Altered Renal Distal Tubule Structure and Renal Na\(^+\) and Ca\(^{2+}\) Handling in a Mouse Model for Gitelman’s Syndrome

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Abstract. Gitelman’s syndrome, an autosomal recessive renal tubulopathy caused by loss-of-function mutations in the thiazide-sensitive NaCl co-transporter (NCC) of the distal convoluted tubule (DCT), is characterized by mild renal Na\(^+\) wasting, hypocalciuria, hypomagnesemia, and hypokalemic alkalosis. For gaining further insights into the pathophysiology of Gitelman’s syndrome, the impact of NCC ablation on the morphology of the distal tubule, on the distribution and abundance of ion transport proteins along its length, and on renal tubular Na\(^+\) and Ca\(^{2+}\) handling in a gene-targeted mouse model was studied. NCC-deficient mice had significantly elevated plasma aldosterone levels and exhibited hypocalciuria, hypomagnesemia, and compensated alkalosis. Immunofluorescent detection of distal tubule marker proteins and ultrastructural analysis revealed that the early DCT, which physiologically lacks epithelial Na\(^+\) (ENaC) and Ca\(^{2+}\) (TRPV5) channels, was virtually absent in NCC-deficient mice. In contrast, the late DCT seemed intact and retained expression of the apical ENaC and TRPV5 as well as basolateral Na\(^+\)-Ca\(^{2+}\) exchanger. The connecting tubule exhibited a marked epithelial hypertrophy accompanied by an increased apical abundance of ENaC. Ca\(^{2+}\) reabsorption seemed unaltered in the distal convoluted tubule (i.e., the DCT and connecting tube) as indicated by real-time reverse transcription-PCR, Western blotting, and immunohistochemistry for TRPV5 and Na\(^+\)-Ca\(^{2+}\) exchanger and micropuncture experiments. The last experiments further indicated that reduced glomerular filtration and enhanced fractional reabsorption of Na\(^+\) and Ca\(^{2+}\) upstream and of Na\(^+\) downstream of the DCT provide some compensation for the Na\(^+\) transport defect in the DCT and contribute to the hypocalciuria. Thus, loss of NCC leads to major structural remodeling of the renal distal tubule that goes along with marked changes in glomerular and tubular function, which may explain some of the clinical features of Gitelman’s syndrome.

The renal distal convoluted tubule (DC) and the connecting tubule (CNT) are co-expressed in the late DCT. High amounts of Ca\(^{2+}\) transporting proteins such as the apical calcium channel (TRPV5/ENaC) and the basolateral Na\(^+\)-Ca\(^{2+}\)-exchanger (NCX) have been revealed in the DCT and CNT [reviewed in (6,7)]. Likewise, proteins implicated in renal Mg\(^{2+}\) handling, such as the apical TRPM6 cation channel (8,9) and the basolateral γ subunit of the Na-K-ATPase (10), are highly expressed in the DC.

NCC loss-of-function mutations cause human Gitelman’s syndrome, an autosomal recessive tubulopathy that is characterized by mild renal Na\(^+\) wasting, hypocalciuria, hypomagnesemia, and hypokalemic alkalosis (11). Numerous NCC mutations, occurring throughout the entire coding sequence of the protein, have been described (12,13). When heterologously expressed in *Xenopus laevis* oocytes, mutated NCC proteins are retained in the endoplasmic reticulum (ER) (14,15) or do not exhibit normal NaCl cotransport activity when they reach...
the cell surface (14). Although the molecular and cellular mechanisms that lead to NCC dysfunction in Gitelman’s syndrome are beginning to be understood, many aspects of the pathophysiologic mechanisms that lead to the characteristic phenotype of the disease are still elusive.

For example, compared with patients with other salt-losing tubulopathies (e.g., pseudohypoaldosteronism type I), patients with Gitelman’s syndrome have only mild renal Na\(^+\) wasting and small, although significant, reduction in BP (16), pointing to yet not well-characterized renal adaptive mechanisms that allow compensation for impaired NCC-mediated NaCl reabsorption. Moreover, it is unclear how mutations of NCC affect the renal handling of divalent cations. Experiments on the acute effect of thiazides on microperfused DCT (17), on DCT cell vesicle preparation (18), and on immortalized DCT cells in vitro (19) pointed to increased Ca\(^{2+}\) transport by DCT cells in response to an acute inhibition of NCC function. On the basis of these experiments, it has been hypothesized that the hypocalciuria in Gitelman’s syndrome is due to increased Ca\(^{2+}\) reabsorption by the DCT. Two mechanisms, neither of which has been proved, have been proposed to explain the hypocalciuria: (1) impaired NaCl entry via NCC may lower the intracellular chloride concentration, hyperpolarize the plasma membrane, and subsequently activate voltage-gated Ca\(^{2+}\) channels in the apical plasma membrane; and (2) reduced intracellular Na\(^+\) may stimulate Ca\(^{2+}\) exit across the basolateral membrane as a result of increased activity of the basolateral NCX (20). The hypomagnesemia is even less readily explained. In DCT segments in vivo (21) and in immortalized DCT cells in vitro (22), inhibition of NCC by thiazides stimulates rather than inhibits Mg\(^{2+}\) transport by DCT cells. Therefore, it has been presumed that hypomagnesemia is not related to altered magnesium handling by the DCT itself but might be secondary to the hypokalemia in Gitelman’s patients (22).

