SNAP23 regulates KCC2 membrane insertion and activity following mZnR/GPR39 activation in hippocampal neurons

Highlights
- Neuronal K⁺/Cl⁻ cotransporter 2 (KCC2) is regulated via interaction with SNAP23.
- Zn⁡²⁺ enhances interaction and membrane insertion of SNAP23, Syntaxin 1A, and KCC2.
- Zn⁡²⁺-dependent mZnR/GPR39 regulation of KCC2 requires SNAP23 phosphorylation.
- Epithelial KCC3 regulation by ZnR/GPR39 also requires SNAP23.
SNAP23 regulates KCC2 membrane insertion and activity following mZnR/GPR39 activation in hippocampal neurons

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SUMMARY

Modulation of the neuronal K⁺/Cl⁻ cotransporter 2 (KCC2) activity, which mediates Cl⁻ export, is critical to neuronal function. Here, we demonstrate that KCC2 interacts with the SNARE protein synaptosome-associated protein 23, SNAP23, an essential component of membrane insertion machinery. Using KCC2 truncated mutants, we show that KCC2 C-terminal domain is essential for membrane targeting and SNAP23-dependent upregulation of KCC2 activity triggered by activation of the Zn²⁺-sensitive receptor mZnR/GPR39 in HEK293 cells. Expression of SNAP23 phosphorylation-insensitive mutants or inhibition of its upstream activator IκB kinase (IKK) prevents mZnR/GPR39 upregulation of KCC2 activity in mouse hippocampal neurons. We further find that SNAP23 interacts with Syntaxin 1A and KCC2, and that all three proteins exhibit increased membrane insertion following mZnR/GPR39 activation in neurons. Our results elucidate a G-protein-coupled receptor-dependent pathway for regulation of KCC activity, mediated via interaction with SNARE proteins.

INTRODUCTION

The K⁺/Cl⁻ cotransporter KCC2 is responsible for Cl⁻ extrusion in mature neurons and thus maintains the inward Cl⁻ gradients essential for generation of hyperpolarizing GABA<sub>A</sub> and glycine responses (Kahle et al., 2016; Mapplebeck et al., 2019; Watanabe et al., 2019). Activity of KCC2 is highly regulated by synaptic signaling, thereby affecting Cl⁻ extrusion capability and neuronal inhibitory tone (Mahadevan and Woodin, 2016; Spoljaric et al., 2019). Regulation of KCC2 is achieved by rapid changes in the phosphorylation status of multiple residues on KCC2, which, in turn, can directly modulate its activity (Cordshagen et al., 2018; Weber et al., 2014), or influence its presence on the plasma membrane (Côme et al., 2019a; Markkanen et al., 2017; Medina et al., 2014). Most prominent, phosphorylation of serine 940 (S940) by protein kinase C (PKC), a pathway activated by multiple G-protein coupled receptors, rapidly increases KCC2 surface stability and thereby Cl⁻ transport (Lee et al., 2007; Leonzino et al., 2016). In contrast, surface expression of KCC2 is reduced by phosphorylation of threonine 1007 and 906, which results in a depolarized GABA<sub>A</sub> reversal potential (Friedel et al., 2015; Heubl et al., 2017). Phosphorylation of these threonine residues is mediated via activation of the with-no-lysine kinase (WNK) and its downstream target STE20/SPS1-related proline/alanine-rich kinase (SPAK) (Zhang et al., 2020). Other putative phosphorylation sites have been identified on both the N- and C-terminal domains of KCC2 and were shown to be selectively activated by various triggers in cells exogenously expressing the transporter (Cordshagen et al., 2018). In neurons, the membrane stability of KCC2 has been associated with modulation of the C-terminus while the N-terminus was suggested to control its delivery into the plasma membrane and is particularly important for its expression in distal dendrites (Friedel et al., 2017).

Zinc ions (Zn²⁺) that are synaptically released from neuronal presynaptic mossy fibers activate a postsynaptic metabotropic Gq-protein coupled receptor, mZnR/GPR39, triggering intracellular Ca²⁺ signaling (Besser et al., 2009). Subsequent activation of the mitogen-activated kinase (MAPK) pathway increases KCC2 surface expression in a soluble N-ethylmaleimide-sensitive factor attachment protein receptor (SNARE)-dependent manner (Chorin et al., 2011; Saadi et al., 2012). Further studies have shown that mZnR/GPR39 knockout (KO) mice exhibit enhanced kainate-induced seizures, compared to wild-type mice, indicating that regulation of KCC2 activity by mZnR/GPR39 plays an important homeostatic role that can dampen epileptic activity (Gilad et al., 2015). The precise mechanism linking mZnR/GPR39 activation...
Figure 1. KCC2 C-terminal is essential for Zn\(^{2+}\)-dependent upregulation of transport
(A) Immunoblot analysis of KCC2, ΔNTD-KCC2, or ΔCTD-KCC2 expression in HEK293 cells. Blots were exposed to a KCC2 antibody together with an anti-FLAG antibody, and normalized to actin as loading control. The KCC2 antibody, which is targeted to the C-terminus, works well with both the WT and ΔNTD
Figure 1. Continued

KCC2. The FLAG antibody recognizes the WT KCC2 and ΔCTD truncated mutant but works poorly with the ΔNTD. White arrows on the blot point to the monomeric forms of each of the constructs at their expected molecular weight. Right panel shows quantification of expression level, and a schematic drawing of the KCC2 constructs used. (p = 0.0008, Welch’s t test)

(B) HEK293 cells co-transfected with GPR39 and KCC2 or KCC2 alone were loaded with the pH-sensitive fluorescent indicator BCECF (1 μM). Cells were washed with K+-free Ringer’s solution, baseline, and addition of NH4Cl (15 mM) resulted in increase in fluorescence due to NH3 diffusion. Subsequent NH4+ transport, used as surrogate by K+ transporter, resulted in acidification (Hershfinkel et al., 2009). Shown are representative fluorescent signal traces from cells treated with Zn2+ (200 μM, 2 min) compared to control cells or cells treated with the KCC inhibitor DIOA (100 μM). Right panel shows averaged rates of NH4+ transport mediated by KCC2, obtained over initial 100 s from maximal signal. (**p < 0.0001, t test).

(C) BCECF fluorescent signal traces from cells co-transfected with GPR39 and ΔNTD-KCC2 or ΔCTD-KCC2, treated with or without Zn2+. Averaged rates of NH4+ transport, representing KCC2 activity, are shown in right panel. (**p = 0.0032, t test). Dotted line indicates the rate of transport in cells expressing GPR39 and WT KCC2 that were treated with Zn2+ as in (B).

