induced and Induced WT, p < 0.0005, and AA, p < 0.05 (Non-induced) and p < 0.0005 (Induced); whereas the comparison between Non-induced vs. Induced AA was p < 0.0005 for ClO and ClO, and p < 0.0005 for SOB. Data represent the mean ± SEM values.

To explore mechanisms contributing to the Cl-independent Kᵢ loss, we performed ion replacement studies and tested inhibitors of several different candidate transport systems (see Supplementary Table 3). KCC3 AA-induced Cl-independent Kᵢ loss was unaffected by the following: Changes in extracellular Na⁺ (data not shown); TEA (2 mM) and clofilium (100 µM), inhibitors of multiple K⁺ channels (Iwatsuki and Petersen, 1985; Arena and Kass, 1988; Gough and El-Sherif, 1989); Ruthenium Red (1 µM), a non-selective inhibitor of Ca²⁺ conductance; and stimulation of NKP. The effects of KCC3 AA induction on NKCC and NKP are abolished when extracellular Na⁺ is replaced with NMDG.

Unexpectedly, DCPIB, a blocker of volume-regulated Cl⁻ channel activity (VRAC, or Iᵥ(Clₐ-well)), and known to inhibit K⁺ loss through intermediate K⁺ (IK, Kᵥₐ₃,₃₉) conductance channels (Lauf et al., 2008) caused partial inhibition of Kᵢ loss during induction (Figure 5). Furthermore, DCPIB inhibited KCC3 AA-induced Cl-independent Kᵢ loss (Figure 6) while inhibiting...
KCC and NKP and activating NKCC (Figure 7). DCPIB, dose-dependently, also increased NKCC and K_i content but inhibited KCC as a function of time when present during the flux (Figure 8).

To further assess the effect of the VRAC inhibitor on the main transporters studied, DCPIB (50 µM) was applied during the 3 main steps of the Rb⁺ influx protocol; i.e., during KCC3 AA induction (I), pre-incubation (P), flux (F), and in different combinations (IP, IF, and IPF) (Figure 9, Supplementary Tables 1, 2). Three main components for K⁺ loss were defined and were calculated as total KCC3 AA-induced K⁺ loss (T), Cl⁻ cotransporter-dependent K⁺ loss (CICT, i.e., through NKCC + KCC, see Figure 10), and K⁺ channel-dependent K⁺ loss (KCH). Figure 9 depicts the total KCC3 AA-induced K⁺ loss (T) through its two components (CIOT and KCH) in the various conditions.

Strikingly, statistically significant inhibition of KCH by DCPIB was observed when the inhibitor was present during induction (I) alone (I + DC in I) (p < 0.05), and complete inhibition when it was present during I + flux (I + DC in IF) (p < 0.005), n = 4 with respect to I alone, although there was not statistically significant difference between groups (See Figure 9 legend for more details). In the latter condition, K_i was maintained at the control level. Similar results were observed when the inhibitor was present during either induction + preincubation (IP), or during induction + preincubation + flux (IPF). These data show KCC3 AA-induced Cl⁻ independent K_i loss is DCPIB-sensitive, suggesting dependence on VRAC activity.

The activities of the Cl-coupled cotransporters (NKCC and KCC) were determined by measuring Rb⁺ influx or K⁺ loss under identical conditions as in Figure 9 and the total Rb⁺ influx or K⁺ loss through NKCC + KCC are represented in Figure 10 as their sum for Rb⁺ influx and for K loss. Note that the sum of the two cotransporters was roughly constant for both Rb⁺ influx and K⁺ loss independently of the increasing exposure to DCPIB, and that K⁺ loss was about twice the Rb⁺ influx across the different conditions. This difference may be explained by the driving forces for K⁺ (∼140 mM K⁺/0 mM K⁺) and Rb₀ (0 mM Rb⁺/10 mM Rb⁺). These findings indicate that constitutive dephosphorylation of KCC3 could lead to cell shrinking when VRAC is inhibited by DCPIB.