Schultheis et al. (23) generated an NCC null mutant mouse model that mimics to a large extent the renal phenotype of the disease. Like affected humans, these mutant mice exhibit hypocalciuria and hypomagnesemia with no apparent signs of hypovolemia as long as the animals are kept on a standard Na\(^+\) diet (16). Kidney Na\(^+\) handling in NCC-deficient mice has been described previously (23). Heterozygous mice (NCC\(^+/−\)) were backcrossed into a homozygous genetic background of C57BL/6 mice (Iffa Credo, Arbresle, France) for >10 generations. Animals were bred in a standard, non-specific pathogen free (SPF) animal facility. Experiments were performed on 2- to 3-mo-old female and, for micropuncture experiments, male wild-type (NCC\(^+/+\)) or null (NCC\(^+/−\)) littermates. All animals had free access to standard lab diet (containing 0.24% Na\(^+\)) and tap water and were housed either in groups of six animals or individually in metabolic cages to allow recording of 24-h urinary volume and ion excretion. All experimental procedures were in accordance with the Guide for the Care and Use of Laboratory Animals (Institute of Laboratory Animal Resources, National Academy of Sciences, Bethesda, MD) and complied with the legal stipulations of the countries in which the experiments were performed.

**Blood Analysis**

For plasma K\(^+\), Ca\(^{2+}\), Mg\(^{2+}\), and aldosterone measurements, mice were anesthetized by intraperitoneal administration of ketamine/xylazine (0.1 and 0.01 mg/g body wt, respectively). Blood was collected by puncture of the retrobulbar venous plexus. Plasma K\(^+\) was measured by automated enzymatic methods (Kodak Biolyzer; Eastman Kodak, Rochester, NY). Plasma Ca\(^{2+}\) and Mg\(^{2+}\) were determined by atomic absorption spectrophotometry (model 3110; Perkin Elmer, Norwalk, CT). Plasma aldosterone concentration was measured by RIA (Kit Aldo RIA, Sanofi Diagnostics, Pasteur, France). Blood collected from retrobulbar venous plexus of awake mice was immediately analyzed for gases and pH.

**Western Blot Analysis**

Kidneys were homogenized in extraction buffer (250 mM sucrose, 150 mM NaCl, 30 mM Tris [pH 7.5], and 0.5 mM PeFabloc; Roche, Rotkreuz, Switzerland). Supernatants from two centrifugations at 1500 × g were pooled and centrifuged at 100,000 × g for 1 h. The pellet was resuspended in 100 μL of extraction buffer. Equal amounts of protein (∼70 μg) from kidneys of wild-type and knockout mice were diluted in reducing sample buffer (4× NuPAGE-LDS; Invitrogen, Basel, Switzerland), heated for 10 min at 70°C, and loaded on a 4 to 12% polyacrylamide gel (Bis-Tris-Gels; NuPAGE; Invitrogen). After electrophoretic separation, proteins were transferred to a polyvinylidene difluoride membrane (BioRad, Reinach, Switzerland). The membrane was blocked for 30 min in 5% nonfat dry milk in a Tris-NaCl-Tween-Buffer (TNT) and was then incubated at 4°C for 16 h with either a rabbit anti-NCC antibody (26) or a rabbit anticanine NCX (Swant, Bellinzona, Switzerland) antibody, each diluted 1:2000 in TNT buffer with 5% dry milk, followed by incubation for 2 to 3 h with a 1:2000 dilution of horseradish peroxidase–conjugated
goat anti-rabbit IgG. Sites of antibody binding were visualized with the ECL Western blotting analysis system (Amersham Pharmaceutica, Otelfingen, Switzerland).

**Light and Electron Microscopy**

The kidneys of anesthetized mice were fixed by vascular perfusion with 3% paraformaldehyde and 0.05% picric acid as described previously (27). Parts of the kidneys were postfixed for 24 h in the same fixative to which 1.0% glutaraldehyde was added. Afterward, the tissue was embedded in Epoxy resin (Epon). Semithin and ultrathin sections were cut with an ultramicrotome (Reichert Jung, Vienna, Austria) and stained with 1% methylene blue and 1% azure II, and lead citrate and uranyl acetate, respectively. Sections were studied with a Polysar microscope (Reichert Jung, Vienna, Austria) and a Philips CM 100 electron microscope, respectively.

