NKCC1 Does Not Accumulate Chloride in Developing Retinal Neurons

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NKCC1 does not accumulate chloride in developing retinal neurons. J Neurophysiol 98: 266–277, 2007. First published May 9, 2007; doi:10.1152/jn.00288.2007. GABA excites immature neurons due to their relatively high intracellular chloride concentration. This initial high concentration is commonly attributed to the ubiquitous chloride cotransporter NKCC1, which uses a sodium gradient to accumulate chloride. Here we tested this hypothesis in immature retinal amacrine and ganglion cells. Western blotting detected NKCC1 at birth and its expression first increased, then decreased to the adult level. Immunocytochemistry confirmed this early expression of NKCC1 and localized it to all nuclear layers. In the ganglion cell layer, staining peaked at P4 and then decreased with age, becoming undetectable in adult. In comparison, KCC2, the chloride extruder, steadily increased with age localizing primarily to the synaptic layers. For functional tests, we used calcium imaging with fura-2 and chloride imaging with 6-methoxy-N-ethylquinolinium iodide. If NKCC1 accumulates chloride in ganglion and amacrine cells, deleting or blocking it should abolish the GABA-evoked calcium rise. However, at P0-5 GABA consistently evoked a calcium rise that was not abolished in the NKCC1-null retinas, nor by applying high concentrations of bumetanide (NKCC blocker) for long periods. Furthermore, intracellular chloride concentration in amacrine and ganglion cells of the NKCC1-null retinas was \( > 30 \) mM, same as in wild type at this age. This concentration was not lowered by applying bumetanide or by decreasing extracellular sodium concentration. Costaining for NKCC1 and cellular markers suggested that at P3, NKCC1 is restricted to Müller cells. We conclude that NKCC1 does not serve to accumulate chloride in immature retinal neurons, but it may enable Müller cells to buffer extracellular chloride.

INTRODUCTION

During brain and retinal development, GABA’s action shifts from excitatory to inhibitory (Ben Ari 2001, 2002; Catsicas and Mobbs 2001; Fischer et al. 1998; Sernagor et al. 2003; Stellwagen et al. 1999; Zhang et al. 2006b; Zhou 2001). This shift results from a decline of intracellular chloride concentration from \( \sim 30 \) to \( \sim 15 \) mM which reduces the chloride equilibrium potential \( (E_{CI}) \) from above to below the resting potential \( (E_{rest}) \) (Zhang et al. 2006b). Maintaining an equilibrium potential different from the resting potential requires an active transport of chloride: extrusion for \( E_{CI} < E_{rest} \) and accumulation for \( E_{CI} > E_{rest} \).

The extruder is probably the neuron-specific \( K^+ - Cl^- \) cotransporter, KCC2. This transporter normally extrudes chloride with the outward potassium gradient (Payne et al. 1996), and evidence supporting its function as the main chloride extruder is strong. It upregulates during development (Li et al. 2002; Vu et al. 2000; Zhang et al. 2006a), and in adult, it broadly localizes to GABA-receptive neurons (Li et al. 2002; Lu et al. 1999; Vardi et al. 2000; Williams et al. 1999). Additionally, pharmacological blockade, gene deletion, or inhibition of expression with siRNA, all result in a higher than normal \( E_{CI} \) (Balakrishnan et al. 2003; DeFazio et al. 2000; Gutacsi et al. 2003; Hubner et al. 2001; Jarolimek et al. 1999; Rivera et al. 1999; Zhu et al. 2005).

The accumulator is thought to be the ubiquitous \( Na^+ - K^+ - Cl^- \) cotransporter NKCC1, which uses the inward sodium gradient to accumulate chloride along with sodium and potassium (reviewed by Russell 2000). The evidence supporting NKCC1 as the chloride accumulator in certain mature neurons is good. NKCC1 localizes to neurons known to maintain high intracellular chloride such as olfactory epithelial cells and retinal horizontal cells (Reisert et al. 2005; Vardi et al. 2000); the olfactory epithelial cells in the NKCC1-null mouse show a reduced calcium-activated inward chloride current due to reduced intracellular chloride (Reisert et al. 2005), and in dorsal root ganglion cells, blocking NKCC1 with bumetanide, manipulating sodium concentrations, or deleting the gene encoding NKCC1 reduces intracellular chloride concentrations (Rohrbough and Spitzer 1996; Sung et al. 2000).

In contrast, evidence supporting NKCC1 as the chloride accumulator in immature neurons is weak and relies mostly on the downregulation of NKCC1 mRNA and protein during development (Ikeda et al. 2003; Kanaka et al. 2001; Li et al. 2002; Mikawa et al. 2002; Plotkin et al. 1997b). Functional experiments were only performed in neocortex where the downward shift of high \( E_{CI} \) with bumetanide was correlated with NKCC1 mRNA expression (Yamada et al. 2004). No study has utilized the NKCC1 knockout mouse to test the transporter’s function in early development except for its effect on epileptic seizure (Dzhala et al. 2005). Here we show that in immature retinal amacrine and ganglion cells, NKCC1 blockade or deletion does not affect intracellular chloride. Furthermore, in the first postnatal week, NKCC1 appears to be restricted to Müller cells, and is probably not localized to any retinal neurons.