**Effect of KCC3 Thr991/Thr1048 Dephosphorylation on Cell Volume Regulation in Hypotonic Osmotic Stress**

Given the potent Cl-dependent and Cl-independent K_i loss induced by KCC3 AA, we speculated these cells might exhibit different swelling properties compared to those expressing KCC3 WT. We therefore examined cell volume changes and RVD in response to hypotonic challenge in KCC3 WT and KCC3 AA cells (see Materials and Methods for details). At baseline, in isotonic conditions, both cell lines exhibited similar cell volumes (Figure 11A). However, hypotonic conditions [HEPES-MEM (150 mOsm/kg H₂O)] elicited a ∼3.2 ± 0.5-fold increase in cell volume (peak values) in KCC3 WT cells, compared

![FIGURE 5 | Constitutive KCC3 dephosphorylation elicits rapid and potent cellular Kᵢ loss, which is time-dependent and partially sensitive to DCPIB. Kᵢ content was measured as described in Materials and Methods during induction with doxycycline (1.0 µg/ml) and in the presence and absence of DCPIB (50 µM), an inhibitor of volume regulated Cl⁻ channels (VRACs) and intermediate K⁺ channels. Black squares, Control, i.e., no doxycycline or DCPIB; red circles, doxycycline alone, and green triangles, doxycycline + DCPIB. Time points where there was a significant change were compared vs. the Control. n = 8 independent determinations. Total number of independent identical experiments N = 2. *p < 0.001; data represent the mean ± SEM values.](Image 314x533 to 541x710)

![FIGURE 6 | Constitutive KCC3 Thr991/Thr1048 dephosphorylation elicits rapid and potent cellular Kᵢ loss which is time-dependent and partially sensitive to DCPIB. Kᵢ content was measured as described in Materials and Methods during induction with doxycycline (1 µg/ml) and in the presence and absence of DCPIB (50 µM). (A) Kᵢ content in Cl⁻-free medium, sulfamate (S) replacement without (SOB, black squares) and with DCPIB (SOBD, red circles). (B) Kᵢ content in Cl⁻-media ± ouabain (0.1 mM) ± bumetanide (10 µM) without (Cl⁻, black squares, red circles, and green triangles, respectively) and with DCPIB (Cl⁻, black squares, red circles, and green triangles, respectively) and with Cl⁻ (Cl₀, red circles, and green triangles, respectively) and with DCPIB (Cl₀B, blue triangles point down, cyan diamonds, and magenta triangles point left, respectively). Mean ± SE, n = 3–6 individual determinations. (A) For SOB vs. SOBD, there was a statistically significant difference between groups as determined by One-Way ANOVA [F(8, 21) = 51.65, p < 0.0005, n = 30], and confirmed by paired t-test, *p ≤ 0.01 at 3 and 6 h and **p ≤ 0.005 at 9 h. (B) For Cl⁻, Cl₀, SOB vs. Cl₀D, Cl₀B, and SOB, respectively, there was a statistically significant difference between groups as determined by Kruskal–Wallis One-Way ANOVA [F(8, 21) = 64.42, 77.64, and 45.44, p < 0.0005, n = 30], and confirmed by paired t-test, *p < 0.05 at 3 h and **p < 0.005 at 6 and 9 h. Data represent the mean ± SEM values. Representative results from 4 similar experiments.](Image 96x589 to 238x709)
to significantly less swelling in KCC3 AA cells [1.2 ± 0.3-fold increase in cell volume (peak values)] (Figures 11A, B). Two phases of cell volume changes were further analyzed in Figures 11C, D. First, KCC3 WT cells showed a sharp swelling increase upon hypotonic challenge (at a rate of 1.8 ± 0.3 % cell volume/min); in contrast, KCC3 AA cells exhibited a significantly slower rate (1.05 ± 0.2 % cell volume/min). Secondly, KCC3 WT cells triggered RVD at a rate of 0.13 ± 0.01 % cell volume/min (Figure 11D) and recovered to ∼50 ± 0.2 % of their original volume in ~20 min; in contrast, KCC3 AA cells exhibited a slower RVD rate (0.04 ± 0.007 % cell volume/min) (Figure 11D). These data suggest that constitutive KCC3 Thr991/Thr1048 dephosphorylation renders cells less prone to acute swelling in hypotonic osmotic stress.