(D) Biotinylation assay of KCC2 variants in HEK293 cells co-transfected with GPR39 and KCC2 or ΔNTD-KCC2 or ΔCTD-KCC2 (upper panel). Comparison of surface expression between cells treated with Zn2+ (200 μM, 2 min) and control cells, using KCC2 and FLAG antibodies (as in A). Loading controls for surface expression were the membrane protein transferrin receptor and cytoplasmic tubulin. Total protein levels (lower panel) determined in cell lysates were used for normalization. Tubulin was used as loading control. The arrows mark the protein isoform, quantified in the right panel, at the expected molecular weight. Quantification of normalized membrane expression of KCC2 variants in Zn2+-treated cells versus control (**p = 0.0064 KCC2 with and without Zn2+, **p = 0.0012 ΔNTD-KCC2 with and without Zn2+, Mann-Whitney test). All bar graphs values are averages ± SD, with all data points presented.

with KCC2 function, however, has not been fully delineated. Several other pathways that activate metabolic neuronal signaling via G-protein coupled receptors also upregulate KCC2 activity (Mahadevan and Woodin, 2016). Among them, mGluR1 and mGluR5 (Banke and Gegelashvili, 2008), 5-HT2A serotonin receptors (Bos et al., 2013), the oxytocin receptor (Leonzino et al., 2016), and kainate receptors (Garand et al., 2019) all activate Ca2+-dependent PKC signaling that directly enhances KCC2 S940 phosphorylation and, subsequently KCC2 surface expression. We show here that the G-protein coupled receptor mZnR/GPR39 activates a distinct pathway that enhances interaction between the SNARE protein SNAP23 and KCC2 to regulate the membrane expression and activity of the transporter.

RESULTS

KCC2 C-terminal domain is essential for regulation by mZnR/GPR39 signaling

We first asked if mZnR/GPR39 regulation of KCC2 activity is mediated via its N-terminal domain (NTD) or the C-terminal domain (CTD). For this purpose, we used HEK293 cells co-transfected with mZnR/GPR39 and the truncated isoforms ΔNTD-KCC2 or ΔCTD-KCC2, lacking, respectively, the NTD and CTD (Friedel et al., 2017). Transfection of HEK293 cells with the ΔNTD-KCC2 (deletion of amino acids 1–100) resulted in low, albeit measurable expression of the transporter (Figure 1A), while ΔCTD-KCC2 (deletion of amino acids 654–1114) yielded similar expression levels when compared to wild-type (WT) KCC2 (Figure 1A), in agreement with previous results (Li et al., 2007). To study KCC2-dependent transport, we determined rates of cellular acidification with BCECF fluorescence using established methodology (Chorin et al., 2011; Hershfinkel et al., 2009). Application of NH4Cl to the extracellular solution induces cellular alkalization due to passive entry of NH3. This is followed by pronounced KCC2-dependent transport that acts in reverse mode and mediates influx of NH4+, a surrogate to K+, leading to intracellular acidification and markedly decreased BCECF fluorescence (Figure 1B). We monitored acidification rates in control cells treated with 200 μM Zn2+ for 2 min, a condition that activates mZnR/GPR39 without affecting intracellular Zn2+ levels (Besser et al., 2009). As previously noted (Chorin et al., 2011), cells treated with Zn2+ showed increased rates of NH4+ transport when compared to acidification rates in vehicle-treated controls (Figure 1B). Treatment of cells with the KCC2 inhibitor DIOA (100 μM) prevented Zn2+-dependent upregulation of transport activity, confirming that KCC2 mediates the NH4+-dependent acidification. Zn2+ treatment of HEK293 cells expressing KCC2 without mZnR/GPR39 did not influence the rates of NH4+-dependent acidification, indicating that mZnR/GPR39 is essential for enhancing KCC2-dependent transport by Zn2+ (Figure 1B). Next, we measured NH4+ transport rates in cells expressing mZnR/GPR39 together with the truncated ΔNTD-KCC2 or ΔCTD-KCC2 variants. We noted slower baseline rates of NH4+ transport activity in cells expressing ΔNTD-KCC2 (Figure 1C), when compared to the full-length KCC2 (Figure 1B), consistent with the lower expression levels of this variant. Nonetheless, NH4+ transport rates in cells expressing ΔNTD-KCC2 were still significantly enhanced following Zn2+ treatment (Figure 1C), suggesting that the N-terminus is not necessary for transporter regulation by mZnR/GPR39. In contrast, transport rates in ΔCTD-KCC2-expressing cells were completely insensitive to Zn2+ (Figure 1C), indicating that the C-terminal domain is required for mZnR/GPR39-mediated regulation of KCC2 activity.
mZnR/GPR39-dependent upregulation of KCC2 activity is not mediated through phosphorylation sites on the C-terminus of the transporter

The results presented thus far indicate that mZnR/GPR39-dependent regulation of KCC2 transport activity requires the C-terminus, which contains numerous regulatory phosphorylation sites (Côme et al., 2019b). As mentioned earlier, various Gq-coupled metabotropic receptors upregulate KCC2 activity via PKC phosphorylation of S940 (Banke and Gegelashvili, 2008; Lee et al., 2007; Leonzino et al., 2016; Mahadevan and Woodin, 2016). We thus asked whether phosphorylation of S940 (Figure 2A) is also required for mZnR/GPR39-dependent upregulation of KCC2. We mutated S940 into a phosphorylation-insensitive alanine residue (S940A), and determined NH4+ transport rate in HEK293 cells co-expressing this mutant and mZnR/GPR39 (Figure 2B). The S940A KCC2 mutant showed Zn2+-dependent upregulation of transport activity similar to the WT KCC2 (Figure 2B). Because other residues on KCC2 may be phosphorylation targets of PKC, we tested whether kinase inhibitors staurosporine (1 nM) and GO6983 (10 μM) could block the modulatory actions of Zn2+ on transporter function. However, cells expressing WT KCC2 and mZnR/GPR39 exhibited similar Zn2+-dependent upregulation of KCC2 activity in the presence of both inhibitors (Figure 2C). Taken together, these results suggest that mZnR/GPR39 upregulation of KCC2 activity does not involve PKC signaling or phosphorylation of KCC2 at S940.

We then explored the possible role of threonines 906 and 1007, previously shown to enhance KCC2 surface expression following dephosphorylation (Friedel et al., 2015). We transfected HEK293 cells with KCC2 mutants that mimic the constitutively phosphorylated state at these sites, T906E or T1007E (Figure 2A), and monitored Zn2+-dependent NH4+ transport rates. In cells expressing either T906E or T1007E, which could not be dephosphorylated to enhance KCC2 membrane expression levels, we still observed Zn2+-dependent upregulation of KCC2 activity (Figure 2D). This suggests that T906 or T1007 are also not involved in mZnR/GPR39 regulation of KCC2.

Previous results showed that similar to neuronal KCC2, the epithelial KCC family members are also activated by mZnR/GPR39 via Ca2+ signaling and activation of MAP kinase (Mero et al., 2019; Sunuwar et al., 2017). To determine if this kinase phosphorylated a previously unknown site on KCC2, we identified putative MAPK phosphorylation sites that are shared between KCC family members using the Group-based Prediction System (GPS) Website (Xue et al., 2011) (http://gps.biocuckoo.cn/online.php). Putative phosphorylation sites receiving a high score were mutated into alanine, namely S988A, S986A, T584, and S868A (Figure 2A). In addition, we deleted a region that is enriched with serine residues (SPVSS, residues 1045–1049) also suspected to be phosphorylated by MAPK. However, we found that all these mutants
Figure 3. SNAP23 phosphorylation is required for ZnR/GPR39-dependent KCC3 and KCC2 upregulation