METHODS

Western blotting

Newborn mice were deeply anesthetized on postnatal days P0 (date of birth) to P6 with halothane and killed by decapitation. Older mice were deeply anesthetized with an intraperitoneal injection of ketamine (85 \( \mu \)g/gm) and xylazine (13 \( \mu \)g/gm).
and 2 EDTA, 2.5 mM Tris, 5 Tris-HCl, 600 g/gm) followed by anesthetic overdose. Animals were treated in compliance with federal regulations and University of Pennsylvania policy. Retinas were collected into two developmental series: series 1: P0, 3, 5, 8, 12, 14, 24, and adult (>3 mo) and series 2: P0, 3, 5, 9, 12, 16, and adult. For ages younger than P5, retinas from two to three littermates were combined. Retinas were frozen immediately with liquid N2 and stored at −80°C. Retinas from each series were processed in parallel: they were homogenized in 0.2–1 ml ice-cold homogenization buffer [5 ml/g wet weight; containing (in mM): 320 sucrose, 5 Tris, 2 Tris-HCl, and 2 EDTA, 2.5 β-mercaptoethanol, pH adjusted to 7.4 at room temperature] containing protease inhibitor cocktails (P8340; Sigma Aldrich, St. Louis, MO), and the homogenates were centrifuged at 6,000 g for 10 min at 4°C to remove debris.

Equal amounts of protein (10 μg) from each age of a developmental series were mixed with NuPAGE LDS sample buffer (Invitrogen, Carlsbad, CA), incubated for 45 min at 37°C, then loaded onto 5.0 or 7.5% Tris-glycine gel (Bio-Rad). Electrophoresis was performed with a mini-protein II electrophoresis cell (Bio-Rad) under reduced denaturing conditions. The resolved proteins were transferred to a polyvinylidene difluoride (PVDF) membrane (Bio-Rad) in transfer buffer (192 mM glycine, 25 mM Tris-Cl, pH 8.3, and 20% methanol) for 40–50 min at 15 V using a Bio-Rad Trans-Blot semi-dry transfer cell. The membrane was blocked with 5–7% skim milk in TBS-T (10 mM Tris-HCl, 150 mM NaCl, 0.1% Tween 20, PH 7.4) for 1 h, and incubated with the primary antibody overnight at 4°C in TBS-T-milk. After extensive washes, the membrane was incubated with the secondary antibody (anti-rabbit Fab fragments linked to HRP, Protos, Burlingame, CA) in TBS-T-milk for 2 h at room temperature. After extensive washes in TBS-T, bound antibody was detected by enhanced chemiluminescence (Pierce, Rockford, IL).

In certain experiments, comparison of NKCC1 and KCC2 expression was done on the same blot. In these experiments, we removed the antibodies from the membrane with a stripping buffer (Pierce, Rockford, IL) and re-probed the membrane with a second antibody. The transporter patterns detected after the first probe was stripped and a second probe applied were similar to those with the fresh probe, but the intensity was reduced. Thus only the freshly probed blots were included in our quantification.

For quantification, chemiluminescence was captured within 20 min of the reaction using a Kodak Imaging Station 440CF system (with a 12-bit cooled CCD camera) and the result was quantified using Kodak 1D imaging analysis software (Sunnyvale, CA). Control experiments have shown that detection of protein is linearly proportional to the amount of protein in this assay. Exposures of each blot were adjusted to ensure that all signals were within the linear range. After subtracting background intensity from each pixel, we summed all the signals within a band to give the net band intensity. For Display, the 12-bit chemiluminescence imaging data were converted to 8-bit after a linear transform with saturation at the high intensities to allow visibility of the low-intensity bands.

Immunostaining

Eyecups from several littermates were grouped into the following postnatal ages, series 1: 0, 1, 3, 5, 7, 13, and 20 and series 2: 0, 2, 5, 7, 10, 12, 16, and 18. For intensity comparison, eyecups were immersion-fixed for 1 h at room temperature in 4% paraformaldehyde and 0.01% glutaraldehyde diluted in 0.1 M phosphate buffer at pH 7.4. For the rest of the immunostaining experiments, fixation varied between 2 and 4% paraformaldehyde fixed for 10–60 min. Retinas were cryoprotected with 30% sucrose in phosphate buffer (overnight at 4°C). Eyes were then frozen in a mixture of Tissue Freezing Medium (Electron Microscopy Sciences, Ft. Washington, PA) and 20% sucrose (1:2), and cryosectioned vertically in 10-μm sections. Cryosections were stained according to a standard protocol: soak in diluent containing 0.1 M phosphate buffer, 10% normal goat serum, 5% sucrose, and 0.3% Triton X-100; incubate in primary antibody overnight at 4°C; wash and incubate in secondary antibody for 3 h at room temperature. Sections were mounted in Vectashield (Vector Laboratories, Burlingame, CA) and visualized with a confocal microscope (Leica, Nussloch, Germany). In double-labeling experiments, sections were incubated with a mixture of primary antibodies followed by a mixture of secondary antibodies.

