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Depolarizing GABA contributes to glutamatergic network rewiring in epilepsy

Running head: GABA fosters glutamatergic rewiring

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Abstract

Objective: Rewiring of excitatory glutamatergic neuronal circuits is a major abnormality in epilepsy. Besides the rewiring of excitatory circuit, an abnormal depolarizing GABAergic drive has been hypothesized to participate in the epileptogenic processes. However, a remaining clinically relevant question is whether early post Status Epilepticus (SE) evoked chloride dysregulation is important for the remodeling of aberrant glutamatergic neuronal circuit.

Methods: Osmotic mini-pumps were used to infuse intracerebrally a specific inhibitor of depolarizing GABAergic transmission as well as a functionally blocking antibody towards the pan-neurotrophin receptor p75 (p75\textsuperscript{NTR}). The compounds were infused between 2 and 5 days after pilocarpine-induced SE. Immunohistochemistry for NKCC1, KCC2 and ectopic recurrent mossy fiber (rMF) sprouting as well as telemetric EEG and electrophysiological recordings were performed at day 5 and 2 months post-SE.

Results: Blockade of NKCC1 after SE with the specific inhibitor bumetanide restored NKCC1 and KCC2 expression, normalized chloride homeostasis and significantly reduced the glutamatergic recurrent mossy fiber sprouting (rMF) within the dentate gyrus. This mechanism partially involves p75\textsuperscript{NTR} signaling as bumetanide application reduced SE-induced p75\textsuperscript{NTR} expression and functional blockade of p75\textsuperscript{NTR} decreased rMF sprouting. The early transient (3days) post-SE infusion of bumetanide reduced rMF sprouting and recurrent seizures in the chronic epileptic phase.

Interpretation: Our findings show that early post-SE abnormal depolarizing GABA and p75\textsuperscript{NTR} signalings foster a long lasting rearrangement of glutamatergic network that contributes to the epileptogenic process. This finding defines promising and novel targets to constrain reactive glutamatergic network rewiring in adult epilepsy.

Keywords: mossy fiber sprouting, epilepsy, pan-neurotrophin receptor p75\textsuperscript{NTR}, GABAergic signaling, NKCC1, KCC2, bumetanide.

Abbreviations: TLE: temporal lobe epilepsy; SE: status epilepticus; DG: dentate gyrus; DGC: dentate gyrus granule cell; rMF: recurrent mossy fiber; p75\textsuperscript{NTR}: pan neurotrophin receptor p75; KCC2: K\textsuperscript{+}-Cl\textsuperscript{-} co-transporter; NKCC1: Na\textsuperscript{+}-K\textsuperscript{+}-2Cl\textsuperscript{-} co-transporter.
Introduction

Plastic remodeling of glutamatergic neuronal networks is one of the major pathophysiological processes observed in epileptic tissue and post-traumatic brain injuries (TBI) both in humans and animal models\(^1,2\). The understanding of the molecular mechanisms involved in this plastic remodeling is of particular importance since it creates a powerful hyperexcitable cerebral focus that drives recurrent disabling seizures\(^3,4\). This phenomenon, called reactive plasticity, is well documented in the dentate gyrus (DG), where ectopic dentate granule cell (DGC) axons (the recurrent mossy fibers [rMFs]) sprout and create a reverberant excitatory circuit between DGCs. This is involved in the formation of epileptiform bursts in patients with temporal lobe epilepsy (TLE) and TBI, as well as in animal models\(^6-13\). Although the molecular mechanisms are unclear, there is evidence supporting an active role of neurotrophin signaling\(^15-18\). However, the early post-SE molecular events that activate neurotrophin pathways and trigger the ectopic rMF growth remain elusive. Besides the rewiring of excitatory circuits, an abnormal depolarizing GABAergic drive has been hypothesized to participate in epileptogenic processes in adult brain. This includes the dysregulation of the neuronal specific K\(^{+}\)-Cl\(^{-}\) co-transporter KCC2 and the Na\(^{+}\)-K\(^{+}\)-2Cl\(^{-}\) co-transporter NKCC1 as reported in resected tissues from human TLE patients\(^19,20\) and animal models\(^21-23\). Indeed, bumetanide, a specific blocker of NKCC1, has been shown to have acute anticonvulsant action on chronic epileptic activity\(^25\). We have also previously shown that depolarizing GABAergic transmission triggers the upregulation of the pan-neurotrophin receptor, p75\(^{NTR}\), in an acute injury model\(^24\).