(A) SHSY-5Y cells were transfected with an siRNA sequence targeted to silence GPR39, or with a scrambled sequence (SCR) to determine the role of mZnR/GPR39 in upregulating transport in these cells. Insert shows mRNA expression level of GPR39 in SCR and siGPR39 cells. NH₄⁺ transport was monitored in...
continued to show Zn\textsuperscript{2+}-dependent upregulation of KCC2 transport activity (Figure 2E). Notably, we found that the S988A mutant showed an increase in baseline transport activity, suggesting that dephosphorylation of S988 may be involved in regulation of KCC2 that is independent of mZnR/GPR39, but mutation of both S988 and S986 to alanine reversed this baseline effect. Importantly, both mutants showed Zn\textsuperscript{2+}-dependent upregulation of transport activity. Moreover, mutation of both S986 and 988 into negatively charged aspartate residues also yielded Zn\textsuperscript{2+}-dependent upregulation of transport, further suggesting that the phosphorylation or dephosphorylation of these sites are not linked to mZnR/GPR39 regulation of KCC2. In addition, mutation of T584 to a phosphorylation-insensitive alanine resulted in an increased effect of Zn\textsuperscript{2+} on transport activity, suggesting that following Zn\textsuperscript{2+}-dependent insertion of KCC2 into the membrane, dephosphorylation of this site may enhance membrane stability of the transporter, as shown for T906 and T1007 (Friedel et al., 2015). Thus, although MAPK inhibition attenuates mZnR/GPR39 activation of endogenous KCC2, mZnR/GPR39 regulation of KCC2 is mediated by SNAP23 (Chorin et al., 2011), we could not identify a direct phosphorylation site of KCC2 that is required for mZnR/GPR39 regulation of transport.

mZnR/GPR39 regulation of KCC2 is mediated by SNAP23

In light of the aforementioned results, we next sought other potential mechanisms that could be responsible for mediating mZnR/GPR39-dependent upregulation of KCC2. In a prior study, we reported that SNARE proteins are essential for regulation of neuronal KCC2 function following mZnR/GPR39 activation (Saadi et al., 2012). In a separate line of experiments, we found that mZnR/GPR39 regulates the epithelial KCC isoforms KCC1 in colon cells and KCC3 in breast cancer cells (Chakraborty et al., 2021; Mero et al., 2019; Sunuwar et al., 2017). As the epithelial SNARE protein SNAP23 regulates surface expression of major ion transporters, including the cystic fibrosis transmembrane conductance regulator (CFTR) and epithelial sodium channel (ENaC) (Cormet-Boyaka et al., 2002; Saxena et al., 2006), we asked if SNAP23 could mediate the enhanced neuronal KCC2 activity following mZnR/GPR39 activation. For this purpose, we used SHSY-5Y neuroblastoma cells, which endogenously express KCC2 and mZnR/GPR39 (Chorin et al., 2011), thus do not require overexpression that can promote promiscuous protein–protein interactions. We first confirmed that mZnR/GPR39 activation enhanced KCC-dependent transport in these cells using mZnR/GPR39-silencing (siGPR39) constructs. We found that Zn\textsuperscript{2+} treatment upregulated NH\textsubscript{4}\textsuperscript{+} transport in cells transfected with a scrambled siRNA (Figure 3A). In contrast, siGPR39 that effectively reduced mRNA expression of mZnR/GPR39 (inset, Figure 3A) abolished the stimulatory actions of Zn\textsuperscript{2+} (Figure 3A). To determine if KCC2 mediates the observed transport, we applied the KCC inhibitor DIOA or the KCC2 selective inhibitor VU0463271. Each of these inhibitors effectively diminished Zn\textsuperscript{2+}-dependent upregulation of NH\textsubscript{4}\textsuperscript{+} transport (Figure S1). We noted, however, that inhibition of KCC2 by VU0463271 showed enhanced baseline transport activity, a potential compensatory mechanism mediated by the sodium/potassium/chloride co-transporter NKCC1. We therefore asked if NKCC may be regulated by Zn\textsuperscript{2+}. But, the NKCC inhibitor bumetanide did not reverse the effect of Zn\textsuperscript{2+} (Figure 3B), indicating that NKCC is not upregulated by mZnR/GPR39 in SHSY-5Y cells. Finally, we found that silencing of KCC2 using siRNA constructs reduced endogenous KCC2 mRNA (inset, Figure 3B) and prevented mZnR/GPR39 upregulation of NH\textsubscript{4}\textsuperscript{+} transport activity (Figure 3B). These results suggest that SHSY-5Y cells can effectively serve as a model system to investigate the pathway for mZnR/GPR39 activation of endogenous KCC2.
**A** Primary hippocampal neurons

- BAY11-7085 (5 μM) + Zn²⁺
- Control
- BAY11-7085 (5 μM)
- Zn²⁺

**B** Primary hippocampal neurons

- NH₄Cl
- SCR peptide
- SNAP23 peptide
- SNAP23 peptide + Zn²⁺
- SCR peptide + Zn²⁺

**C** Control

**D** IP: Anti SNAP23

| WB: Anti KCC2 | kDa | Control | Zn²⁺ | IgG |
|---------------|-----|---------|------|-----|
| Anti SNAP23   | 25  |         |      |     |
| Anti Syntaxin 1A | 35  |         |      |     |

**E** kDa - + Zn²⁺

- Membrane fraction
- Total proteins

- Anti KCC2
- 140 kDa
- Membrane fraction
- Total proteins

- Anti SNAP23
- 25 kDa
- Membrane fraction
- Total proteins

- Anti Syntaxin 1A
- 35 kDa
- Membrane fraction
- Total proteins

Membrane expression following Zn²⁺ (normalized to total)

- KCC2
- SNAP23
- Syntaxin 1A
- No Zn²⁺
Next, we silenced SNAP23 in SHSY-5Y cells (mRNA expression shown in inset, Figure 3C) and monitored rates of neuronal KC2-dependent transport activity. Remarkably, Zn²⁺ failed to enhance the rate of NH₄⁺ transport in SHSY-5Y cells transfected with siSNAP23 (Figure 3C), while expression of a scrambled siRNA construct showed Zn²⁺-dependent increase in transport activity (Figure 3A). This suggests that SNAP23 is required for mZnR/GPR39-dependent modulation of endogenous KC2 activity. Similar siSNAP23 silencing in HEK293 cells expressing mZnR/GPR39 and the epithelial KC3, showed reversal of Zn²⁺-dependent upregulation of NH₄⁺ transport (Figure S2). This finding suggests a general role for SNAP23 in mZnR/GPR39-dependent regulation of KC2 transporters. In mast cells, phosphorylation of SNAP23 by IκB kinase (IKK) induces granule membrane fusion and exocytosis (Naskar and Puri, 2017; Suzuki and Verma, 2008). We therefore investigated whether Zn²⁺-dependent upregulation of NH₄⁺ transport in SHSY-5Y cells is modulated by inhibition of IKK. Pretreatment of SHSY-5Y cells with the IKK inhibitor BAY11-7085 (2 μM) abolished mZnR/GPR39 enhancement of NH₄⁺ transport (Figure 3C), suggesting that IKK phosphorylation of SNAP23 may be a critical component of KC2 regulation by mZnR/GPR39. As IKK phosphorylates serine 95 and serine 120 of SNAP23 to regulate its activity (Hepp et al., 2005), we next tested if a phosphorylation-insensitive mutant of the SNARE protein, in which both serine 95 and serine 120 had been replaced by alanine (595/120A), would diminish mZnR/GPR39 activation of KC2. Because SHSY-5Y cells express an endogenous SNAP23, we used cells in which the endogenous human SNAP23 isoform was silenced using our siRNA construct (Figure 3C). We then transfected the cells with a mouse wild-type SNAP23 (WT-mSNAP23) or a mouse 595/120A mutant SNAP23, which are insensitive to the human isoform-directed siRNA. In SHSY-5Y cells silenced for the human SNAP23, but expressing the mouse SNAP23 isoform (WT-mSNAP23), we observed Zn²⁺-dependent upregulation of acidification rates (Figure 4A). In BAY11-7085 (5 μM)-treated neurons, however, Zn²⁺ did not enhance rates of cellular acidification (Figure 4A), suggesting that the pathway under study is also involved in mZnR/GPR39 regulation of KC2 in this preparation. To specifically determine the role of SNAP23, hippocampal neurons were incubated with a cell-permeable blocking peptide that mimics the N-terminal domain of SNAP23 that is critical for mZnR/GPR39 enhances SNAP23 binding to KC2 to enhance its surface expression in hippocampal neurons