To examine NKCC1 localization at the electron microscope level, fixed tissue was frozen and thawed three times prior to Vibratome sectioning (100 μm thick). Sections were incubated in diluent and primary antibody as in the preceding text (stain was invisible with <0.3% Triton X-100), then in secondary antibody conjugated to HRP. They were then rinsed, developed in 0.05% 3, 3’-diaminobenzidine (DAB) +0.01% hydrogen peroxide in phosphate buffer for 15 min, washed, and postfixed. DAB reaction product was intensified with gold-substituted silver and sections were osmicated, dehydrated, and mounted in Epon 812. Ultrathin sections were poststained with uranyl acetate. Primary antibodies are listed in Table 1. Secondary antibodies: for single labeling, we used anti-rabbit and anti-mouse conjugated to FITC, and for double labeling, we also used anti-rabbit and anti-mouse conjugated to rhodamine or to Cy3. All secondary antibodies were from Jackson Immunoresearch (West Grove, PA).

Calcium imaging

A retina was cleaned of vitreous, detached from the pigment epithelium, and cut into two to three pieces. Retinal pieces were mounted on a filter paper and maintained in bicarbonate-based Ames medium saturated with carbogen (95% O2-5% CO2). In young mice (P0–P6), ganglion cells were often exposed when detaching the vitreous; in older mice, ganglion cells were exposed by gently scratching the vitreal surface with a scalpel tip or a brush. This procedure ensured that fluorescence was collected cleanly from ganglion and amacrine cells somas (Zhou 2001). Retinal pieces were loaded in 5–10 μM Fura-2 AM (Molecular Probes; Eugene, OR) in oxygenated Ames medium for 1 h at 27–30°C. A retina was placed in an optical recording chamber mounted on a fixed-stage, upright microscope (Olympus BX51WI) and continuously superfused (3–4 ml/min) with Ames medium preheated to 30–33°C. Fura-2 was excited alternately at 340 and 380 nm with a xenon arc lamp in lambda DG4 (Sutter Instrument; Novato, CA) and filtered with a 510/40 nm emission filter (set 71000, Chroma; Rockingham, VT), and captured with a Hamamatsu Orca II ER digital CCD camera (C4742-
98-24ER, Hamamatsu; Hamamatsu city, Japan). Wavelength switch and imaging capture were controlled by Openlab software (Improvision; Lexington, MA) running under Mac G4. To minimize potential UV phototoxicity, the illuminated retinal area was restricted by closing the field aperture to match the CCD imaging field (216 x 300 μm), and the pixels were binned (2 x 2). Exposure time was 25–200 ms, and sampling interval was 3–4 s.

**Determining intracellular chloride with a reference method**

To determine intracellular chloride level, we used the reference methods developed earlier (Zhang et al. 2006b). Briefly, fresh 6-methoxy-N-ethyl-1,2-dihydroquinoline (dihydro-MEQ) was synthesized from 5 mg 6-methoxy-N-ethylquinoline iodide (MEQ) according to the protocol provided by Molecular Probe (Eugene, OR). The reduced product was resuspended in dimethyl sulfoxide, and added to Ames medium to yield a loading concentration of 10–15 μM. Intracellular loading was performed at 35°C (Zhang et al. 2006b). The optical setup and acquisition system were the same as that for calcium imaging except that excitation and emission were filtered for MEQ and calcein. Excitation alternated between 345 nm (D345/10x, Chroma) and 485 nm (S485/25x, Chroma), and emission was collected with a dual band filter with peaks at 450 and 535 nm (set 91018, Chroma). Images were collected every 10–15 s. After the unperturbed MEQ fluorescence of cells located at the ganglion cell layer was recorded, the retina was perfused with a reference solution containing 30 mM Cl\(^{-}\) (instead of Na\(^{+}\)), 14 mM K\(^{+}\) (instead of Na\(^{+}\)), 14 mM Cl\(^{-}\)/OH\(^{-}\) exchanger tributyltin (Krapf et al. 1988). We chose a 30 mM reference solution because the reference method is most accurate when the reference solution is close to the predicted intracellular solution, and the intracellular solution in young wild-type ganglion cells is close to this value (Zhang et al. 2006b). Cells were then imaged until intracellular chloride concentration equilibrated and the fluorescent remained relatively stable (~15 min).

**Optical signal analysis**

Imaging and statistical analysis were done with Openlab and Excel (Microsoft; Seattle, WA). The time-lapse image stacks were first registered to correct for retinal movements. Then a region of interest was drawn around a dye-loaded ganglion or amacrine cell in the ganglion cell layer, and the cell's fluorescence was measured at each time point. Background due to dark current, ambient light, and autofluorescence was estimated in control experiments with unloaded retinas and subtracted from the cell’s fluorescence measurement. For calcium imaging with the ratiometric indicator fura-2, we computed the ratio of the cell’s average fluorescence at 340 to that at 380 nm.
with the commonly used T4 monoclonal antibody raised against human NKCC1 (supplemental Fig. 1A). In Western blotting, this antibody recognized a band at the correct MW (155 kDa), which was eliminated in the NKCC1-null mouse, but it also recognized two prominent bands at 110 and 60 kDa that were not eliminated in the null mouse supplemental Fig 1B. In immunostaining, horizontal cells, which were detected by the T4 antibody when applied to rabbit and monkey (Vardi et al. 2000), were not detected in the mouse. Staining with this antibody was present in dendritic tips of on bipolar cells (strongly), and in lateral processes in the inner plexiform layer (weakly). However, this staining pattern appeared similar in the NKCC1-null retina (supplemental Fig. 1, C and D). Further testing revealed that staining of HEK cells transfected with mouse NKCC1 resembled that of untransfected cells (supplemental Fig. 1E). These experiments suggested that the T4 antibody does not recognize mouse NKCC1 under our staining conditions. Consequently, data collected with the T4 antibody were not used.