Here we have tested the hypothesis whereby the early alteration of chloride homeostasis following SE is a precipitating event involved in rMF sprouting establishment via the activation of p75\(^{NTR}\). We examined a novel therapeutic strategy based on the transitory blockade of NKCC1, early after SE (i.e. before the appearance of ictal activities), to reduce the ectopic sprouting and recurrent seizures in the chronic epileptic phase (i.e. several months after SE).

We found that a functional establishment of ectopic rMF sprouting takes place concomitantly with altered expression of NKCC1 and KCC2 as early as day 5 post-SE in the rat pilocarpine model of TLE. Moreover, we were able to demonstrate that p75\(^{NTR}\) is upregulated and is involved in the formation of ectopic rMF sprouting. In turn, NKCC1 inhibition resulted in the decrease of post-SE p75\(^{NTR}\) upregulation and the reduction of rMF sprouting. Finally, transitory blockade of NKCC1 few days after SE resulted in a significant reduction of rMF growth and recurrent seizures at the chronic stage.

In conclusion, our findings pave a way to identify novel therapeutic strategies targeting cation-chloride co-transporters and p75\(^{NTR}\) signaling pathways to constrain abnormal glutamatergic network rewiring and chronic adult epilepsy.

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Experimental procedures

Animals. All experiments were approved by the Institute National de la Santé et de la Recherche Médicale (INSERM) through the local ethic committee (project number 66-15112012) and the European community council directive (2010/63/UE). Adult male Wistar rats had access to food and water ad libitum and were housed under a 12 h light/dark cycle at 22–24°C.

Rat model of temporal lobe epilepsy. An animal model of temporal lobe epilepsy was obtained using adult male Wistar, as previously described\textsuperscript{26}. Briefly, six weeks old male rats (Charles River and Janvier Laboratories) were treated with pilocarpine hydrochloride (360 mg/kg, i.p.; Sigma-Aldrich) 30 minutes after scopolamine methyl nitrate (0,5 mg/kg, i.p.; Sigma-Aldrich). After 3h of Status Epilepticus (Stage V on Racine’s scale), convulsions were suppressed with diazepam (8 mg/kg, i.p.; Roche) and saline solution (0,9% NaCl; Sigma-Aldrich) was administrated subcutaneously. The success rate of the pilocarpine sessions was 67%, which is in accordance with other studies using adult rats, similar dose, and SE duration\textsuperscript{27-28}.

Bumetanide and p75\textsubscript{NTR} antibody infusion. Osmotic minipumps and cannulae were implanted to continuously deliver bumetanide (B3023, Sigma-Aldrich). Rats were anaesthetized with 4.5% isoflurane and maintained with 2.5% isoflurane for surgery and placed on a heating pad to control body temperature. Rats were placed in a stereotaxic apparatus, a scalp incision was made, and a 0,8 mm diameter hole was drilled through the skull (A/P -3.8; L/M +2.5; D/V 3.0) and a 3.0 mm long 28 gauge cannula (Alzet brain infusion kit, Charles River) was inserted and secured to the skull with dental cement (Paladur, Hereaus). An osmotic mini-pump (Alzet mini-pumps 2002, Charles River) and tubing connected to the cannula were implanted subcutaneously over the back. Mini-pumps were filled with vehicle solution containing 0.3% of ethanol in 1M PBS with or without bumetanide (20μM). After surgery, rats were kept warm, received saline (0.9% NaCl, s.c.), and analgesic buprenorphine (0.03 mg/kg, s.c.) was administrated to minimize animal pain after surgery. In all experiments, infusions began at two days post-status Epilepticus and last for 3 days with a delivery rate of 0.5μl/hour ± 0.05 μl at body temperature, resulting into a quantity of bumetanide injected of 86ng/24h during 3 days. For p75\textsuperscript{NTR} antibody infusion, mini-pumps contained vehicle solution (1M PBS), control rabbit IgG (dilution 1:300 in 1M PBS; sc-2027, Santa Cruz Biotechnology), or p75\textsuperscript{NTR} antibody (dilution 1:300 in 1M PBS of 2mg/ml; final concentration of 6.67µg/ml; AB1554; Millipore). The mortality rate was 5-8% at 24h post-surgery.
qRT-PCR. Expressions of NKCC1 and KCC2 mRNA were detected by real-time quantitative PCR. Total RNAs were isolated from in toto hippocampi using Mini RNeasy kit (Qiagen). First strand cDNA was synthetized from 1mg of total cellular RNA Reverse Transcription QuantiTect kit (Qiagen) at 42°C for 15min then 95°C for 3min in solution containing RNAases inhibitors, oligo-DTs, primers and dNTPs. Real-time PCR cycles were carried out for amplification of NKCC1, KCC2, and GAPDH using termal cycles (Eppendorf). Primers were designed and synthetized by Qiagen. RT-PCR was then carried out using 5mL of cDNA in a final reaction volume of 30mL using SYBER Green Real-time PCR Master Mix. Primers sequences are listed in Table 1. The relative copies are expressed in arbitrary unit.