Discussion

Hypotonic cell swelling triggers RVD to maintain cell volume, resulting in K⁺ and Cl⁻ (and obligated water) efflux via the activation of K⁺ channels, VRACs, and KCC3 (Hoffmann and Dunham, 1995). KCC3 Thr991/Thr1048 dephosphorylation accompanies cell swelling (Rinehart et al., 2009; De Los Heros et al., 2014), but the cellular physiology of this event has not been systematically examined. We show constitutive dephosphorylation of KCC3a Thr991/Thr1048, modeled via alanine substitution at the involved residues (preventing phosphorylation), is a potent mechanism of stimulating KCC3a activity, resulting in a rapid (<10 min) and significant (>90%) reduction in K⁺ via both Cl-dependent and Cl-independent (and DCPIB-sensitive) pathways. These effects, along with the associated inhibition of NKCC1 activity and increase in NKP activity, render cells less prone to acute swelling in hypotonic osmotic stress. Our studies with 85Rb⁺ and 39K⁺ under zero-trans conditions used here and developed several decades ago (Adragna et al., 1994; Adragna and Lauf, 1998), are in agreement with previous studies utilizing radioactive 86Rb⁺ under steady state (Rinehart et al., 2009; De Los Heros et al., 2014), and corroborate and extend earlier work demonstrating the potency of KCC3 Thr991/Thr1048 dephosphorylation for transporter activation. Furthermore, and in contrast to previous studies, our findings demonstrating an important role of the KCC3 Thr991/Thr1048 switch as a key regulator of K⁺ content and cell volume in response to hypotonic stress are entirely novel.

In isotonic conditions, KCC3a is robustly phosphorylated at Thr991/Thr1048 and functionally inactive. In these conditions, and consistent with previous findings measuring unidirectional transport with radioactive 86Rb⁺ under steady state (Rinehart et al., 2009; De Los Heros et al., 2014), we found genetic prevention of KCC3 phosphorylation at these sites elicited to a >25-fold increase in transporter activity relative to wild
type KCC3. This constitutive KCC3 activity was associated with an inhibition (and subsequent reversal) of NKCC1 activity. The negative values of NKCC1 activity in tested conditions (Figure 1A) likely represent Rb\(^+\) leak via NKCC1 in the absence of bumetanide, since these conditions resulted in an increase in intracellular Rb\(^+\) (CIO < CIOB). Constitutive KCC3 activity was also associated with increase in the ouabain-sensitive Rb\(^+\) uptake component, consistent with increased NKP activity. Extracellular Na\(^+\) replacement with NMDG eliminated NKP activity (Figure 3), suggesting constitutive KCC3 activity increases cellular Na\(^+\) entry (which maximally activates NKP in Na\(^+\) but not in Na-free medium). These data reveal a close functional association between KCC3a and NKCC1/NKP in the context of KCC3a activation, and suggest that attempts to “specifically” inhibit or activate individual CCCs (e.g., blocking NKCC1 or activating KCC2/3) will have effects on other functionally related transport systems that aim to maintain a steady state in ion content and cell volume.

KCC3 AA induction was also accompanied by a rapid and significant K\(_i\) loss. About 50% of this K\(_i\) loss was recovered in Cl-free medium. Partial inhibition of K\(_i\) loss in sulfamate-containing medium indicated that there are two components to this K\(_i\) loss: a Cl-dependent and a Cl-independent K\(_i\) loss. The Cl-dependent K\(_i\) loss was inhibited by either replacement of extracellular Rb\(^+\) with K\(^+\) or furosemide (2 mM), suggesting a dependence on KCC3 and NKCC1-mediated K\(^+\) efflux. While the Cl-dependent component could thus be explained, the Cl-independent K\(_i\) loss associated with KCC3 activation required further investigation. We attempted to define this component by inhibiting it with extracellular ion replacement studies or different inhibitors of candidate transport pathways. TEA, clofilium, Ruthenium Red, BAPTA, EDTA, RN-1734, and several other inhibitors had no effect on the Cl-independent K\(_i\) loss under the conditions tested (Supplementary Table 3). Because of the large K\(_i\) loss in such a short period of time, a Ca-activated K\(_i\) channel, such as a BK channel, was thought to be likely involved, as seen in human red blood cells and in vascular smooth muscle, coronary arteries and lens epithelial cells. Thus, 2 mM TEA, which selectively inhibits BK channels, was used, in contrast to 10 mM, which is the concentration used for most Ca-sensitive K\(^+\) channels of intermediate or smaller conductance. To test the basic idea that Ca\(^+\) was involved, different approaches, i.e., with inhibitors and chelators, were used, with negative results. Next, clofilium was used as a Ca-independent K\(^+\) channel blocker, but also with no effect. However, it is possible that BAPTA AM/EDTA under the conditions used in this study may not have effectively depleted other internal sources of Ca\(^+\).