In contrast, another antibody against NKCC1 (a polyclonal antibody raised against the C-terminus of the mouse protein; sequence is shown in supplemental Fig. 1A) did not stain untransfected HEK cells and gave a membrane-associated staining in cells transfected with NKCC1 (supplemental Fig. 1, F and G). Western blotting with this antibody gave a doublet at the expected molecular weight (~150 kDa; Fig. 1A). Neither band was present in the NKCC1-null retina. Immunostaining of the adult wild-type retina was primarily restricted to horizontal cells, where somas and processes stained strongly (Fig. 1B); processes of on bipolar cells were unstained (not shown). Immunostaining of the young wild-type retina showed staining throughout the retina (Fig. 1C). The intensities of both staining patterns, in the adult and young retinas, were diminished in the NKCC1-null retinas (n > 3; Fig. 1, B and C). Thus the polyclonal anti-NKCC1 is specific and can be used for further localization and quantification. For KCC2, we used an antibody

Genotyping

The NKCC1 knockout mouse was generated in a mixed C57BL/6J-DBA/2J background and homozygous animals were obtained from heterozygous breeding (Pace et al. 2000). Genotyping was done by PCR with two sets of primers: one detecting the wild-type NKCC1 allele, the other detecting the mutant NKCC1 allele (neomycin resistant gene cassette). NKCC1 upper primer 5'-CGTGTGTTGTGGATGCCA and NKCC1 lower primer 5'- GTTGTTCTATTAGCCCGTAAAGACA generate 106-bp PCR product. Neo upper primer 5'-ATGGATTTGACGCAAGTTCT and Neo lower primer 5'-CCTTGAGCCTGGCCAACGAT generate 512-bp PCR product.

RESULTS

Specificity of antibodies

Initial experiments to localize NKCC1 were performed with the commonly used T4 monoclonal antibody raised against human NKCC1 (supplemental Fig. 1A). In Western blotting, this antibody recognized a band at the correct MW (155 kDa), which was eliminated in the NKCC1-null mouse, but it also recognized two prominent bands at 110 and 60 kDa that were not eliminated in the null mouse supplemental Fig 1B. In immunostaining, horizontal cells, which were detected by the T4 antibody when applied to rabbit and monkey (Vardi et al. 2000), were not detected in the mouse. Staining with this antibody was present in dendritic tips of on bipolar cells (strongly), and in lateral processes in the inner plexiform layer (weakly). However, this staining pattern appeared similar in the NKCC1-null retina (supplemental Fig. 1, C and D). Further testing revealed that staining of HEK cells transfected with mouse NKCC1 resembled that of untransfected cells (supplemental Fig. 1E). These experiments suggested that the T4 antibody does not recognize mouse NKCC1 under our staining conditions. Consequently, data collected with the T4 antibody were not used.

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Fig. 1. Polyclonal anti-NKCC1 is specific in retina. A: Western blots of wild-type retina immunoreacted with the polyclonal antibody against NKCC1 shows a doublet (~150 kDa, which is absent from the NKCC1-null (-/-) retina. The doublet likely represents NKCC1 protein with different glycosylation levels (Plotkin et al. 1997a,b). B: confocal fluorescence micrograph of vertical adult sections immunostained for NKCC1. Horizontal cells (hc) are strongly stained in the wild-type retina but not in NKCC1-null retina. In B and C, staining parameters and image acquisition were identical for wild-type and null retinas. PRL, photoreceptor layer; ONL, outer nuclear layer; OPL, outer plexiform layer; INL, inner nuclear layer; IPL, inner plexiform layer; GCL, ganglion cell layer; NFL, nerve fiber layer. C: confocal fluorescence micrograph of vertical juvenile sections (P4-5) immunostained for NKCC1. Staining is present throughout the wild-type retina and is absent from the null retina. No 1°. Primary antibody was omitted.
that has been established to be specific in rodent retina (Vardi et al. 2000; Vu et al. 2000).

**NKCC1 and KCC2 upregulate during development**

In mouse, intracellular chloride concentration decreases at about P6 (Zhang et al. 2006b), so we used Western blots and immunohistochemical staining to map the pattern of chloride cotransporters across P0-24 aged retinas. Western blots showed that NKCC1 was present as early as P0, maintained a constant level through P5, then rapidly increased reaching a plateau between P12 and 24, and finally declined to a much lower adult level (Fig. 2). In contrast, KCC2 was undetectable during P0-5, clearly visible at P8, then rapidly increased reaching a peak between P17-24, and finally dropped slightly to the adult level.