Western Blots. Animals were killed by decapitation after isoflurane anaesthesia and the hippocampi were quickly dissected out and flash-frozen in liquid nitrogen and stored at -80°C until processed. Brain tissue was homogenized in RIPA buffer (50mM Tris-HCl pH: 8; 150mM NaCl; SDS 0,1%; Deoxycholic Acid 0,5%; 1% triton X-100), containing 1 Complete Protease/Phosphatase Inhibitor Tablet (Thermofisher) and loaded with Laemmli 3X loading buffer. The samples were separated in 7,5-20% SDS–PAGE gel (Criterion gel, Bio-Rad) and transferred to nitrocellulose membrane (Whatman). After blockade in Tris-buffered saline plus 5% BSA, we exposed membranes overnight at 4°C to polyclonal rabbit KCC2-specific antibody diluted 1:2000 in blocking solution. Chemiluminescent detection was performed using horseradish peroxidase-conjugated rabbit-specific antibody diluted 1:3000 in blocking solution for 2h at room temperature and revealed using ECL-plus western blotting reagents (ECL-plus kit, Pierce Biotech). We measured signal intensities with the image analysis software G box (Syngene). Then, membranes were stripped using DTT/SDS in Tris-HCl, pH 7.0, and probed with rabbit anti-β3-tubulin or α-tubulin (Sigma, 1:20000) for normalization and signal detection and revelation following the same procedure. Quantification is performed using Get Plot Analyser plugin (Fiji). The antibodies are listed in Table 1. The relative expression levels are expressed in arbitrary unit.

Immunohistochemistry. Rats were anaesthetized with isoflurane followed by pentobarbital (65mg/kg) and transcardiacaally perfused with cold PBS (1M) prior to cold 4% paraformaldehyde in phosphate buffer saline (PBS). Brains were post-fixed overnight in 4% paraformaldehyde in PBS at 4°C and then coronaly cut with a Leica VT1200S vibratome (60μm thick-sections). Sections (6 slices per condition) were then permeabilized and blocked in PBS with 0.3% Triton X-100 and 5% normal goat serum (NGS) for 1h at room temperature. Incubation with primary antibodies diluted in PBS with 5% NGS and 0.1% Triton X-100 was carried out at 4°C overnight (SPO was incubated additional 24h at 4°C) using different primary antibodies (Table 1). After rinsing 3 times in PBS, slices were incubated with the corresponding Alexa Fluor-conjugated secondary antibodies diluted in PBS.
(1:1000, Invitrogen) for 1h (SPO was incubated additional 2h) at room temperature and finally
counterstained for 5 min with Hoechst 33258 (10μg/ml in PBS, Sigma-Aldrich).
Quantification of KCC2 analysis was performed in blind. For plot analysis of KCC2 the intensity and
distribution associated with the membrane or cytosolic regions in control, pilocarpine rats,
pilocarpine vehicle-treated rats and pilocarpine bumetanide-treated rats were performed using 20X
objective and Plot analysis using Fiji software. The same straight-line length (10µm) was applied from
the nucleus to the external cell compartment of 50 cells from each condition. The plot profile values
were analyzed and calculated the S.E.M. (error bars). The total fluorescence intensity analysis of
KCC2 has been normalized to the number of NeuN-positive cells contained in the ROIs. The 3D object
counter pluggin from Fiji software was used to count the number of cells.