To assess the role of SNAP23 in the modulation of KC2 in a more relevant model, mouse hippocampal neuronal cultures were used. In agreement with our previous results (Chorin et al., 2011; Saadi et al., 2012), Zn²⁺ enhanced NH₄⁺ transport, monitored by cellular acidification of BCECF-loaded neurons (Figure 4A). In BAY11-7085 (5 μM)-treated neurons, however, Zn²⁺ did not enhance rates of cellular acidification (Figure 4A), suggesting that the pathway under study is also involved in mZnR/GPR39 regulation of KC2 in this preparation. To specifically determine the role of SNAP23, hippocampal neurons were incubated with a cell-permeable blocking peptide that mimics the N-terminal domain of SNAP23 that is critical for...
activation of this SNARE protein (Cheng et al., 2013). We found significant Zn²⁺-dependent upregulation of KCC2 activity in cells incubated with a scrambled peptide, used as control (Figure 4B). In contrast, the SNAP23 blocking peptide completely abolished upregulation of KCC2 activity by Zn²⁺ (Figure 4B). Altogether, these results support a role for IKK and SNAP23 in mZnR/GPR39-mediated regulation of KCC2 in primary neurons.

We then asked if Zn²⁺ treatment could promote interaction between SNAP23 and KCC2 in neurons. We first studied the interaction between KCC2 and SNAP23 using a proximity ligation assay (PLA) (Krall et al., 2020; Söderberg et al., 2006), which allows detection of fluorescent puncta when target proteins are at least within 40 nm from each other. We found that in hippocampal neurons in culture Zn²⁺ treatment, which activated mZnR/GPR39 (Chorin et al., 2011), led to a significant increase in fluorescent PLA puncta (Figure 4C). The Zn²⁺-triggered enhanced proximity likely resulted from increased interaction between KCC2 and SNAP23. The interaction between these two proteins was further tested by immunoblot analysis of membrane protein fractions from hippocampal tissue slices immunoprecipitated with SNAP23 antibodies. Indeed, KCC2 immunoreactivity was observed in control and Zn²⁺-pretreated cells (Figure 4D), indicating that SNAP23 and KCC2 are found in a protein complex. We also found that syntaxin 1A, previously shown to interact with SNAP23 (Cormet-Boyaka et al., 2002; Saxena et al., 2006), was present in the SNAP23-immunoprecipitated fraction that contained KCC2 (Figure 4D), suggesting all three of these proteins are found within the complex. While co-IP cannot provide an accurate quantitative measure of expression levels, due to differences in antibody pull-down efficiency, we did note that in both control and Zn²⁺-treated tissue, expression of KCC2 and syntaxin 1A is seen in the SNAP23-immunoprecipitated fractions (Figure 4D). To determine if Zn²⁺-pretreatment enhances membrane insertion of SNAP23 and syntaxin 1A, as we have shown previously for KCC2 (Chorin et al., 2011), we studied the expression level of these proteins in membrane-enriched fractions. Immunoblot analysis of KCC2, SNAP23, and syntaxin 1A in non-immunoprecipitated membrane-enriched fractions from hippocampal tissue slices indicated that expression of all three proteins was indeed increased in Zn²⁺-treated tissues compared to control (Figure 4E). Thus, our results indicate that in neurons, Zn²⁺ treatment enhances the interaction between SNAP23, syntaxin 1A, and KCC2, as well as their expression in membrane-enriched fractions.

DISCUSSION

Mechanisms for modulation of KCC2 transport activity, directly or via its trafficking to the cell membrane, have been the focus of numerous studies due to the major role of this transporter in influencing neuronal excitability (Mahadevan and Woodin, 2016). Phosphorylation of KCC2 is considered a fundamental feature of the function of this transporter, required for rapid regulation of its membrane insertion and stability (Côme et al., 2019a). In the current study, we identify a previously unrecognized molecular component that regulates KCC2 membrane insertion via a transporter phosphorylation-independent pathway, namely SNAP23. We show that the interaction of KCC2 with the SNARE proteins SNAP23 and syntaxin 1A is essential for the mZnR/GPR39-dependent upregulation of KCC2 activity.

Modulation of neuronal inhibitory tone by the Gq-coupled mZnR/GPR39 is mediated via increased membrane expression of KCC2 (Chorin et al., 2011; Gilad et al., 2015). Signaling activated by mZnR/GPR39 involves release of Ca²⁺ from intracellular stores, similar to the cellular signaling cascade activated by the metabotropic glutamate receptor (mGLUR1) (Banke and Gegelashvili, 2008). Previous studies demonstrated that metabotropic Gq-coupled receptors, among them mGLUR1, mGLUR5, or 5-hydroxytryptamine type 2A serotonin receptors (5-HT2Rs), regulate KCC2 activity via PKC-dependent phosphorylation of serine 940 (S940) (Banke and Gegelashvili, 2008; Bos et al., 2013; Garand et al., 2019; Lee et al., 2007; Mahadevan and Woodin, 2016). Therefore, it seemed likely that mZnR/GPR39 would activate a similar pathway to regulate KCC2 activity (Mahadevan and Woodin, 2016). Surprisingly, mZnR/GPR39-dependent upregulation of transport activity was maintained in the presence of the phosphorylation-insensitive S940A KCC2 mutant or PKC inhibitors. Moreover, other major phosphorylation sites on KCC2 were similarly not involved in mZnR/GPR39 regulation of KCC2 transport. In contrast, we find that mZnR/GPR39 activation of IKK and SNAP23 are required for increased transport activity. Thus, despite shared Ca²⁺ signaling by these Gq-coupled receptors, mZnR/GPR39 and the mGLURs activate distinct, possibly context-dependent signaling pathways that can regulate KCC2 activity via distinct phosphorylation-independent and dependent components. It thus would be interesting to determine if these pathways can synergistically enhance KCC2 activity, especially since Zn²⁺ and glutamate co-exist in synaptic vesicles in a large number of excitatory synapses (Krall et al., 2021; McAllister and Dyck, 2017).
In a previous study, we demonstrated that mZnR/GPR39 upregulation of KCC2 activity in neurons is prevented by a botulinum toxin C1-expressing plasmid (Saadi et al., 2012). This toxin cleaves SNARE proteins SNAP25 and syntaxin 1A, but not SNAP23 (Gu et al., 2016; Pantano and Montecucco, 2014; Saadi et al., 2012). Our present results show that KCC2 interacts with SNAP23 and syntaxin 1A in a protein complex, suggesting that our previous observations can be explained by the cleavage for syntaxin 1A, preventing from complexing with SNAP23 and KCC2. In neurons, expression of SNAP25 might be more prominent in presynaptic membranes, where it regulates synaptic vesicle trafficking and membrane fusion to induce neurotransmitter release (Rizo and Südhof, 2012; Tao-Cheng et al., 2000; Zamponi, 2003). In contrast, SNAP23 is found in postsynaptic dendritic spines where it regulates surface expression of NMDA (Suh et al., 2010; Washbourne et al., 2004) and GABA \\
 receptors (Gu et al., 2016), via its interaction with receptor-containing vesicles. Interestingly, SNAP23 was also shown to directly interact with mGluR1 and regulate its membrane expression (Raynaud et al., 2018). Our results point to another role for SNAP23, namely, regulation of KCC2 surface expression following activation of mZnR/GPR39, likely by recruiting Syntaxin 1A. Moreover, our results also show that mZnR/GPR39 signaling activates SNAP23-dependent up-regulation of transport mediated by the epithelial KCC3 isoform. Thus, we suggest that SNAP23 interaction with KCC transporters may be a ubiquitous pathway for regulation of this family of transporters.