**Immunohistochemistry**

The remaining parts of the kidneys were frozen in liquid propane and processed for immunohistochemistry as described previously (27). The following primary antibodies were used: rabbit anti-rat bumetanide-sensitive Na-K-2Cl co-transporter (NKCC2) (28); rabbit anti-rat NCC diluted 1:8000; rabbit anti-rat α-subunit, β-subunit, and γ-subunit of ENaC (29) diluted 1:500 to 1:1000 (α,βENaC) or 1:20,000 (γENaC); rabbit anti-parvalbumin (PV; Swant) diluted 1:2000; guinea pig anti-rabbit TRPV5 diluted 1:500 (30); rabbit anti-canine NCX (Swant) diluted 1:1000; rabbit anti-rat calbindin D28K (CB; Swant) diluted 1:20,000; mouse anti-chicken CB (Swant) diluted 1:40,000; and mouse anti-bovine H⁺-ATPase (31) diluted 1:4.

Binding sites of the primary antibodies were detected with Cy3-conjugated donkey anti-rabbit IgG (Jackson ImmunoResearch Laboratories, West Grove, PA), Cy3-conjugated goat anti–guinea pig (Jackson ImmunoResearch Laboratories), and FITC-conjugated goat anti–mouse IgG (Jackson ImmunoResearch Laboratories), diluted 1:1000, 1:500, and 1:40 in PBS/BSA 1%, respectively. For control of nonspecific antibody binding, the primary antibodies were omitted or replaced by a nonimmune rabbit serum.

**Identification of Tubular Segments in Mice**

Cortical distal tubular segments were identified according to immunohistochemical (Table 1) and standard morphologic criteria that are described elsewhere (25,32). The kidneys of nine mice per group (from two independent breedings) were histologically analyzed by three experienced investigators (J.L., D.L., B.K.), who were blinded to the genotype of the animals. Qualitative judgments regarding tubular morphology and immunostainings were similar for all investigators.

**Morphometric Measurements**

Consecutive cryosections were stained with polyclonal antibodies against PV, NCX, ENaC, and aquaporin 2 (AQP2). Each section was also co-stained with a monoclonal antibody against CB. Overviews were taken from each section with the ×10 objective of a Polysar microscope (Reichert-Jung) using a CDD camera. After printing of the micrographs, distal tubular segments were identified according to their specific antibody-staining pattern (Table 1). Because NCX expression, which was the primary characteristic of the DCT in wild-type mice, was absent in knockout mice, we defined the early DCT for both wild-type and knockout mice by the high abundance of PV and low, if any, NCX and CB immunostaining. The late DCT was defined by the presence of ENaC and strong NCX and CB immunostaining. CNT were defined by coexpression of ENaC and AQP2, intermediate NCX and/or CB immunostaining, and their characteristic location within the cortical labyrinth. The CCD was classified by coexpression of ENaC and AQP2 but undetectable NCX and weak CB immunostaining. The fractional cortical tubular volumes for early DCT, late DCT, CNT, and CCD were determined from the printed micrographs by planimetric point-counting methods according to Weibel (33).

**RNA Isolation and Quantitative PCR**

Total RNA from kidney was isolated using Trizol Reagent (Life Technologies BRL, Life Technologies, Breda, The Netherlands) according to the manufacturer’s protocol. RNA was treated with DNase to prevent contamination of genomic DNA and finally resuspended in diethylpyrocarbonate-treated milliQ water. Total RNA (2 μg) was subjected to reverse transcription using Moloney Murine Leukemia Virus Reverse Transcriptase (Life Technologies BRL) as described previously (34). Expression levels of renal TRPV5, calbindin-D28K, and NCX1 mRNA were quantified by real-time quantitative PCR, using the ABI Prism 7700 Sequence Detection System (PE Biosystems, Rotkreuz, Switzerland). With the use of standard curves, the amount of copy numbers of the target genes in each sample was measured.

**Table 1. Segmental distribution of proteins used for the identification of tubular segments in C57BL/6 mice**

|            | TAL | DCT early | DCT late | CNT | CCD |
|------------|-----|-----------|----------|-----|-----|
| NKCC2      | +++ | +         | +        | +   | +  |
| NCC        | +++ | +         | +        | +   | +  |
| ENaC       | +   | +         | +        | +   | +  |
| AQP2       | +   | +         | +        | +   | +  |
| TRPV5      | +   | +         | +        | +   | +  |
| NCX        | +   | +         | +        | +   | +  |
| CB         | +   | +         | +        | +   | +  |
| PVb        | +   | +         | +        | +   | +  |

*Protein abundance: ++ weak; + intermediate; ++ + strong; TAL, thick ascending limb; DCT, distal convoluted tubule; CNT, connecting tubule; CCD, cortical collecting duct; NKCC2, Na⁺-K⁺-2Cl⁻ cotransporter 2; NCC, Na⁺-Cl⁻ cotransporter; ENaC, epithelial Na⁺ channel; TRPV5, transient receptor potential, vanilloid-receptor related ion channel 5; NCX, Na⁺-Ca²⁺ exchange; CB, calbindin D28k; PV, parvalbumin.