**Acute slices preparation.** Animals were deeply anaesthetized with chloral hydrate (350 mg/kg, i.p.)
and decapitated. The brain was removed rapidly, the hippocampi were dissected, and transverse
400µm slices were cut in oxygenated (95% O2 and 5% CO2) modified artificial cerebrospinal fluid
(mACSF) containing the following (in mM): 132 choline, 2.5 KCl, 1.25 NaH2PO4, 25 NaHCO3, 7 MgCl2,
0.5 CaCl2, and 8 D-glucose; using a Leica VT1000S tissue slicer (Leica VTS1200S, Germany). Slices
were then transferred for rest at room temperature (1-3h) or 34°C (1h) for electrophysiological
recordings in oxygenated (95% O2 and 5% CO2) normal artificial CSF (ACSF) containing the following
(in mM): 126 NaCl, 3.5 KCl, 1.2 NaH2PO4, 26 NaHCO3, 1.3 MgCl2, 2.0 CaCl2, and 10 D-glucose, pH 7.4.

**In vitro electrophysiological recordings.** Hippocampal slices were individually transferred to a
recording chamber maintained at 30–32°C and continuously perfused (2 ml/min) with oxygenated
normal or adapted ACSF containing 5 µM gabazine. Synchronized burst discharges were evoked by
antidromic stimulation of the mossy fiber pathway in the CA3b area while recording local field
potentials in the dentate granular layer using a DAM-80 amplifier (lowfilter, 0.1 Hz; highpass filter, 3
KHz; World Precision Instruments, Sarasota, FL). Signals were analyzed off-line using Clampfit 9.2
(PCclamp) and MiniAnalysis 6.0.1 (Synaptosoft, Decatur, GA). Whole-cell patch-clamp recordings were
done using a MultiClamp 700B amplifier (Molecular Devices, USA). The patch pipette solution
contained the following (in mM): 18 KCl, 111 K-gluconate, 0.5 CaCl2, 2NaOH, 10 glucose, 10 HEPES, 2
Mg-ATP, and 5 BAPTA, (pH adjusted to 7.3 with KOH). Patch pipettes were fabricated from
borosilicate glass (Harvard Apparatus, Edenbridge, UK), and their resistance ranged from 6.5 to 7.5
MΩ. Only cells with access resistances between 10 and 20 MΩ were accepted for analysis.
Membrane potential was clamped at 60 mV. All membrane potential values were corrected for a
liquid-junction potential of 10 mV. The reversal potential of GABAergic currents ($E_{GABA}$) was
determined from the current–voltage (I–V) relation obtained by sequentially clamping the
membrane at different voltage levels. GABA was photolyzed from the α-carboxy-2-nitrobenzyl (DPNI)-caged GABA compound (Tocris) using local uncaging as described previously. Briefly, caged GABA (2.5 mM) was dissolved in the physiological solution and delivered at a flow rate of 1-3 μl/min to the vicinity of the patch-clamped cell using an Aladdin Programmable syringe pump (WPI, Sarasota, FL) and a syringe with an inner tip diameter of 100 μm. For local photolysis of caged GABA, the 405 nm output of a IQ Series laser diode module (Power Technology Inc.) was delivered to the slice via a multimode optical fiber through an Leica LUMPlanFl 40X water-immersion objective. The UV beam yielded an uncaging spot of 10 μm in diameter that was focused either at the soma or at dendrite at a distance of 100 μm from the soma. At each location, current–voltage relationship was determined by varying the holding potential from −90 to −40 mV (10 mV steps). Duration of UV light stimuli was 50 ms.

In vivo electrophysiological recordings. Scalp electroencephalographic (EEG) recordings were performed in freely-moving rats. To implant recording electrodes, rats were anaesthetized with 4.5% isoflurane and maintained at 2.5%, placed in stereotaxic frame, placed on a heating pad to control body temperature and prepared for aseptic surgery. A hole was drilled 2 mm posterior and 1.5 mm right to bregma. A screw connected to EEG transmitter was placed in the skull, and reference electrode was placed rostrally in the cerebellum. The screws were attached to the skull with dental cement and scalp closed. After surgery, rats were kept warm and received saline (0.9% NaCl, s.c.), and analgesic buprenorphine (0.03 mg/kg, s.c.) was administrated to minimize animal pain after surgery. After a 24h recovery, EEG (amplified (1000X), filtered at 0.16–97 Hz pass, acquired at 500 Hz) was monitored using a telemetric system (Data Sciences International, St. Paul, MN) for 5 days, 24 h per day; food and water were given ad libitum. Recordings were analyzed offline using Spike2 8.0. Tonic seizures were defined on the EEG as single or multiple 5-30Hz poly-spike of 5 fold SD amplitude and 10-30 s duration.