**NKCC1 expression in the ganglion cell layer is higher in developing retina than in adult**

By immunohistochemical staining, NKCC1 was already detected at birth throughout the retina with most somas outlined (Fig. 3). In the inner retina, the ganglion cell layer and the nerve fiber layer just below it were prominently stained by P3; the inner plexiform layer only stained weakly. By P20, staining of the ganglion cell layer and the inner plexiform layer dropped to background levels. In the outer retina, staining for NKCC1 was first detected at P5, the time that this layer becomes distinct. By P7, staining localized to somas and processes at the top of the newly formed inner nuclear layer. On the basis of location and morphological criteria, these appear to belong to horizontal cells. In the following week, staining intensity increased by roughly 2.5-fold and then declined slightly to the adult level. Thus in the layers where ganglion and amacrine cell somas reside, NKCC1’s expression is higher in the early retina than in the adult, consistent with it serving as a chloride accumulator in immature neurons.

**KCC2 expression localizes primarily to synaptic layers and increases during development**

Contrary to Western blotting results, immunocytochemistry showed that KCC2 was already detectable at birth (Fig. 4). Apparently immunostaining is more sensitive, possibly because detection in Western blotting is limited by the amount of protein that can be loaded into a lane. The staining at birth was very weak and confined to the developing inner plexiform layer. This staining was punctate, it distributed fairly homogeneously throughout the inner plexiform layer, and steadily increased with age. At P7, it became possible to detect the somas of amacrine and ganglion cells. By P10, staining in the inner plexiform separated into two wide bands, roughly corre-
sponding to the ON and OFF sublayers with a thin band between strata 2 and 3 devoid of staining. Staining for KCC2 in the outer retina was detected as soon as the outer plexiform layer formed (P5-7) and this staining continued to increase with age. This data confirms that KCC2 upregulates during development approximately following the order of differentiation of different cell types, consistent with findings in rat, ferret, and turtle (Leitch et al. 2005; Vu et al. 2000; Zhang et al. 2006a).

NKCC null retina has normal gross morphology and physiology

To test the effect of NKCC1 on GABA’s function and chloride accumulation in early retina, we used an NKCC1-null mouse. This mouse is deaf, infertile, impaired in pain perception and in locomotion, and may have compromised olfactory sensation (Delpire et al. 1999; Flagella et al. 1999; Pace et al. 2000; Reisert et al. 2005; Sung et al. 2000). To evaluate the gross morphology and physiology of the NKCC1-null retina, we performed immunostaining. The lack of NKCC1 expression has been established in the preceding text (Fig. 1). Retinas of the NKCC1-null mouse had a thickness similar to those of wild type and contained all the layers and cell types (Fig. 5). Thus photoreceptors and ON bipolar cells stained for recoverin, rod bipolar cells stained for Gα13 and Gαo, all showed normal densities and stratifications. Similarly, amacrine cells stained for calbindin, GABA, and ChAT, and ganglion cells stained for parvalbumin and neurofilament had normal morphology and stratifications. Müller cells stained with an antibody against glutamine synthetase showed radial processes that, as in the adult, stained strongly in the outer nuclear layer and in their terminals in the nerve fiber.

![FIG. 4. KCC2 immunoreactivity increases during postnatal maturation. Confocal radial sections from one developmental series immunostained for KCC2. All sections were imaged and displayed with the same parameters. At P0-5, the KCC2 staining was weak and confined to the IPL; at P7, staining was easily detectable in ganglion cell somas; and by P13, it was strong in the OPL, INL, IPL, and GCL. Insets: higher contrast of IPL or OPL. For P1 and P3, the insets were from the area enclosed by - - -. For P7, the IPL inset was taken from a different image to show the stained somas in the GCL (→), and the OPL inset was placed over its correct location in the low contrast image. For P13 and P20, insets show IPL at twice the magnification to show the punctate nature of the stain.

![FIG. 5. Gross morphology of NKCC1-null retina is normal. Confocal fluorescence micrograph of adult NKCC1-null retina (vertical sections) stained with different markers. A type of off-bipolar cell (off-b) labeled with anti-recoverin terminated in the OFF sublamina; the rod bipolar cell (rb) labeled with anti-PKC terminated in sublamina 5 of the IPL; the ON bipolar cells (on-b) labeled with anti-Gαo showed normal high-density up in the INL; the horizontal cell (hc) and amacrine cell (ac) labeled with anti-calbindin had normal appearance with 3 strata in the IPL; the amacrine cells labeled with anti-GABA were numerous in the lower tiers of the INL and more sparse in the GCL; and ganglion cells (gc) labeled with anti-parvalbumin sent processes to all sublaminas of the IPL. Müller cells labeled with an antibody against glutamine synthetase showed radial processes that, as in the adult, stained strongly in the outer nuclear layer and in their terminals in the nerve fiber.](http://jn.physiology.org/10.1152/jn.01224.2007)
layer and weakly in the inner plexiform layer. Müller cells with an antibody against vimentin appeared normal in only two of four retinas where strongly stained radial processes crossed the inner plexiform layer and terminated in the nerve fiber layer. In the other two retinas, occasional processes traveled in the inner plexiform layer laterally; this abnormality was never seen in wild type. Physiologically, NKCC1-null retina exhibited normal spontaneous calcium transients during the first postnatal week (see Fig. 6 in the following text). Thus lack of NKCC1 does not affect the gross morphology and physiology of the retina.