However, DCPIB, an inhibitor of VRACs (and also intermediate K\(^+\) conductance channels), completely suppressed the Cl-independent K\(_i\) loss when used during induction + flux conditions. These results show that constitutive KCC3

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FIGURE 9 | Cl-independent K\(_i\) loss triggered by constitutive KCC3 Thr991/Thr1048 dephosphorylation is DCPIB-sensitive. Bars indicate total doxycycline-induced K\(_i\) loss (T, red). K\(_i\) loss through Cl-dependent cotransporters (NKCC + KCC, CICOT, green), and K\(^+\) channel-sensitive K\(_i\) loss (KCH, blue) when DCPIB, an inhibitor of volume regulated Cl\(^-\) channels (VRACs) and intermediate K\(^+\) channels, was applied in the induction (I), preincubation (P), and/or flux (F) stages of the ion flux study. DCPIB inhibits Cl-sensitive K\(_i\) loss. Representative results from a typical experiment done in quadruplicates. Total number of similar experiments, N = 4. For T, there was a statistically significant difference between groups as determined by One-Way ANOVA (F\(_{1,24}\) = 5.16, p < 0.001, n = 32) or by Kruskal–Wallis One-Way ANOVA (F\(_{1,24}\) = 4.46, p < 0.005, n = 32). There was a statistically significant difference between groups as determined by Kruskal–Wallis One-Way ANOVA (F\(_{1,27}\) = 12.01, p < 0.0005, n = 36) for CICOT but not for KCH (F\(_{1,24}\) = 1.41, p > 0.2, n = 32. Data represent the mean ± SEM values. However, there was a statistically significant difference for KCH, *p < 0.05 and #p < 0.005; and for TD (p < 0.0005) by paired T-Test for the different conditions shown in the figure and described in the text.

FIGURE 10 | Cl-dependent Rb\(^+\) influx and K\(_i\) loss triggered by constitutive KCC3 Thr991/Thr1048 dephosphorylation is DCPIB-sensitive. Bars indicate Rb\(^+\) influx through Cl-dependent cotransporters (NKCC + KCC, cyan) and K\(_i\) loss (NKCC + KCC, magenta), when DCPIB was applied in the induction (I), preincubation (P), and/or flux (F) stages of the ion flux study. DCPIB inhibits Cl-dependent Rb\(^+\) influx and K\(_i\) loss depending on the conditions. Representative results from 4 similar experiments. For Rb\(^+\) (NKCC + KCC), there was a statistically significant difference between groups as determined by One-Way ANOVA (F\(_{1,27}\) = 48.75, p < 0.0005, n = 36). For K\(^+\) (NKK + KCC), there was a statistically significant difference between groups as determined by Kruskal–Wallis One-Way ANOVA (F\(_{1,27}\) = 20.51, p < 0.0005, n = 36). Data represent the mean ± SEM values.