Statistical analysis. Data were expressed as means +/- S.E.M. Statistical analyses were performed using GraphpadPrism (GraphPad software 6.01). For comparison between groups with normal distribution, the two-sample unpaired or paired Student’s t test was used for two groups. When data were not normally distributed, the Mann–Whitney rank-sum test (for unpaired data) or the two sample Kolmogorov–Smirnov rank-sum test (for paired data) was used. The one-way ANOVA all-Tukey test was used to compare multiples groups.
Results

NKCC1 and KCC2 expression is altered at day 5 post-SE

The expression of NKCC1 and KCC2 is known to be altered in TLE\textsuperscript{19-23}. However, the temporal changes of these chloride transporters at early stages after SE remain elusive. We found that mRNA transcripts extracted from \textit{in toto} hippocampi display an initial upregulation of NKCC1 and downregulation of KCC2 at day 1 post-SE (d1pSE) that returns to control levels 14 days post-SE (d14pSE) (Figure 1A; NKCC1; ctrl: 1 ± 0.11; d1: 2.10± 0.41; \(p<0.05\); d5: 1.09± 0.02; d14: 0.95± 0.2; KCC2; ctrl: 1 ± 0.28; d1: 0.46 ± 0.02; \(p<0.05\); d5: 0.73 ± 0.10; d14: 1.16± 0.35).

Using western blotting we found that the NKCC1 protein levels were significantly increased by 45% and 70% of control levels at d1pSE and d5pSE respectively with no significant alteration at d14pSE (Figure 1B; ctrl: 1 ± 0.08; d1: 1.45± 0.13; \(p<0.01\); d5: 1.70± 0.24; \(p<0.05\); d14: 1.62± 0.36, when normalized to β3-tubulin and ctrl: 1 ± 0.13; d1: 1.29± 0.12; \(p<0.01\); d5: 2.07± 0.22; \(p<0.05\); d14: 1.23± 0.41, when normalized to β-tubulin) when normalized to either β-tubulin or β3-tubulin. In contrast, the neuron-specific KCC2 protein was significantly reduced to 60% of the control levels at d5pSE with a tendency to a partial recovery at d14pSE (Figure 1C; ctrl: 1 ± 0.09; d1: 1.10± 0.17; d5: 0.58± 0.08; \(p<0.01\); d14: 0.77± 0.16).

Immunohistochemistry towards KCC2 in pilocarpine-treated rats sections showed a preferential cytoplasmic immunoreactivity pattern at d1pSE with a tendency to return to control conditions at d14pSE but with lower fluorescence intensity (Figure 1D,E). This is in contrast to control conditions where KCC2 immunolabeling showed a strong perisomatic pattern. At d5pSE, KCC2 staining was very low and diffuse within the neuron cell bodies. These findings, showing an overall change in KCC2 neuronal distribution, were consistent within the DG, CA3, and CA1 regions (Figure 1D,E).

Taken together, our qRT-PCR, immunoblot and immunofluorescence results demonstrate that the most drastic dysregulation of both NKCC1 and KCC2 expressions occurred at d5pSE.

Bumetanide functionally reduces early established ectopic MF sprouting

A major reorganization of hippocampal network observed in both human patients and animal models of TLE is the ectopic MF sprouting\textsuperscript{7,30-32}, a phenomenon partly responsible for the generation of epileptiform activities in human patients and animal models\textsuperscript{11,31} of TLE. However, although extensively studied during the chronic phase\textsuperscript{19,33}, it has been poorly assessed during the early stages after SE\textsuperscript{29,30}. Thus, we asked whether ectopic MF sprouting occurs in parallel to the robust changes in
KCC2 and NKCC1 expression observed at d5pSE (Figure 1). Immunohistochemical visualization of rMF using synaptoporin (SPO) staining \textsuperscript{13,34} revealed a marked ectopic band in the molecular layer of DG and strong staining in the granular cell layer in pilocarpine-treated rats at d5pSE, while we found no ectopic band stained in control rats (Figure 2A; arrowheads ctrl: 1 ± 0.05; d5pSE: 5.05 ± 0.04, \(p<0.001\)). Thus, these results demonstrate that the ectopic MF sprouting is established at d5pSE, correlating with the pathological KCC2 and NKCC1 expression pattern.

We next asked whether abnormal chloride homeostasis might be involved in the early establishment of the ectopic MF sprouting. In order to achieve this, we use bumetanide to functionally block NKCC1. Because its blood-brain-barrier penetration has been challenged \textsuperscript{35,36}, we infused bumetanide at a dose of 20\(\mu\)M, in accordance with previous pharmacokinetic studies \textsuperscript{37,38} intracerebrally and continuously from d2 to d5pSE, using osmotic minipumps (Figure 2B).