NKCC1 deletion and bumetanide application in early ganglion and amacrine cells do not prevent GABA from evoking a Ca\(^{2+}\) rise

One of the hallmarks of GABA’s excitatory action in developing retinal ganglion and amacrine cells is its ability to depolarize cells via chloride efflux, activate voltage-dependent calcium channels, and consequently evoke a rise in intracellular calcium (Zhang et al. 2006b). If NKCC1 accumulates chloride in retinal ganglion and amacrine cells, perturbing its function would lower intracellular chloride concentration to the level of passive distribution, thus abolishing GABA-evoked depolarization and calcium rise. Using fura-2 calcium imaging, we tested the effect of deleting NKCC1 on the GABA-evoked calcium rise in the P0-5 ganglion cell layer. In both wild-type and NKCC1-null retinas, GABA (50 \(\mu\)M) evoked a calcium rise in over 98% of cells (Fig. 6, A and D). Because all cells behaved in a similar manner, it is safe to conclude that neither amacrine cells nor ganglion cells of the NKCC1-null retinas were affected. GABA-evoked responses were blocked by the ionotropic GABA receptor antagonist, picrotoxin (100 \(\mu\)M; Fig. 6A), suggesting that these responses were due to GABA\(_{\lambda}\)-mediated chloride currents.

One possible explanation for the lack of a deletion effect is the potential upregulation of another chloride accumulator. We therefore tested the effect of the loop diuretics bumetanide on the GABA-evoked calcium rise. Bumetanide reversibly blocks NKCC (both NKCC1 and NKCC2) by competing with the second chloride binding site with a half-inhibitory constant of \(\sim 0.1\) \(\mu\)M. When used at or <10 \(\mu\)M, the inhibition is specific for NKCC (Russell 2000); but to ensure full blockade, we used it at 10–50 \(\mu\)M. Bumetanide was applied to either wild-type or NKCC1-null retina for up to 40 min before testing the GABA response. This did not block the GABA-evoked calcium rise (Fig. 6, B–D). Perhaps intracellular chloride remained high because chloride efflux is slow and the incubation time with bumetanide was not sufficient to reduce it. Thus we facilitated chloride efflux by applying GABA repeatedly and incubating the retina in bumetanide for a longer time (up to 5 h), or by incubating the retina in a mixture of bumetanide and GABA. However, neither condition led to a change in response amplitude (Fig. 6, C and D). Thus neither NKCC1 deletion nor application of NKCC blocker affects GABA’s early excitatory action in retinal neurons.
FUNCTION OF NKCC1 IN RETINA

NKCC1 deletion and bumetanide application do not lower intracellular chloride in developing ganglion and amacrine cells

Next, we examined the effect of NKCC1 blockade and deletion on the intracellular chloride concentration itself. Cells in the ganglion cell layer were loaded with the chloride indicator MEQ, and the basal fluorescence with unperturbed chloride concentration was recorded. The retina was then subjected to a reference buffer solution containing 30 mM chloride. The retina was then exposed to a 30 mM reference solution, which is the steady-state level as indicated in Fig. 7. The relevant fluorescence change (ΔF30Cl/F) was proportional to the difference between the reference concentration and the initial chloride level in the cell (Zhang et al. 2006b). The relevant fluorescence change (ΔF30Cl/F) is higher than 30 mM and a negative value indicates [Cl\(^-\)]i lower than 30 mM. Numbers of animals, experiments, and cells are given in brackets above each group.

Our finding that NKCC1 is not required for chloride accumulation in the immature ganglion and amacrine cells was....

Fig. 7. Deleting or blocking NKCC1 does not affect [Cl\(^-\)], in immature ganglion and amacrine cells [6-methoxy-N-ethylquinolinium iodide (MEQ) chloride imaging]. A–D: corrected MEQ fluorescence change in response to clamping intracellular chloride concentration at 30 mM (see METHODS). Each trace represents recording from a single cell. The initial transient drop is an artifact of high potassium (see Zhang et al. 2006b). The relevant fluorescence change (ΔF30Cl/F) is the steady-state level as indicated in D. △F30Cl/F in D, minimum, midrange and maximum ΔF30Cl/F in this experiment. A new steady-state higher than the baseline fluorescence (as in A and B) indicates that fluorescence increased due to a reduction in chloride. Hence, the cell’s initial chloride level was >30 mM. Responses remaining at the dashed line (0 fluorescence change) indicate that the initial [Cl\(^-\)]i was around 30 mM. In C, retina was incubated in bumetanide (10 μM) and GABA (50 μM) for 40 min prior to testing its MEQ response to 30 mM chloride. E: summary of [Cl\(^-\)]i, measured in experiments similar to those shown above. Each vertical line represents the range of ΔF30Cl/F for a particular experiment as shown in D. ●, mean ± SE of all midrange points within a group. In a Gaussian distribution, midrange = median = mean. - - (no change in MEQ fluorescence) corresponds to [Cl\(^-\)]i = 30 mM; a positive value indicates [Cl\(^-\)]i, higher than 30 mM and a negative value indicates [Cl\(^-\)]i, lower than 30 mM. Numbers of animals, experiments, and cells are given in brackets above each group.