Rarely, weak PV immunostaining is also present in TAL and late DCT cells, whereas strong PV immunostaining is always visible in early DCT cells. These observations are consistent to two previous studies on C57BL/6 mice (27) (9), but are at variance to reports on NMRI mice that described high levels of PV in both early and late DCT (54).
calculated and expressed as a ratio to the hypoxanthine-guanine phosphoribosyl transferase gene. Primers and probes targeting the genes of interest were designed using Primer Express software (Applied Biosystems, Foster City, CA) as described previously (35).

**Clearance and Micropuncture Experiments**

Mice were prepared for micropuncture under inactin/ketamine anesthesia as described (36). For assessment of GFR of both kidneys and of single nephrons, [1H]inulin was infused intravenously. Urine collections were performed using a bladder catheter. The left kidney was prepared for micropuncture. On the kidney surface, the last loop of proximal tubules (LPT) or the DC were identified and punctured for quantitative collections of tubular fluid. Tubular fluid volumes were determined from column length in a constant bore capillary. The concentrations in tubular fluid of Na+ and K+ were determined by a microflow photometer (Department of Pharmacology, University of Tübingen, Germany) (36) and of Ca2+ by a flow-through microfluorometer (NanoFlo; WPI, Sarasota, FL) using Fluo-3 (MoBiTec, Gottingen, Germany) for detection (37).

**Statistical Analyses**

Data are given as means ± SEM. Statistical differences between means were evaluated by unpaired t test (two tails). Differences were considered to be significant at \( P < 0.05 \).

**Results**

*Physiologic Data*

As previously reported (23), NCC+/+ mice exhibited hypocalciuria and hypomagnesemia (Table 2). Moreover, the animals had a mild compensated alkalosis as indicated by the increased plasma bicarbonate concentration. Plasma levels for potassium did not differ between NCC+/+ and NCC−/− mice. Plasma aldosterone levels were significantly elevated when compared with NCC+/+ mice. This is at variance with the initial characterization of NCC−/− mice (23) and might be related to differences in the genetic background (mixed and C57BL/6 in the previous and the present study, respectively) or animal husbandry.

**Lack of NCC Protein in Kidneys of NCC−/− Mice**

To confirm the absence of NCC protein from the kidneys of NCC−/− mice, we performed Western blot analysis and immunohistochemistry with affinity-purified anti-NCC IgG. In NCC+/+ mice, Western blot analysis of kidney homogenates revealed a single band at 190 kD and occasionally weaker additional bands at higher molecular weights, likely representing multimeric NCC complexes (Figure 1a); none of the bands was detectable in NCC−/− mice (Figure 1a). In immunohistochemical experiments with the NCC antiserum, a number of tubular profiles were brightly stained in the renal cortex of NCC+/+ mice but not in kidney sections of NCC−/− mice (Figure 1b).

**Distribution of Transport Proteins and Structure along the Cortical Distal Nephron**

The binding patterns of the antibodies that were used and the structure along the distal nephron of kidney sections of NCC+/+ mice were identical to those described previously (27). Loss of NCC had no apparent effect on TAL cell structure and NKCC2 abundance. In mice of both genotypes, NKCC2 immunostaining was visible in the apical plasma membrane of TAL cells and ceased abruptly at the transition from TAL to DCT (Figure 2).

In NCC+/+ mice, the abrupt termination of NKCC2 and the beginning of NCC expression coincided with a marked rise in epithelial height and strong cytoplasmic PV expression that extended for a considerable length along the DCT. In overviews of the renal cortex, a number of PV-positive tubular profiles were consistently visible in NCC+/+ mice (Figure 2, a and d) but almost absent in NCC−/− mice (Figure 2, b and f). In NCC−/− mice, the epithelium after the point at which NKCC2 expression terminated remained as thin as that of the preceding TAL, and immunostainings revealed none of the

![Figure 1](image-url) NaCl-co-transporter (NCC) protein abundance in kidneys of wild-type and NCC knockout mice. (a) Total kidney homogenates of wild-type (+/+) and knockout (−/−) mice were subjected to SDS-PAGE analysis and incubated with an affinity-purified rabbit anti-mouse NCC antibody, followed by a horseradish peroxidase-coupled donkey anti-rabbit IgG and subsequent ECL detection. (b) Cryosections; overviews of renal cortex of wild-type (+/+) and knockout mice (−/−) immunostained with an affinity-purified rabbit anti-mouse NCC antibody, followed by a Cy3-conjugated goat anti-rabbit IgG. Bar = ~200 μm.
tested proteins, except occasional minute amounts of PV and/or CB (Figures 2f and 3). The cells in this short tubular portion were analyzed further by electron microscopy. The cells revealed a pronounced structural atrophy when compared with the early DCT cells of NCC+/+ mice (Figure 2, g and h). In NCC−/− mice, intercalated cells, identified by their bright immunofluorescent staining with antibodies against the H⁺-ATPase, were consistently found in close proximity to the transition from TAL to DCT (Figure 3, g and h). This is in contrast to wild-type animals, in which the most proximal intercalated cells appear distant from the TAL-DCT transition. Taken together, the findings from immunostaining and morphological studies suggest that the early DCT of knockout mice is drastically atrophied and shortened.