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K\(_i\) loss was inhibited by either replacement of extracellular Rb\(^+\) with K\(^+\) or furosemide (2 mM), suggesting a dependence on KCC3 and NKCC1-mediated K\(^+\) efflux. While the Cl-dependent component could thus be explained, the Cl-independent K\(_i\) loss associated with KCC3 activation required further investigation. We attempted to define this component by inhibiting it with extracellular ion replacement studies or different inhibitors of candidate transport pathways. TEA, clofilium, Ruthenium Red, BAPTA, EDTA, RN-1734, and several other inhibitors had no effect on the Cl-independent K\(_i\) loss under the conditions tested (Supplementary Table 3). Because of the large K\(_i\) loss in such a short period of time, a Ca-activated K\(_i\) channel, such as a BK channel, was thought to be likely involved, as seen in human red blood cells and in vascular smooth muscle, coronary arteries and lens epithelial cells. Thus, 2 mM TEA, which selectively inhibits BK channels, was used, in contrast to 10 mM, which is the concentration used for most Ca-sensitive K\(^+\) channels of intermediate or smaller conductance. To test the basic idea that Ca\(^+\) was involved, different approaches, i.e., with inhibitors and chelators, were used, with negative results. Next, clofilium was used as a Ca-independent K\(^+\) channel blocker, but also with no effect. However, it is possible that BAPTA AM/EDTA under the conditions used in this study may not have effectively depleted other internal sources of Ca\(^+\).

However, DCPIB, an inhibitor of VRACs (and also intermediate K\(^+\) conductance channels), completely suppressed the Cl-independent K\(_i\) loss when used during induction + flux conditions. These results show that constitutive KCC3
activation elicits a rapid and pronounced decrease in $K_i$ via $Cl$-dependent/CCC-mediated pathways, as well as $Cl$-independent pathways possibly dependent on VRAC activity, such as $I_{Cl\text{swell}}$ (Voss et al., 2014), or possibly via a yet-unidentified $K^+$ conductance that is not inhibited by either TEA or clofilium. In any event, the results obtained with DCPIB seem to shed light on the mechanism by which $K^+$ is lost independently of $Cl^-$ when the KCC3 AA mutant is overexpressed. Identifying this $K^+$ transport pathway will be an interesting topic of detailed future investigation.

The mechanisms by which over-expression of KCC3 AA, resulting in constitutively active $K^+-Cl^-$ cotransport, leads to the associated changes in NKCC and NKP, and $K_i$ and cell volume, are summarized in the model in Supplementary Figures 3, 4. Supplementary Figure 3 shows the potential mechanisms accounting for our experimental findings during KCC3 AA induction with doxycycline and preincubation. These experiments were performed in the presence of 5 mM extracellular $K^+$ ($K_o$). During these time intervals, KCC3 AA induction (step 1) would trigger KCC-mediated outwardly-directed $K^+$ and $Cl^-$ loss, and consequently, cell shrinkage. This would initially stimulate NKCC1 (step 3), increasing intracellular $Na^+$ and $Cl^-$ (step 4). The increase in intracellular $Na^+$ would stimulate NKP, bringing more $K^+$ into cells. The increase in intracellular $Na^+$ and $Cl^-$ (and associated influx of osmotically-obliged water) would then lead to a “rebound” transient cell swelling and membrane depolarization (step 5), activating DCPIB-sensitive VRACs (i.e., Lrrc8-dependent $I_{Cl\text{swell}}$) (step 6), and indirectly, inducing $K_i$ loss (step 7). During the flux procedure (Supplementary Figure 4), which measures $Rb^{+}$ influx through KCC, NKCC and NKP in the presence of 10 mM $Rb$, and $K_i$ content at 0 $K_o$, the nominally infinite outward gradient for $K^+$ causes an outward efflux of $K^+$ and $Cl^-$ through KCC (step 1), and enhanced entry of $Rb^{+}$ through KCC, NKCC and NKP (step 2). $Rb^{+}$ will enter up to a certain level after which back-flux ensues (Sachs, 1967), exiting through NKCC1 (i.e., in a reversal of activity) and blocking its further entry through inwardly directed NKCC1 (step 3). Treatment with DCPIB during induction alone, or induction and pre-incubation, and afterwards during flux, would likely directly inhibit $I_{Cl\text{swell}}$ and indirectly inhibit $Cl$-sensitive $K^+$ loss, as well as the aforementioned pathways for $Rb^{+}$ and $K^+$ transport (KCC and NKP), whereas its effect on NKCC1...
would appear to be inhibitory or stimulatory depending on the conditions (Figures 6–8).