Exposing the retinas to a 30 mM reference solution, which is far from the actual intracellular chloride concentration in the wild-type adult retina, and may thus carry a larger error for the adult than the young retinas (Zhang et al. 2006b).

To test the effect of bumetanide, we incubated the retina with this blocker (10 μM, 40–120 min) with or without GABA (50 μM, to facilitate chloride efflux), then added a reference solution of 30 mM chloride and recorded the MEQ fluorescence. In all P0-5 cells tested, intracellular chloride remained >30 mM (36.2 ± 4.2 mM, P > 0.05 with the same age groups tested above; Fig. 7, C and E). To test whether chloride accumulation depends on external sodium levels, we incubated wild-type retina with sodium-free Ringer (substituted with choline) and found that low sodium did not change the corrected baseline fluorescence (not shown) and thus does not affect intracellular chloride. Taken together, the data strongly suggests that neither NKCC1 nor NKCC2 contributes to chloride accumulation in ganglion and amacrine cells in early development.

Early expression of NKCC1 in mouse inner retina appears restricted to Müller cells

Our finding that NKCC1 is not required for chloride accumulation in the immature ganglion and amacrine cells was...
surprising because it appears inconsistent with the expression profiling that could support the commonly accepted hypothesis that this transporter accumulates chloride in immature neurons. It therefore became essential to examine NKCC1 localization at higher resolution and with specific cell markers. This seemingly simple and crucial experiment proved to be difficult because cells of the young retina don’t necessarily express certain markers and staining for the purpose of ultrastructural localization of NKCC1 revealed additional staining (see following text) that incurred by the DAB-intensification method.

First, we wished to test if NKCC1 localizes to ganglion cells using an antibody against neurofilament that marks adult ganglion cells (Haverkamp and Wässle 2000). However, at P3, this antibody rarely gave staining in somas of the ganglion cell layer (not shown). To test if NKCC1 localizes to Müller cells (the processes of which envelop most cells), we used the known markers of these cells, vimentin and glutamine synthetase (Robinson and Dreher 1990; Haverkamp and Wässle 2000). Staining for NKCC1 at P3 looked similar to that for vimentin with stained endfeet (arrowhead), but the vertical processes in the IPL are not evident by anti-NKCC1 (radial view). Note the punctate staining in the nerve fiber layer (e.g., puncta enclosed by the bracket); these probably correspond to fimbriae. C and D: whole-mount. Immunostaining for NKCC1 at P3 with focus on the nerve fiber layer. Staining in the wild-type (C) carpets the whole layer with much stronger intensity than in the NKCC1-null retina (D). Scanning of these retinas was done sequentially with identical parameters.

**DISCUSSION**

**Localization of NKCC1 in the adult mouse retina**

Using the specific anti-NKCC1 polyclonal antibody, we report here that staining in the adult was mostly restricted to horizontal cells. The inner plexiform layer stained very weakly, and it is currently not known which processes may express NKCC1. Staining of horizontal cells is in agreement with previous reports in monkey and rabbit (Vardi et al. 2000, 2002). Strikingly, dendritic tips of ON bipolar cells stained with the monoclonal T4 antibody (Vardi et al. 2000) remained unstained by the polyclonal antibody used in this study. Be-
cause the staining obtained with the T4 antibody was similar in wild-type and NKCC1-null mouse, we have to conclude that this staining in mouse does not represent NKCC1. However, because the T4 antibody was raised against human NKCC1, its staining in monkey retina may still be valid. Indeed, unlike mouse, monkey retina horizontal cells did stain, and the inner plexiform layer was largely unstained, suggesting that it does not exhibit the same cross-reactivity in monkey as it does in mouse. The polyclonal antibody gave no staining in monkey (unpublished data) or ferret (Zhang et al. 2006a), so the localization of NKCC1 in these species could not be verified. Lack of staining in ON bipolar dendritic tips suggests that mouse ON bipolar dendrites do not express NKCC1, but it is highly likely that they express an NKCC1-like transporter that is recognized by the T4 antibody. This interpretation is supported by physiological recordings that show that ON bipolar dendritic tips have relatively high intracellular chloride concentration, which is not completely diminished by bumetanide (Varela et al. 2005; Duebel et al. 2006).

**NKCC1 does not accumulate chloride in most immature neurons**

We show here that in the first postnatal week, neither the nature of GABA action nor the intracellular chloride concentration in ganglion and amacrine cells is altered by genetic deletion of NKCC1. It might be proposed that genetic deletion of NKCC1 triggers developmental compensation and upregulation of another protein or mechanism that can maintain a high intracellular chloride. However, pharmacologically blocking NKCC transporter in early wild-type retinas with bumetanide did not alter intracellular chloride concentration either. Furthermore, high-resolution confocal microscopy and electron microscopy does not support NKCC1’s presence in plasma membrane of immature retinal neurons.