Interestingly, at d5pSE, we observed a 45% decrease in SPO fluorescence intensity in bumetanide-treated rats as compared to the vehicle-treated rats (Figure 2C; vehicle-treated rats: 5.03 ± 0.04 vs bumetanide-treated: 2.83 ± 0.049; \(p<0.001\), normalized to control). In order to find out whether this rMF sprouting is functional as early as d5pSE, we performed the antidromic stimulation of the mossy fibers pathway in the CA3b area while recording local field potentials in the DG cell layer (Figure 2D).

Importantly, we found that the evoked population burst was strongly reduced in bumetanide-treated rats as compared to vehicle-treated pilocarpine rats (Figure 2D; vehicle-treated vs bumetanide-treated: 14326.31± 3853 vs 5252.66 ± 1608; \(p<0.05\)).

Together these results demonstrate that a functional recurrent MF network is already in place at d5pSE. Strikingly, the use of bumetanide \textit{in vivo} functionally reduced this ectopic MF sprouting.

Bumetanide blocked NKCC1 upregulation and rescued KCC2 function at d5pSE.

Several \textit{in vitro} and \textit{in vivo} studies have shown that bumetanide can affect the functional expression of NKCC1\textsuperscript{21,22}. To better understand the mechanism by which bumetanide reduced the ectopic MF sprouting and its aberrant network activity, we examined the effect of bumetanide on NKCC1 and KCC2 expression and function. Here, we show that intracerebral infusion of bumetanide partially blocked NKCC1 upregulation. Surprisingly, total KCC2 expression levels were fully restored in bumetanide-treated pilocarpine rats as compared to vehicle-treated rats (Figure 3B; NKCC1: vehicle-treated rats: 2.28 ± 0.10 vs 1.68 ± 0.14; \(p<0.05\), normalized to control. KCC2: vehicle-treated vs bumetanide-treated rats: 0.47 ± 0.15 vs 1.40 ± 0.23; \(p<0.05\), normalized to control). Using histological analysis, we show that KCC2 displays a preferentially perisomatic staining in bumetanide-treated rat similar to vehicle-treated rats at d5pSE in the DG, CA3 and CA1 regions (Figure 3D).
We next investigated whether bumetanide also restored KCC2 function. To investigate KCC2 ability to extrude Cl⁻ we used the whole-cell patch-clamp approach coupled to GABA uncaging in hippocampal slices in the CA1 region (Figure 3E). The approach consists of inducing a somatic load of chloride through the patch pipette followed by uncaging of GABA at the soma and at a distance of 100µm in the apical dendrite. In vehicle-treated rats, we observed that the $E_{\text{GABA}}$ gradient was significantly smaller (Figure 3E, -0.12 ± 0.005mV/100µm) than in bumetanide-treated rats at d5pSE (Figure 3E, -0.17 ± 0.007mV/100µm). The later was almost identical to the levels previously measured for this age.²⁸

Although there is ample evidence for the specificity of bumetanide as inhibitor of NKCC1 uptake activity, when used at low concentration, the effect observed could be derived from nonspecific action of this compound due to tissue accumulation. Interestingly, high concentration of bumetanide (100µM) exacerbated the KCC2 downregulation observed at d5pSE (Figure 4; DG: vehicle-treated rats: 0.44 ± 0.09 vs bumetanide 20-treated: 0.71 ± 0.1, p<0.01 vs bumetanide 100-treated: 0.44 ± 0.08, p<0.001; CA3: vehicle-treated rats: 0.31 ± 0.12 vs bumetanide 20-treated: 0.53 ± 0.12, p<0.001 vs bumetanide 100-treated: 0.01 ± 0.11, p<0.001 and CA1: vehicle-treated rats: 0.60 ± 0.10 vs bumetanide 20-treated: 0.81 ± 0.13, p<0.01 vs bumetanide 100-treated: 0.35 ± 0.11, p<0.001). This result shows that nonspecific effects due to either accumulation of bumetanide and derived inhibition of KCC2, would result in opposite effects. Our findings show that intracerebral bumetanide infusion from d2 to d5pSE rats at low concentration is specific, able to partially block NKCC1 expression, and restored not only KCC2 protein expression but also its function at d5pSE within the hippocampus.