**Intact Late DCT in NCC Knockout Mice**

The short, hypoplastic early DCT segment in knockout mice abruptly transitions to a high epithelium (Figures 3 and 4) with cell nuclei in an apical position and with numerous mitochondria (Figure 4b). By immunostaining, these cells exhibited weak expression of ENaC (Figure 3b) and high expression of TRPV5 (Figures 3d and 4d) and CB (Figures 3f and 4, a, c, and e). AQP2 was detected only in the very last cells of this tubular portion (Figure 4, e and f). A few intercalated cells, apparent as unstained dark spots within the brightly CB-immunostained epithelium (Figure 5e), were observed consistently. By these collected features, the epithelium corresponded to the late DCT in wild-type mice. In knockout mice, it showed no apparent signs of structural atrophy.

**Hypertrophied CNT in NCC Knockout Mice**

The late DCT of knockout mice is followed by a tubular portion that corresponded by its antibody staining pattern and histotopographic localization to the CNT in wild-type mice (Table 1, Figures 5 and 6). In comparison with wild-type mice, CNT profiles from knockout mice revealed a marked epithelial hypertrophy (Figure 6, a and b) that was accompanied by an increased apical abundance of all three ENaC subunits, suggesting enhanced Na⁺ transport rates. In wild-type mice, α-, β-, and γENaC subunits were seen almost exclusively in the cytoplasm of CNT cells, whereas in knockout mice, they were shifted toward the apical plasma membrane as shown exemplary for βENaC in Figure 6, c and d. The apical translocation of α-, β-, and γENaC was limited to the CNT and not visible in the CCD of NCC knockout mice as visible for γENaC in Figure 5f.

Figure 2. Early distal convoluted tubules (DCT) in the renal cortex of wild-type (+/+), and NCC knockout (−/−) mice. (a through f) Cryosections. (g and h) Epon thin sections. (a and b) Overviews of the kidney cortex immunostained by a polyclonal rabbit anti-rat parvalbumin (PV) antiserum followed by a Cy3-conjugated goat anti-rabbit IgG. (c through f) Transitions from thick ascending limb (TAL) to early DCT shown in consecutive cryosections (c and d, and e and f) stained either by a rabbit anti-rat Na-K-2Cl co-transporter (NKCC) antiserum or by a rabbit anti-rat PV antiserum, followed by a Cy3-conjugated goat anti-rabbit IgG; bright apical NKCC2 immunostaining characterizes the TAL (T) and ceases abruptly (arrows) at the transition to the early DCT (D), exactly where PV immunolabeling starts. (g and h) Electron microscopic images of early DCT cells from wild-type (+/+) and knockout (−/−) mice; transitions from TAL (T) to DCT (arrows) were identified on semithin sections (insert in h) and analyzed at the electron microscopic level in the successive ultrathin section. The DCT cell shown in h is marked by an asterisk in the insert. In wild-type mice, the early DCT cells are conspicuous by the dense alignment perpendicular to the basement membrane of elongated mitochondrial profiles, narrowly enveloped by basolateral plasma membranes. The cell nucleus is located on top of the row of mitochondria, just below the apical plasma membrane (g). In knockout mice, the height of the early DCT cells is approximately one third that of the cells in wild-type mice, with only a few mitochondria, basolateral plasma membrane infoldings, or apical microprojections (h). Bars = ~200 μm in b, ~20 μm in f, ~2 μm h.
Morphometry

The fractional cortical tubular volumes of the early DCT, the late DCT, the CNT, and the CCD in wild-type and knockout mice (Figure 6) were consistent with the above-described qualitative observations. In knockout mice, the fractional volume for the early DCT was drastically lower than in wild-type mice. The fractional volumes of the late DCT were similar in both genotypes. The fractional volume of the CNT was significantly greater in knockout mice than in wild-type mice. The CCD volume was similar for mice of both genotypes. Although not specifically addressed in the present study, the morphologic changes along the CNT most likely comprise epithelial hypertrophy and hyperplasia. Numerous previous studies revealed that cellular hypertrophy and hyperplasia both contribute to the adaptation of the distal nephron to an enhanced tubular workload [reviewed in (1,25)].