The extent to which NKCC1 localizes to premature brain neurons is still unclear. In the lateral superior olive, a system where the GABA’s switch was rigorously studied, the major neurons during the period of GABA excitation do not express NKCC1 mRNA (Balakrishnan et al. 2003). In cerebellum, NKCC1 mRNA expression originally thought to localize to Purkinje cells (Plotkin et al. 1997a) was later found to localize to the glial cells surrounding the Purkinje cells (Kanaka et al. 2001). Most studies documenting NKCC1 presence in immature neurons were done at a low resolution. To our knowledge, the function of NKCC1 in normal immature neurons has so far been demonstrated only in certain neocortical neurons (Yamada et al. 2004). Together with the report by Balakrishnan et al. (2003), our data challenge the standing hypothesis that NKCC1 is the primary chloride accumulator in juvenile neurons. Thus a new accumulator for these cells remains to be found.

**Possible candidates for a chloride accumulator in early retinal neurons**

Transporters and exchangers that can accumulate chloride include the kidney-specific Na\(^+\)/Cl\(^-\) cotransporter, the Na\(^+\)/K\(^+\)/Cl\(^-\) cotransporter (NKCC), the Cl\(^-\)/HCO\(_3\)\(^-\) anion exchanger (Boron 2001), pendrin (similar to sulfate transporter, but transports iodine and chloride) (Scott et al. 1999), and possibly certain types of ClC chloride channels, recently shown to transport Cl\(^-\) against its thermodynamic gradient (reviewed by Miller 2006). The Na\(^+\)/Cl\(^-\) cotransporter and NKCC2 are unlikely candidates not only because they were documented to be specific to kidney but also because in our...
experiments, bumetanide did not block chloride accumulation, and sodium-free solution did not affect intracellular chloride concentrations. Also Western blots with two antibodies against NKCC2 were negative in retina (unpublished results).

As for the Cl/HCO₃⁻ anion exchanger AE3, its mRNA appears in the lateral superior olive early in development (Becker et al. 2003). In retina, the long splice variant of this exchanger localizes to adult rat Müller cells and the short splice variant to horizontal cells. The long splice variant expressed by Müller cells was detected in Western blots during the first postnatal week, but it upregulates during development, peaking at the adult (Kobayashi et al. 1994). Furthermore, EₐGABA measured in bicarbonate- and HEPES-based medium are similar, suggesting bicarbonate may not change EₐCl either (Zhang et al. 2006b). Thus this exchanger is not a likely candidate to accumulate chloride in early ganglion cells. Because the rest of the chloride accumulators have not yet been tested in retina, there is presently no obvious candidate for early retinal cells.

Possible function of transient NKCC1 expression in Müller cells

Expression of NKCC1 in Müller cells is higher in the developing retina than in the adult. Downregulation of NKCC1 was also reported in the vimentin-positive radial glial cells of the ventricular zone of the subcortical brain regions. There, NKCC1 is expressed transiently, peaking at E14.5 and disappearing later at E17–P0 (Li et al. 2002). Thus the transient pattern in mouse Müller cells is not an isolated event. Because the radial glial cells are neuron progenitor cells (Noctor et al. 2001), Li et al. (2002) proposed that NKCC1 might regulate cell cycle-related volume change. However, in retina, developing Müller cells are not known to be progenitor cells with the exception of chicken Müller cells that have been implicated in neural regeneration in response to acute damage (Fischer and Reh 2003).

Here we propose an alternative hypothesis that links expression of NKCC1 in Müller cells to GABA excitation in juvenile retina. This hypothesis is based on observation in the adult brain, in which GABA applied to hippocampal pyramidal cells changes extracellular chloride concentrations, and this change can be blocked by bicucullin (Muller et al. 1989). This, together with evidence that cortical glial cells express GABA_A receptors and display a relatively high intracellular chloride concentration of ~40 mM, prompted a suggestion that a delayed activation of the GABA_A receptors on the glial cells would cause a chloride efflux that would raise the extracellular chloride concentration back to normal (Bormann and Kettenmann 1998; MacVicar et al. 1989).

If in retina, as in brain, extracellular chloride concentration changes with extensive activation of GABA_A receptors located on the neurons, then buffering mechanism may be required. We suggest that NKCC1 expressed by Müller cells can contribute to such buffering of extracellular chloride in the developing retina. When a GABA_A channel opens, chloride flows out of the cell and temporarily raises extracellular chloride concentration. This activates NKCC1, which transports the chloride into the Müller cell, thus removing excess of chloride from the extracellular space. Müller cells are ideally positioned to buffer extracellular contents with their intimate interaction with neurons, their radial processes that can serve as a pipeline, and their expanded endfeet that could siphon the chloride into the vitreous humor. As NKCC1 takes up chloride into Müller cells, it could also remove excess potassium from the extracellular space, another potentially hazardous byproduct of neuronal excitation, and thus efficiently maintain a healthy microenvironment. A role for NKCC1 in clearing extracellular potassium was also proposed in cerebral ischemia (Chen and Sun 2005). Because the challenge of extracellular chloride accumulation occurs only in early retinal development when GABA causes chloride efflux, the expression of NKCC1 in Müller cells is necessary only transiently.

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