We examined cell volume changes and RVD in response to acute hypotonic challenge in KCC3 WT and KCC3 AA cells. In these experiments, KCC3 AA cells exhibited significantly less acute cell swelling than KCC3 WT cells in hypotonic conditions. However, KCC3 WT cells exhibited a mildly brisker RVD rate than KCC3 AA cells. These data suggest that constitutive KCC3 activity, achieved via tonic Thr991/Thr1048 dephosphorylation, might render cells less prone to cell swelling in response to hypotonic osmotic stress—particularly during the early phases of physiological perturbation. It remains to be determined whether the reduced acute cell swelling in KCC3 AA cells in these conditions is due solely from increased KCC3-mediated ionic efflux, or from the combined effects of this and other volume-regulated transport proteins.

Multiple diseases exhibit dysregulation of cell volume due to aberrant transmembrane ionic transport (Epstein et al., 1995). For example, sickle cell vaso-occlusive events caused by sickle cell dehydration have been associated with hyperactive KCC-mediated cellular K\(^+\) and Cl\(^-\) efflux, suggesting KCC inhibition might be a pharmacotherapeutic strategy to prevent sickle cell dehydration and HB\(S\) polymerization (Brugnara, 2001). This could be achieved by promoting the inhibitory phosphorylation of KCC3 at Thr991/Thr1048 via PP1/PP2a phosphatase inhibition. Conversely, ischemic and traumatic cerebral edema is associated with glial cell swelling due in part to NKCC1-dependent Na\(^+\), K\(^+\), and Cl\(^-\) intracellular accumulation (Su et al., 2002; Lenart et al., 2004; Gagnon et al., 2007). In this context, stimulating KCC3 by antagonizing Thr991/Thr1048 phosphorylation would be expected to not only facilitate K\(^+\) and Cl\(^-\) efflux and cell shrinkage, but also secondarily decrease the pathological elevation in NKCC1 activity (see Figures 2, 4). The necessity for KCC3-mediated cell volume regulation in the human nervous system has been demonstrated by the autosomal recessive human disease Andermann syndrome, caused by loss-of-function mutations in KCC3, which results in axonal swelling, periaxial fluid accumulation, and neurodegeneration due to defective RVD (Boettger et al., 2003; Byun and Delpire, 2007).

In elucidating the normal physiological functions of the key phospho-regulatory sites of KCC3, our study suggests that antagonizing the inhibitory phosphorylation of KCC3 at Thr991/Thr1048 would activate KCC3, decrease cellular K\(^+\) content, and resist acute cell swelling. This could be achieved by inhibiting the WNK-SPAK protein kinase complex (Alessi et al., 2014). SPAK phosphorylates Thr1048 directly, but not Thr991 (Rinehart et al., 2009; De Los Heros et al., 2014), and WNK1 is necessary for both Thr991 and Thr1048 phosphorylation (De Los Heros et al., 2014). Inhibiting the WNK-SPAK pathway would have the added benefit of concurrently decreasing the stimulatory phosphorylation of NKCC1 at Thr212/Thr217, sites required for NKCC1 activity (Thastrup et al., 2012) and pathologically up-regulated in ischemic edema (O’Donnell, 2014). This powerful coincident reduction in NKCC1-mediated ion influx and stimulation of KCC3-mediated ion efflux would seem ideally suited to counter acute pathological cell swelling, particularly in glia, in hypotonic swelling conditions as seen in acute hyponatremia from water intoxication. Activation of this KCC3 switch might also be of use in isosmotic swelling, which results from activity-dependent transporter and channel-mediated increases in cellular ion load (Kahle et al., 2009; Kaila et al., 2014), as in ischemic stroke. Critical to these considerations will be a determination of how constitutive KCC3 Thr991/Thr1048 dephosphorylation might affect long-term cell viability, and CNS and PNS structure and function, in the CNS in vivo. We are currently evaluating this in knock-in KCC3 mice carrying double-alanine mutation at Thr991/Thr1048. Lastly, dephosphorylation of the homologous residues in KCC2 (Thr906/Thr1107) might also be a novel method of stimulating KCC2 in diseased neurons to promote Cl\(^-\) extrusion and restore GABAergic inhibition (Kahle et al., 2013).

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Supplementary Material

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