Abundance of Ca\(^{2+}\) Transporting Proteins

The expression of TRPV5 and NCX mRNA, as well as the protein abundance of NCX, was not different between wild-type and NCC-deficient mice (Figure 7). Consistent with previous studies (26,34,37), TRPV5 could not be revealed by Western blot analysis of kidney homogenates, perhaps because of the comparably low abundance of the channel in total kidney preparations. Immunofluorescence revealed a slightly decreased abundance of TRPV5 and NCX along the luminal and basolateral membrane, respectively, of late DCT and CNT of NCC\(^{-/-}\) mice that became most apparent at high antibody dilutions (Figure 7c). The lowered TRPV5 and NCX abundance in individual late DCT and CNT cells may explain the unchanged expression levels of TRPV5 and NCX in total kidney homogenates despite the significant hypertrophy of the CNT in NCC-deficient mice.

Clearance and Micropuncture Experiments

NCC\(^{-/-}\) mice presented normal mean arterial BP but lower GFR and as a consequence lower glomerular filtration of Na\(^{+}\), K\(^{+}\), and Ca\(^{2+}\) than in NCC\(^{+/+}\) mice (Table 3). The lowered GFR but unaffected fractional whole kidney tubular reabsorption of Na\(^{+}\) and Ca\(^{2+}\) (Figure 8) resulted in a modestly reduced urinary excretion of Na\(^{+}\) and Ca\(^{2+}\) in NCC\(^{-/-}\) mice compared with NCC\(^{+/+}\) mice (Table 3). Under balanced conditions when urinary Na\(^{+}\) excretion reflects body Na\(^{+}\) intake, it...
is expected that urinary Na⁺ excretion is not different between NCC−/− and NCC+/+ mice. Thus, the neurohumoral activation associated with anesthesia and surgery may have induced a modestly greater renal Na⁺ retention in NCC−/− mice. This could be the consequence of the documented CNT enlargement that may have created a Na⁺ transport machinery in the CNT of NCC−/− mice that is more responsive to neurohumoral stimulation. Micropuncture experiments confirmed that single-nephron GFR was lower in NCC−/− than in NCC+/+ mice in collections from both the last loop of the proximal tubule on the kidney surface (LPT; 5.4 ± 0.2 versus 6.7 ± 0.4 ml/min; P < 0.05) and DC (5.2 ± 0.4 versus 6.7 ± 0.4 ml/min; P < 0.05). As depicted in Figure 8, fractional delivery of fluid, Na⁺, K⁺, and Ca²⁺ to the LPT were reduced, and thus fractional reabsorption up to this site increased in NCC−/− compared with NCC+/+ mice. Fractional delivery of fluid and Na⁺ remained reduced in NCC−/− versus NCC+/+ mice up to the DC, whereas the fractional delivery of K⁺ and Ca²⁺ to this site was not different between genotypes. The lower fractional delivery of Na⁺ but constant fractional delivery of K⁺ to the DC puncture sites resulted in a raised K⁺ to Na⁺ ratio in the tubular fluid in the DC of NCC−/− mice (Figure 9). These findings and the persistently greater K⁺ to Na⁺ ratio in the urine of NCC−/− than in NCC+/+ mice point to an enhanced functional activation of the aldosterone-sensitive distal nephron in NCC−/− mice (Figure 9).

Because K⁺ secretion and Na⁺ reabsorption occur along the distal nephron sites accessible to micropuncture (together with water reabsorption in CNT and CCD), the distal luminal K⁺ to Na⁺ ratio was used as an indicator of the distal collection site (Figure 10). Consistent with intact Ca²⁺ reabsorption along DC and confirming previous experiments that related fractional Ca²⁺ delivery to luminal K⁺ concentration (37), both NCC+/+ and NCC−/− mice showed a rapid fall in fractional Ca²⁺ delivery with increasing ratios of K⁺ to Na⁺ (Figure 10). The shift of the curve to the right in NCC−/− mice may reflect the greater functional activation of the aldosterone-sensitive distal nephron upstream from the puncturing sites. Consistent with some Ca²⁺ reabsorption between late DC accessible to micropuncture and urine, fractional deliveries of Ca²⁺ in late DC (high K⁺ to Na⁺ ratio) were modestly higher than values found in urine for both genotypes.

**Discussion**

In this study, we used morphologic, biochemical, and functional techniques to determine the impact of genetic NCC ablation on the morphology and the function of the renal distal tubule. Our data reveal a pronounced epithelial remodeling of the DCT and CNT in NCC-deficient mice that likely reflects the preceding DCT portion and shows easily detectable AQP2. The labeling of tubular basement membranes and interstitial cells in c and e is due to binding of the secondary FITC-labeled anti-mouse IgG to endogenous mouse immunoglobulins. Bars = ~20 μm.