Various signaling pathways induce SNAP23-dependent vesicle release, mediated by the phosphorylation of the SNARE protein at distinct residues. For example, LPS-dependent cytokines release from mast cells is triggered by Toll-like TLR4 receptor via intracellular Ca²⁺ rise and IκB kinase (IKK) activation, which, in turn, induces SNAP23 phosphorylation at serines 120 and 95 (Suzuki and Verma, 2008). Astroglia signaling via a Ca²⁺-dependent pathway induced a similar pattern of SNAP23 phosphorylation important for exocytosis (Yasuda et al., 2011). Our results show that mZnR/GPR39 signaling, via IKK pathway and phosphorylation of serine 120 and serine 95 of SNAP23, enhances KCC2 membrane expression. In contrast, the histamine H1 receptor in HeLa cells regulates exosome secretion via release of Ca²⁺ and activation of PKC, leading to phosphorylation of SNAP23 serine 110 (Verweij et al., 2018). Exosome release from cancer cells is triggered by pyruvate kinase type M2 (PKM2) phosphorylation of serine 95 on SNAP23, enhancing metastasis and invasion of the cancer cells through the extracellular matrix (Wei et al., 2017). Hence, distinct upstream events may differentially regulate SNAP23 phosphorylation on sites that induce unique protein–protein interactions and control exocytosis of vesicles or specific proteins.

Previous studies suggested that KCC2 membrane insertion is determined by the N-terminal domain under baseline expression conditions and that mutants lacking the C-terminus are more rapidly internalized (Friedel et al., 2017). While the surface expression of the N-terminal truncated protein was previously shown to be ambiguous in HEK-293 cells (Friedel et al., 2017; Li et al., 2007), our results show that mZnR/GPR39 regulation of KCC2 is maintained in the absence of the N-terminus. Although baseline activity of the ΔNTD-KCC2 mutant was low, activation of mZnR/GPR39 increased its membrane localization and facilitated an increase in transport activity. In contrast, we show that the C-terminal is necessary for SNAP23-mediated membrane insertion. Thus, in addition to the known phosphorylation sites that regulate KCC2 membrane stability (Friedel et al., 2017), the C-terminus of KCC2 likely also has a binding site for SNAP23 that controls its membrane insertion. Interestingly, previous results have revealed an interaction of the KCC2 C-terminus with the actin cytoskeleton, leading to enhanced neuronal differentiation and migration (Horn et al., 2010).

Taken together, our results indicate that SNAP23 interacts with KCC2 to enhance its surface expression, a pathway critical for modulating inhibitory postsynaptic responses in hippocampal neurons.

Limitations of the study
We found that the C-terminal domain of KCC2 is essential for membrane insertion of the transporter by SNAP23. Future studies should identify the precise interaction domains on KCC2 (Krall et al., 2020; Raynaud et al., 2018), which will enable selective targeting of this interaction to control neuronal inhibitory tone. In addition, previous studies found that KCC2 regulation is mediated by phosphorylation, which can be triggered by Gq-coupled receptors signaling. It would be interesting to determine the role of SNAP23 activation by other GPCRs.

STAR* METHODS
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ACKNOWLEDGMENTS
We thank Dr. Igor Medina for critical reading of the manuscript and his advice and support throughout this project. This work was funded by a joint National Science Foundation-US-Israel Binational Science Foundation grant NSF-IOS-BSF 1655480 (MH, EA). MH is also funded by Israel Science Foundation grant 812/20.

AUTHOR CONTRIBUTIONS
H.A and M.H designed experiments, H. A., N.G., and M.B. performed experiments and analyzed data, H.A., I.S., E. A., and M.H. interpreted the data, wrote and edited the manuscript.

DECLARATION OF INTERESTS
The authors declare no competing interests.

Received: September 20, 2021
Revised: November 25, 2021
Accepted: January 6, 2022
Published: February 18, 2022

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**KEY RESOURCES TABLE**

| REAGENT | SOURCE | IDENTIFIER |
|---------|--------|------------|
| **Antibodies** | | |
| Anti SNAP23 | Synaptic System | 111–213, RRID:AB_10805651 |
| Anti SNAP23 | Synaptic System | 111–202, RRID:AB_887788 |
| Anti KCC2 (D1R2R) | Cell Signaling Technology | Y4725, RRID:AB_2800232 |
| Anti-KCC2 [N1/12] | Abcam | Cat# ab134300 |
| Anti Syntaxin 1A | Synaptic Systems | 110–111, RRID:AB_887848 |
| Anti FLAG | Genscript | A 00187, RRID:AB_1720813 |
| Anti Transferrin receptor | Thermofisher scientific | 13–6800, RRID:AB_2533029 |
| β-Actin | MP Biomedicals | Cat# 0869100 |
| β-Tubulin | Santa Cruz Biotechnology | sc5274, RRID:AB_2288090 |
| **Chemicals, Peptides and recombinant proteins** | | |
| Dulbecco’s Modified Eagle’s Medium (DMEM) | Biological Industries | 01-055-1A |
| Fetal Calf Serum | Biological Industries | 04-001-1A |
| pen-strep | Biological Industries | 03-031-1B |
| L-Glutamine | Biological Industries | 03-020-1B |
| Zinc sulfate heptahydrate | Sigma- Aldrich | z4750 |
| BCECF-AM | Tef Labs | Not available |
| BSA | MP Biomedicals | 216006980 |
| TransIT-X2 | mirusbio | MIR 6000 |
| DioA | Sigma-Aldrich | D129 |
| 2-[(2-butyl)-6,7-dichloro-2-cyclopentyl-1-oxo-3H-inden-5-yl]oxy]acetic acid | Sigma-Aldrich | B3023 |
| Bumetanide | Sigma-Aldrich | G1918 |
| Staurosporine | Santa Cruz Biotechnology | sc3510A |
| Go6983 | Sigma-Aldrich | 4719 |
| BAY11-7085 (E)-3-(4-tert-butylphenyl)sulfonylprop-2-enenitrile | APEXBio | B3033 |
| VU0463271 | Tocris | 4719 |
| Iodoacetamide | Sigma-Aldrich | I6125 |
| 2-iodoacetamidine | Sigma-Aldrich | K1007 |
| Protease inhibitor cocktail | Worthington | LS033126 |
| Papain | Worthington | 02-015-18 |