**p75NTR upregulation drives the ectopic MF sprouting and is blocked by bumetanide**

We have previously shown that, in axotomized organotypic hippocampal slices, altered NKCC1/KCC2 balance induced p75NTR upregulation that could be prevented by bumetanide.²⁶ P75NTR is also known to play a key role in neurite outgrowth and is crucial for axon formation in the DG under physiological conditions.⁴⁰ Therefore, we asked whether bumetanide could regulate p75NTR expression to mediate the effects observed on the ectopic MF sprouting at d5pSE (Figure 5A). Using western blot from in toto hippocampi, we found that p75NTR expression level was reduced in bumetanide-treated rats as compared to vehicle-treated rats at d5pSE (Figure 5B, vehicle-treated vs bumetanide-treated rats: 1.42 ± 0.13 vs 0.90 ± 0.13, p<0.05, normalized to control 1 ± 0.09). Then, using immunohistochemistry, we showed that as previously described, p75NTR was expressed at low level in DGCs in control conditions. Interestingly, bumetanide application strongly reduced the
SE-induced abnormal p75NTR expression within the granule cells at d5pSE (Figure 5C, vehicle-treated vs bumetanide-treated rats: 1.64 ± 0.04 vs 1.14 ± 0.02; p<0.001, normalized to control).

Finally, in order to establish if p75NTR upregulation is involved in the formation of ectopic MF sprouting, we decided to infuse intracerebrally a previously validated24 p75NTR functionally blocking antibody (p75NTR-Ab) from d2 to d5pSE (Figure 5D for the experimental design). Interestingly, SPO staining revealed significantly decreased fluorescence intensity in p75NTR Ab-treated as compared to vehicle- or IgG-treated rats at d5pSE (Figure 5D, IgG-treated rats: 6.27 ± 0.57; vehicle-treated vs p75NTR Ab-treated rats: 5.06 ± 0.5 vs 3.17 ± 0.24; p<0.001, normalized to control).

Altogether these results suggest that bumetanide-mediated functional reduction of ectopic MF sprouting is, at least partly, mediated by p75NTR upregulation blockade in vivo at d5pSE in the pilocarpine model of TLE.

**Early bumetanide infusion reduced seizure frequency during the chronic phase of TLE**

The transient application of bumetanide had profound positive effects on morphological and functional reorganization as well as on functional expression of chloride homeostasis regulatory proteins within the hippocampal formation during the early post-SE window.

To examine whether transitory application of bumetanide early after SE had a long-lasting effects, we used a similar approach to the one use above with the difference that sprouting is analyzed during the chronic and not the early phase of TLE (Figure 6A). The examination of the cortical EEG (24h/d) from d2 to d5 post-SE did not reveal interictal and ictal activities under the present experimental conditions in vehicle nor bumetanide infused animals (data not shown). However, following bumetanide infusion from d2 to d5 post-SE, SPO staining revealed a reduction of the ectopic band in the molecular layer of DG in bumetanide-treated rats as compared to vehicle-treated rats at 2 months post-SE (Figure 6B; vehicle-treated vs bumetanide treated: 7.34 ± 0.35 vs 6.11 ± 0.41; p<0.05, normalized to control). This interesting long-lasting effect of bumetanide on ectopic MF sprouting prompted us to analyze the seizure activity with telemetric scalp EEG recordings at 2 months post-SE after the rats underwent the same experimental paradigm as in the previous experiments (Figure 6A). In this chronic condition, we observed that the number of ictal events was reduced by 50% in the bumetanide-treated group as compared to vehicle-treated group (Figure 6C-E: vehicle-treated vs bumetanide treated: 13.94 ± 0.74 vs 7.05 ± 0.39; p<0.05, normalized to control), without a significant change in their duration (Figure 6D; vehicle-treated vs bumetanide treated: 18.59 ± 0.71 vs 17.65 ± 1.04; p>0.05).
Discussion