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**Figure 4.** Immunofluorescent (a, c through f) and morphologic (b) characterization of late DCT in NCC knockout mice. (b) Epon semithin section; all others are cryosections immunostained with rabbit antiserum (a) and mouse monoclonal antibodies (c and e) against CB, affinity-purified guinea pig antiserum against TRPV5 (d), or affinity-purified rabbit anti-rat aquaporin-2 (AQP; f). c and d, e and f are double immunostainings; the transition (arrows in a and b) from early (1) to late (2) DCT is characterized by the sharp increase in CB immunostaining and in the epithelial cell height (a and b). The late DCT (2) has high cytoplasmic CB (a, c, and e) and apical TRPV5 expression (d) but lacks detectable AQP2 along most of its lengths (f). Only at the transition from late DCT (2) to CNT (3), single cells express high CB levels and AQP2 (f), consistent with previous findings in wild-type mice showing that the very last NCC-positive DCT cells coexpress AQP2 [see Figure 3 in (25)]. The distally localized connecting tubule (3) exhibits weaker CB and TRPV5 expression than...
altered sodium transport activities in the respective segments. Moreover, our data suggest that altered renal Na\(^+\) handling upstream and downstream of the DCT provides compensation for the Na\(^+\) transport defect in the DCT and that hypocalciuria in NCC-deficient mice is primarily the consequence of altered renal Ca\(^{2+}\) handling upstream of the DCT.

Numerous studies, particularly on the renal DCT, revealed the close interrelationship between transepithelial ion transport activity and epithelial structure [reviewed in (1,32)]. Prolonged increases in NaCl transport rates in the DCT epithelium are followed by a notable epithelial hypertrophy, whereas a prolonged decrease in transport rates leads to DCT hypotrophy with reductions of mitochondrial volume and basolateral membrane area (1,32). In NCC-deficient mice, the alterations in epithelial structure of the DCT are compatible with those that result from markedly reduced transepithelial transport rates. We found a decrease in the fractional cortical tubular volume for the entire DCT (early and late parts) from ~13% in wild-type mice to ~4.5% in knockout mice. This reduction in fractional volume matches well the ~60% decrease of DCT cell number as detected previously by electron microscopic analysis (23). Remarkably, the atrophy of the DCT is limited to its early portion, which is almost absent from the kidneys of NCC\(^{-/-}\) mice. In contrast, the late DCT of NCC\(^{-/-}\) mice seems structurally intact and retains its typical DCT cell morphology and, with the exception of the NCC, the expression of ion transport proteins. This is consistent with our previous findings in thiazide-treated rats. In these rats, only the early DCT but not the late DCT undergoes apoptosis in response to NCC inhibition by thiazide treatment (38). The presence of additional ion transporters (e.g., ENaC and TRPV5 and possibly others) may enable the late DCT cells to escape the detrimental effect of the genetic or pharmacologic loss of NCC function.