(Continued on next page)
Continued

| REAGENT                                                                 | SOURCE                          | IDENTIFIER |
|------------------------------------------------------------------------|---------------------------------|------------|
| HEPES                                                                  | Biological Industries           | 03-025-1C  |
| 2-[(4-(2-hydroxyethyl)piperazin-1-yl)ethanesulfonic acid               |                                 |            |
| poly-D-lysine                                                           | Sigma-Aldrich                   | p0899      |
| Neurobasal medium                                                       | GIBCO                           | 21103049   |
| Fetal Bovine Serum                                                     | HyClone                         | sh30070.03 |
| B-27                                                                  | GIBCO                           | 17504044   |
| Glutamax-100X                                                          | GIBCO                           | 35050038   |
| Gentamicin                                                             | Biological Industries           | 03-035-1C  |
| Veri blot IP detection reagent                                        | Abcam                           | ab131366, RRID:AB_2892718 |
| protein A magbeads                                                     | GenScript                       | L00273     |
| qPCR bio probe blue mix                                                | PCR biosystem                   | pb20.25-05 |
| Cytosine β-D-arabinofuranoside (AraC)                                  | Sigma- Aldrich                  | c6645      |
| SNAP23 blocking peptide                                                | GRKRRQRRRRPQMDOLSPEEIQLRAHQVDSEQ| GenScript USA Inc |
| SCR blocking peptide                                                   | GRKRRQRRRRPQFAESLFQSIKESGFSHGCG| GenScript USA Inc |

**Critical commercial assays**

| Bio-Rad protein assay                                                  | Bio-Rad                          | 50000006   |
| QS Site-Directed Mutagenesis Kit                                      | New England Biolabs              | E05545     |
| EZ-Link Sulfosuccinimide-NHS-Biotinylation Kit                         | Pierce Biotechnology, Thermoscientific | PIR-89881 |
| Dualink in situ red starter kit                                       | SigmaAldrich                     | DU092101   |
| PureLink™ RNA Mini Kit                                                | Invitrogen                       | 12183018a  |
| cDNA synthesis kit                                                     | Quanta bio                       | 95047      |

**Experimental models: Cell lines**

| HEK293                                                                | ATCC                            | CRL-1573   |
| SHSY-5Y                                                               | ATCC                            | CRL-2266   |

**Experimental models: Organisms/strains**

| C57BL/6jRccHsd mice                                                   | Envigo                          | 043        |

**Oligonucleotides**

| siGPR39                                                               | 5’-CCAUGGAGUUUCUACAGCAU-3'       | Sigma- Aldrich |
| siSNAP23                                                             | 5’-CAACACAGAAGCUAUUGAAG-3'       | Integrated DNA Technologies, IDT |
| siKCC2                                                               | 5’-CAACAGAUGCAUUUACCAAGAT-3'     | Integrated DNA Technologies, IDT |
| SCR                                                                  | 5’-GCCCAGAUCCCGUGACGU-3’         | Sigma- Aldrich |
| KCC2 S940A                                                           | 5’-GTCAGCCCGCAATCCGGAG-3’        | 5’-TCCTCTGTGATACCTCTGGTGGTCTC-3’ |
| KCC2 S988A                                                           | 5’-CAGCTCCCGGCCCCGAGGGGA-3’      | 5’-CTGGGCGAGCTGGAGACCTC-3’       |
| KCC2 S986A                                                           | 5’-CCCAGCGCGCCCGCTCCC-3’         | 5’-CAAGTTGGAAGACTCTGATGCTGG-3’   |
| KCC2 T584A                                                           | 5’-GTCAGCTCGGAGCGACTCTGGATG-3’   | 5’-GTCAGCGAGGACCTCTGGATC-3’      |
| KCC2 T906E                                                           | 5’-CTATCGAGGCTGTTGTGGAGG-3’      | 5’-GTGAAACCTGAGATGCTG-3’         |
| KCC2 T1007E                                                          | 5’-GTCATCTGACTCGGTGCAGAAGG-3’    | 5’-TTCCTGGATCTGCTTCC-3’          |
| KCC2 S868A                                                           | 5’-GAGATGACATCCATCTCAGAAGG-3’    | 5’-ATCTGGGGCCACGTGAG-3’          |
| SPV5S                                                                | 5’-GAGGGCGTACAGAGGCTT-3’         | 5’-GGGGCCCTATTCTTC-3’            |
| ΔNCD-KCC2                                                            | 5’-ATGGGCGTGATCCCTGAGC-3’        | 5’-TTTACATCATCATCATTTTATATACCTGACT-3’ |
| ΔCTD-KCC2                                                            | 5’-TGCTCTAGAGCATCATATC          | CCATGCACCACACTCC-3’             |

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**RESOURCE AVAILABILITY**

**Lead contact**
Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact: Michal Hershfinkel (hmichal@bgu.ac.il)

**Materials availability**
Reagents and plasmids generated in this study are available from the lead contact upon request.

**Data and code availability**
All data needed to evaluate the conclusions in the paper are present in the paper and/or the supplemental information. All data reported in this paper will be shared by the lead contact upon request.

This paper does not report original code.

**EXPERIMENTAL MODEL AND SUBJECT DETAILS**

**Cell cultures**
HEK293 and SHSY-5Y cell lines, purchased from ATCC and not authenticated. Cells were grown in Dulbecco’s Modified Eagle’s Medium (DMEM, Biological Industries, Israel) supplemented with 10% (v/v) fetal calf serum (Biological Industries, Israel), 1% pen-strep, 1% L-Glutamine solution. All cells were grown in 5% CO2 humidified atmosphere at 37°C.

**Primary mouse neuronal cultures**
Primary hippocampal neurons were prepared from postnatal day 0 C57BL/6JRccHsd mouse neonates of both sex, using a protocol approved by the Institucional Animal Care and Use Committee of the Ben-Gurion University of the Negev. Hippocampi were treated with papain (Worthington, Lakewood, NJ) and dissociated by mechanic pipetting in Hanks’ balanced salts solution (HBSS) supplied with 20mM HEPES (pH 7.4). Cells were seeded on 50 μg poly-D-lysine-coated coverslips (13 mm) in 24-well plates at the density of 5 × 10⁶ cells/mL (coverslip). During initial 24h cells were cultured in neurobasal medium (GIBCO), supplemented with 5% fetal bovine serum (FBS), 2% B-27, 1% Glutamax-100X, and 1 μg/mL Gentamicin, after which the medium was changed to serum-free medium. To suppress glial growth, 5 μM Cytosine β-D-arabinofuranoside (AraC) was added to the medium on DIV5. All cells were cultured in a humidified atmosphere of 5% CO2 at 37°C.

**Mouse tissues for co-IP**
C57BL/6JRccHsd male mice of both sex (18 days) were sacrificed, hippocampal slices were sectioned (Chorin et al., 2011), and incubated at 37°C in O2 bubbled ACSF buffer (124 mM NaCl, 26 mM NaHCO3, 1.25 mM NaH2PO4, 2 mM MgSO4, 2 mM CaCl2, 3 mM KCl, and 10 mM glucose, at pH 7.4), for 30 min.