Studies on TLE show that a major glutamatergic circuit reorganization within the DG, the ectopic rMF, is responsible for an increase in hippocampal excitability and recurrent seizures\(^1,143\). Equally strong evidence has been found showing that adult seizures are generated in a circuit in which GABA\(_A\)-R-mediated inhibition is impaired\(^21,44\). Within this context, compromised GABAergic inhibition resulting from chloride homeostasis dysregulation in epileptic-like conditions has been extensively studied both in animal models and human resections in TLE\(^19-23\). In these studies, acute use of bumetanide, leading to restoration of chloride homeostasis, has been shown to have potent anticonvulsant effects in the adult brain. However, whether GABAergic depolarization plays a role in the precipitating events of epileptogenesis and could exert long-term effect on morphological and functional features of glutamatergic network rewiring and subsequently affect seizure activity has not been assessed. Here, we have addressed these issues by using a pilocarpine model of TLE. First, we found that early ectopic rMF sprouting and subsequent rMF-driven network aberrant activity was established as early as 5 days after the SE. This time point coincided with NKCC1 upregulation and KCC2 downregulation. Importantly, we showed that functional inhibition of NKCC1 with the loop diuretic bumetanide from d2 to d5 post-SE resulted in restoration of chloride homeostasis and robust attenuation of rMF-driven network activity during early epileptogenesis. These effects extended to the chronic epileptic phase and resulted in strongly reduced seizure frequency. The mechanism appears to involve upregulation of the pan neurotrophin receptor p75\(^{NTR}\) as inhibition of NKCC1 activity significantly reduced the SE-induced upregulation of this receptor. Previous studies have suggested the involvement of NGF/TrkA/p75\(^{NTR}\) signaling on MF sprouting in a kindling animal model\(^17\). We now also demonstrate that p75\(^{NTR}\) is a key factor in the early establishment of the ectopic rMF sprouting in TLE as: i) in accordance to previous studies\(^45,46\) p75\(^{NTR}\) is upregulated at d5pSE, ii) while bumetanide strongly reduced the ectopic rMF sprouting it also blocked p75\(^{NTR}\) upregulation, iii) a functionally blocking p75\(^{NTR}\) antibody reduced the ectopic rMF sprouting. Furthermore, previous reports suggest a role of GABA\(_A\)-R-mediated depolarization on axonal elongation during development and adult neurogenesis\(^47\). Our results show that early post-SE bumetanide application reduces ectopic rMF, p75\(^{NTR}\) expression, and chronic seizures. These effects are associated with impairment of GABA\(_A\)-R-mediated depolarization by bumetanide through its specific inhibitory action on the chloride co-transporter NKCC1. Future investigations are required to elucidate the causal relationship between altered GABAergic transmission and aberrant neurotrophin signaling under pathological conditions. The reported results also suggest that the ectopic MF sprouting, promoted by chloride homeostasis dysregulation and p75\(^{NTR}\) signaling pathway during early epileptogenesis, plays a key role for the
development of chronic seizures since post-SE application of bumetanide reduced the number of spontaneous and recurrent seizures in chronic epileptic rats.

It is plausible that the effects exerted by bumetanide are mediated through an effect on post-SE remnant seizure activity. Indeed, bumetanide has been shown to have anticonvulsant action in numerous seizure models. Within the early period following SE (2-3 days post-SE), sporadic remnant interictal and ictal activities can occur. However, in the present experimental paradigm, the pilocarpine-treated rats exhibit a seizure-free period from day 2 to day 5 post-SE (data not shown). Nonetheless, we cannot exclude an effect of bumetanide on sporadic hippocampal interictal events that could not be reliably detected from cortical EEG recordings in contrast to seizure activity. Further studies are necessary to dissect if and how GABA depolarization is involved in post SE network activity during early epileptogenesis.

Altogether, the present study suggests that SE-induced depolarizing GABAergic signaling could play a mechanistic role in the early and long lasting rewiring of glutamatergic networks that is implicated in the severity of chronic seizures. These results also prompt to investigate optimal conditions for better and more efficient restriction of seizures in the chronic phase using this approach.

In the clinical setting of symptomatic and unprovoked epilepsy the use of anti-epileptic drug prophylaxis during the acute period following SE is recommended. In addition, symptomatic epilepsy specially involving the temporal lobe often displays pharmaco-resistance to available antiepileptic drugs. Thus, the results presented here prompt us to conclude that normalizing chloride homeostasis with the FDA-approved drug bumetanide during the acute posttraumatic period could be able to sustainably reduce the pathological glutamatergic network rewiring and subsequent chronic adult epilepsy. Moreover, prophylactic approaches to correct dysregulation of GABAergic transmission and p75NTR signalling are novel clinically relevant targets that could complement the current therapy following acute symptomatic seizures.

Acknowledgments

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Author contributions


Potential conflict of interest

A patent (EPI15306384) has been filled by INSERM Transfert “Methods and pharmaceutical composition for the treatment of post-traumatic epilepsies”.

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Figure legends

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