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### REAGENT SOURCE IDENTIFIER

| REAGENT            | SOURCE                      | IDENTIFIER                                                                 |
|--------------------|-----------------------------|----------------------------------------------------------------------------|
| SNAP23 S95/120A    | 5'-GAACCTTTGAGCTGGAAGAAGAC-3' | 5'-TTTGCCTATTACAAGGG-3'                                                   |
|                    | 5'-TAAGCAACCCGCCGATTACAAATG-3' | 5'-GATACTACATTGCTAGGTG-3'                                                 |

| Reagent Source DNA | Source                          | Identifier                                                                 |
|--------------------|---------------------------------|---------------------------------------------------------------------------|
| Human KCC2 cDNA    | Prof. Gamba                     |                                                                           |
| Mouse SNAP23       | OriGene                         | MR202271                                                                  |
| Human GPR39        | OriGene with mcherry addition and deletion of FLAG-MYC | www.origene.com/catalog/cdna-clones/expression-plasmids/rc219212/gpr39-nm_001508-human-tagged-orf-clone |
METHOD DETAILS

Cell culture preparation and transfection

For imaging experiments, cells were seeded at 1 x 10^6 cells/mL on 10 mm glass coverslips, and used after 24–48 h. For siGPR39, siKCC2, siSNAP23 silencing, HEK293 or SHSY-5Y cells were seeded in 60 mm plates, 24 h prior to transfection. Cells were transfected with siRNA constructs (Sigma- Aldrich, Israel, and Integrated DNA Technologies, IDT, IOWA, USA) using the CaPO4 precipitation protocol or Mirus transit-X2, according to the manufacturer’s protocol. Sequences of siRNA used for GPR39 silencing were: 5'-CCAU GGAUUCUACGCAU-3', for SNAP23: 5'-CCAACAGAAGGAUAUUGA-3', for KCC2 5'-CAAACAGA CAGAUACAAAT-3' and the control siRNA (scrambled) sequence: 5'-GGCGAUGCCCUGU ACGU-3'. At 48 h after transfection, cells were used for imaging experiments or immunoblots.

Inhibitors

DIOA, #D129 SigmaAldrich. Bumetanide, #B3023 Sigma Aldrich. Staurosporine, #sc3510A Santa Cruz Biotechnology. Go6983, #G1918 SigmaAldrich. BAY11-7085, #B3033 APExBio. VU0463271, #4719 Tocris. All inhibitors were applied in Ringer’s solution at concentrations and incubation times shown in the relevant figure legend.

Molecular tools

The human KCC2 cDNA coding plasmid was a kind gift from Prof. Gamba (Song et al., 2002). The mouse SNAP23 plasmid was purchased from OriGene (OriGene Technologies Inc. MR202271). All site-directed mutagenesis was performed using Q5 Site-Directed Mutagenesis Kit, according to the manufacturer’s protocols (New England Biolabs). Primers used for generation of mutations were as follows: KCC2 point mutations: Ser940 to Ala (S940A; 5'-GTCACGAGGCGCAATCCGGAG-3', 5'-TCATCTGTGATACTCTGGATCTC-3'); Ser988 to Ala (S988A; 5'-CAGCTCCCCGGCCCCAGGGGA-3', 5'-CTGGGGCAGCTGGGAAGCACTC-3'); Ser986 to Ala (S986A; 5'-CCCCAGCAAGCCCGCTCCCCC-3', 5'-CAGCTGGGAGCACTCTGATG-3'); Thr584 to Ala (T584A; 5'-GCTGCTGAGGGCACCCAACTG-3', 5'-GTCTGCACTGCGAGGCC-3'); Thr906 to Glu (T906E; 5'-CTATGAGAAGGAGTTGG TGATGGAG-3', 5'-GTGTAAGCTGAGACTGTCG-3'); Thr1007 to Glu (T1007E; 5'-GGTGCATCTCGAGTGGACCAAGGACAAG-3', 5'-TTCTCCGGA TCTGTCTCC-3'); Ser868 to Ala (S868A; 5'-GGATGACAATGCCATCCAGATGAAGAAG-3', 5'-ATCTGGGCCACAGTGGAAG-3'), SPVSS deletion (SPVSS; 5'-GAGGGCATCAAGGACTTC-3', 5'-GGGGCCCTTTAACAACCGATCCGGAGCCAGATC-3'). Deletion of 1–100 amino acids ΔNCD-KCC2; 5'-ATGCGCTGTTGAGGATCTGG-3', 5'-TTTACTTCATTCTCT-3', Ser95 to Ala (594/120A; 5'-GAACCTGAGCTGGAAAAGAAAC-3', 5'-TTTGTCTCTATTCAAGG-3', 5'-TAAGCAACCGCAGAGGTATCAATATTG-3', 5'-GATACTACATTTCTAGGTG-3'). All mutations were verified using sequence analysis.

qPCR assay

Silencing efficiency of SNAP, GPR39 or KCC2 in SHSY-5Y cells was quantified using qPCR. Cells were seeded on 60 mm plates, 24 h prior to transfection. Cells were transfected with siRNA constructs, and after 48 h the total RNA was purified using RNeasy Mini Kit (Invitrogen) as described by the manufacturer. 1 μg RNA was converted to cDNA using cDNA synthesis kit as described by the manufacturer (Quanta bio). cDNA was diluted 1:10 (as per calibration) with stop codon and Xba restriction enzyme site at the C-terminal domain (ΔCTD-KCC2; 5'-TGCTCTAGAGCATCATATCCCATCGCCCCACTCC-3'), ΔTTA TCATCATCATCTTTTATAACCATGCCCCCATTT-3', Alfa and Ser120 to Ala (594/120A; 5'-GAACCTGAGCTGGAAAAGAAAC-3', 5'-TTTGTCTCTATTCAAGG-3', 5'-TAAGCAACCGCAGAGGTATCAATATTG-3', 5'-GATACTACATTTCTAGGTG-3'). All mutations were verified using sequence analysis.
Fluorescent imaging

For measurement of KCC-dependent transport, we used the NH4Cl paradigm (Chorin et al., 2011) to cells loaded with a pH-sensitive dye, 2′,7′-Bis-(2-Carboxyethyl)-5-(and-6)-Carboxyfluorescein (BCECF). Cells grown on coverslips were loaded for 15 min at room temperature with 1 μM BCECF-AM (Tef Labs) in Ringer’s solution containing 0.1% BSA and were subsequently incubated in Ringer’s solution containing 0.1% BSA for additional 15 min at room temperature. Inhibitors were added in some experiments to the incubating solution, as described in figure legends. For SNAP23 inhibition, a cell permeant blocking peptide for the N-terminus (SNAP23: GRKKRRQRRRPQMDDLSPEEIQLRA HQVTDSEQ or SCR: GRKKRRQRRRP QGFAESLFOQIESGFCG) was designed to include a trans-activator of transcription (TAT) cell-penetrating peptide (YGRKKRRQRRR) (Cheng et al., 2013; Krall et al., 2020). Primary cultured neurons were incubated with the peptide, or the scrambled control, for 6 h prior to BCECF loading for imaging. Treatment with extracellular Zn2+ for mZnR/GPR39 activation was done by addition of 200 μM Zn2+ for 2 min prior to imaging. Coverslips were then mounted in a perfusion chamber and BCECF was excited at 440 and 470 nm and imaged with a 510-nm long-pass filter. Briefly, the NH4Cl paradigm consists of perfusion of the cells with K+-free Ringer’s solution containing NH4Cl (15 mM, HEK293 or 5 mM, SHSY-5Y and Hippocampal neurons primary culture). This triggers initial alkalization that is induced by NH3 diffusion, resulting in an increase in fluorescence signal. Subsequently, activation of ion transport systems induces transport of NH4+, which is a surrogate to K+, and results in cellular acidification that is representing NH4+ efflux rate. The rate of acidification was determined using a linear fit obtained during a 100-s period following the peak fluorescent signal.

The imaging system consisted of an Axiovert 100 inverted microscope using a 20 objective (Zeiss, Germany), Polychrome V monochromator (TILL Photonics, Germany) and a SensiCam cooled charge-coupled device (PCO, Germany). Fluorescent imaging measurements were acquired with Imaging Workbench 5 (Indec, USA).

Protein expression and co-immunoprecipitation

To monitor changes in KCC2 membrane expression HEK293 cells were seeded in 10 cm plates and cotransfected with 1 μg of the GPR39-mcherry and 8 μg of the either the WT-KCC2, ΔCTD-KCC2 or ΔNTD-KCC2 plasmids using the TransIT-X2 Transfection Reagent (Mirus). Cell surface biotinylation assay was performed using the EZ-Link Sulfo-NHS-Biotinylation Kit (Pierce Biotechnology, Thermoscientific). Cells were treated with 200 μM Zn2+ for 2 min then incubated with sulfo-NHS-Biotin for 30 min at 4°C, cells were then lysed (50 mM HEPES, pH 7.4, 150 mM NaCl, 1 mM EGTA, 1 mM EDTA, 10% glycerol, 1% Triton X-, 100 mM MgCl2, 10 mM iodoacetamide and protease inhibitor cocktail, A2S) 250 μg lysates were then incubated with Neutravidin Gel (Pierce) overnight at 4°C. Sample were resolved on 8–15% gradient SDS-PAGE followed by immunoblot analysis.

Membrane expression and interaction of KCC2, SNAP23 and Syntaxin-1A were studied using membrane fractions of hippocampal brain section. C57BL/6JRccHsd male mice (18 days) were sacrificed, hippocampal slices were sectioned as described (Chorin et al., 2011), and incubated at 37°C in O2 bubbled ACSF buffer (124 mM NaCl, 26 mM NaHCO3, 1.25 mM NaH2PO4, 2 mM MgSO4, 2 mM CaCl2, 3 mM KCl, and 10 mM glucose, at pH 7.4), for 30 min. Slices were then washed with phosphate-free ACSF, followed by 200 μM Zn2+ treatment for 1 min and hippocampi were isolated and lysed in fractionation buffer (20 mM HEPES, pH 7.4, 10 mM KCl, 2 mM MgCl2, 1 mM EDTA, 1 mM EGTA with protease inhibitors), lysates were centrifuged at 720 g for 5 min. Supernatant, containing total cell lysates, was used to measure the total cell protein. Protein concentrations were determined using Bio-Rad protein assay only for the total cell protein lysate. The supernatant was further ultracentrifuged at 140,000 g for 1 h, the pellet that contains membrane proteins was resuspended in fractionation buffer and used to evaluate membrane protein levels with immunoblots. Briefly, SDS sample buffer was added, and samples were heated to 40°C for 5 min. Proteins were separated on gradient gel 8–15% SDS-PAGE and blotted onto nitrocellulose membranes.

In addition, the membrane-enriched fractions of the hippocampal slices, described above, was further used for co-immunoprecipitation. Membrane fractions were incubated overnight at 4°C with an antibody against the SNAP23 (111–213, Synaptic System), then subjected to protein A magbeads (L00273, Genescript) for 1 h at room temperature. Beads were washed and proteins were isolated from the beads with sample buffer containing beta-mercaptoethanol. Proteins were separated on gradient gel 8–15% SDS-PAGE and blotted onto nitrocellulose membranes.
Proteins were detected using antibodies raised against KCC2 (94725, Cell Signaling Technology), SNAP23 (111–202, Synaptic Systems), Syntaxin 1A (110–111, Synaptic Systems) and FLAG (A00187, Genscript), transferrin receptor (13–6800, Thermofisher scientific), β-Actin (0869100, MP), β-Tubulin (sc5274, Santa Cruz Biotechnology). For quantification of KCC2 expression levels the monomeric form of the protein was quantified from immunoblots and normalized as mentioned in the figure legends. Note that KCC2 oligomers may be associated with other proteins and in ectopic expression system their levels may depend on the preparation (Medina et al., 2014). To minimized detection of light and heavy chains of the IgG we used the Veriblot IP detection reagent (ab131366, Abcam).

**Proximity ligation assay**

PLA assay was performed using on PFA fixed neuronal cultures that were treated with Zn\(^{2+}\) (100 μM, 30 s) or without it as control, and the Duolink in situ red starter kit (DU092101, SigmaAldrich) was applied as per the manufacturer protocol, using antibodies against KCC2 (ab134300, Abcam) and against SNAP23 (111–213, Synaptic Systems). Coverslips seeded with cells were treated with Zn\(^{2+}\) to activate mZnR/GPR39, and then PFA 4% for 10 min, rinsed in PBS (PBS), and then permeabilized with 0.1% Triton X-in PBS. Coverslips were incubated in Dualink secondary antibodies (anti-rabbit SNAP23 and anti-mouse KCC2), which were conjugated with complementary oligonucleotides and PLA probes, and fluorescently labeled. Fluorescent images were acquired with appropriate filters using Nikon Eclipse Ti confocal microscope using an X60 objective. PLA puncta were quantified by counting the fluorescent particles using ImageJ software module in each field of view. PLA experiments were performed on three independent cultures, and n refers to the number of images used for quantification.

**QUANTIFICATION AND STATISTICAL ANALYSIS**

All imaging traces were generated and fitted, using a linear curve fit, in Microsoft Excel. Rates of KCC2 transport were determined by the decrease in fluorescence signal during a 100s period following the peak fluorescent signal, all acquired rates are presented in bar graphs with mean ± SD (SD). Primary neurons imaging data are presented as boxplots, with min and max, showing median values and all data points. All imaging experiments were done on at least three independent cultures. For western blots, quantification was done using ImageJ module for blot analysis (Gels) and data from all gels are presented in bar graphs with mean ± SD or boxplots, min to max, with median value. PLA puncta were quantified by counting the fluorescent particles using ImageJ software module in each field of view. PLA experiments were performed on three independent cultures, and n refers to the number of images used for quantification.

All data are expressed as means ± SD in graphs generates by Microsoft Excel. Statistical significance between the groups was determined following Shapiro-Wilk normality test (GraphPad Prism 8.4.2), using the Student’s t test or ANOVA with multiple comparisons, as appropriate (SPSS). In addition, Mann-Whitney or Independent-Samples Kruskal-Wallis Test, as appropriate, were used for analysis of non-parametric data (GraphPad Prism 7.4.2). The sample sizes for each experiment (n) is shown in all graphs and extent of significance is included in figure legends. *p<0.05; **, p<0.01; ***, p<0.001.