Although ~5% of the filtered sodium load is thought to be reabsorbed in the DCT by NCC, NCC-deficient mice show only little, if any, renal salt wasting (23). As long as the mice are kept on a sufficient Na\(^+\) intake, the BP remains in the normal range and the mice show no apparent signs of hypo-
The reduced GFR in NCC knockout mice is consistent with previous micropuncture and microperfusion studies that reported a decline in the GFR in response to pharmacologic inhibition of NCC by chlorothiazide (17,40). The reason for the reduced GFR in response to genetic or pharmacologic ablation of NCC function is unclear. It is probably not related to a direct activation of the tubuloglomerular feedback mechanism, because the primary Na\(^+\) transport defect lies downstream of the macula densa. Also extracellular volume depletion cannot account for the significant drop in GFR. The BP of NCC-deficient and wild-type mice did not differ significantly in the present study or in a previous study (23). Likewise, thiazide diuretics reduce the GFR even when extracellular volume depletion is prevented by intravenous replacement of salt and fluid losses (17). Whatever the underlying mechanism is, the reduced GFR seems not to be sufficient to compensate fully for the NCC loss, because aldosterone-dependent stimulation of ENaC-mediated sodium transport seems to contribute as well. Knepper’s group (41) showed by immunoblotting-techniques that the abundance of a lower molecular weight form of the \(\gamma\)-subunit of ENaC is increased in kidneys of NCC\(^{-/-}\) mice, whereas the abundances of the major apical sodium transporting proteins of the proximal tubule and the TAL (NHE3 and NKCC2, respectively) are unchanged. The low molecular weight form of \(\gamma\)ENaC has been proposed to be indicative of enhanced ENaC activity and to represent \(\gamma\)ENaC subunits cleaved by luminal proteases (42). The observed hypertrophy of the CNT, the increased apical localization of ENaC along the CNT, and the enhanced Na\(^+\)-K\(^+\) exchange along the DC establish that ENaC-mediated Na\(^+\) reabsorption in the CNT is increased in the kidneys of NCC-deficient mice. The elevated plasma aldosterone levels most likely play a role in these adaptive changes in the CNT. Aldosterone stimulates Na\(^+\) transport in the renal collecting system (43), induces CNT and CD cell hypertrophy [e.g., (44,45)], shifts ENaC from intracellular compartments to the apical plasma membrane (42,46), and stimulates Na\(^+\) and K\(^+\) exchange in the CNT and CD (43). Surprisingly, we found no morphologic evidence for enhanced ENaC activity in the collecting ducts of NCC\(^{-/-}\) mice. In mice of both genotypes, ENaC subunits were predominately localized at intracellular sites of CCD cells, and the fractional CCD volume did not differ between wild-type and NCC-deficient mice. These morphologic data do not exclude a stimulation of ENaC in the CCD, but they are consistent with previous patch-clamp (47), ion transport (48–50), and immunohistochemical (4,46) studies that revealed a several-times higher sodium transport rate and apical activity/abundance of ENaC in the CNT than in the CCD. The salient importance of nephron portions upstream of the CD for the maintenance of sodium homeostasis was highlighted by the recent development of mice with a targeted inactivation of \(\alpha\)ENaC only in the CD but not in the CNT and late DCT (51). Unlike mice and humans with generalized inactivation of ENaC, these mice are able to maintain Na\(^+\) homeostasis even when challenged by sodium restriction (51).
Sodium transport via ENaC is electrogenic and coupled to K+ secretion via luminal K+ channels such as ROMK. In fact, conditions of an increased ENaC activity (e.g., hyperaldosteronism) are often associated with renal K+ losses (43). Likewise, patients with Gitelman’s syndrome frequently develop hypokalemia that can be corrected by treatment with mineralocorticoid-receptor antagonists (52). Despite the apparent hyperaldosteronism and the activation of ENaC, NCC-deficient mice have normal plasma K+ levels. The reason for the absence of hypokalemia is unclear but could be related to species differences in distal nephron potassium handling or to a comparatively higher dietary K+ intake in mice than in humans.

The structural changes along the DCT and CNT may also have significant impact on the handling of ions other than Na+. Micropuncture and microperfusion experiments reported high transepithelial Ca2+ and Mg2+ transport rates along the DCT [reviewed in (1,2)]. The marked atrophy of the DCT in NCC−/− mice suggests a marked reduction of the luminal plasma membrane area available for transepithelial cation transport. This might explain the renal Mg2+ wasting but seems to be at odds with the reduced urinary Ca2+ excretion in NCC-deficient mice, which is thought to result from an increased Ca2+ reabsorption in the DCT (see introduction). Micropuncture studies in wild-type and TRPV5-deficient mice demonstrated that intact Ca2+ reabsorption in the DC requires the presence of TRPV5 (37). In wild-type mice, the highest abundance of TRPV5 as well as NCX and plasma membrane Ca2+-ATPase (PMCA) is found in the late DCT (27), which remains intact in NCC−/− mice. Thus, the selective atrophy of the early DCT does not necessarily rule out the hypothesis of an increased Ca2+ transport in the DCT. However, chronic stimulation of transepithelial Ca2+ transport would be expected to go along with an enhanced expression of the Ca2+ transporting proteins. Indeed, stimulation of distal nephron Ca2+ transport by calcitriol is associated with an induction of distal nephron Ca2+ transporting proteins (34). Conversely, inhibition of Ca2+ reabsorption by targeted deletion of TRPV5 is associated with reduced expression of CB and NCX (37). In the present study, reverse transcription–PCR, Western blot, and immunohistochemistry did not detect any increased abundance of NCX and TRPV5 in kidneys of NCC-deficient mice. Moreover and most important, the in vivo micropuncture experiments indicate similar Ca2+ reabsorption rates along the DC of wild-type and NCC knockout mice. In mice of both genotypes, fractional Ca2+ delivery decreases steeply at DC

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**Table 3. Blood pressure, glomerular filtration and urinary excretion of NCC+/+ (n = 7) and NCC−/− (n = 7) mice**

|                        | NCC+/+ | NCC−/− |
|------------------------|--------|--------|
| Body wt (g)            | 27.2 ± 0.5 | 28.0 ± 0.6 |
| Mean arterial pressure (mmHg) | 99 ± 3  | 97 ± 3  |
| GFR (μl/min/g bw)      | 6.0 ± 0.5 | 4.3 ± 0.3a |
| Na+-filtered (μmol/min/g bw) | 0.91 ± 0.09 | 0.66 ± 0.04a |
| K+-filtered (nmol/min/g bw) | 38 ± 3  | 26 ± 3a  |
| Ca2+-filtered (nmol/min/g bw) | 11.0 ± 0.1 | 7.6 ± 0.6a |
| Urinary flow (nl/min/g bw) | 79 ± 14 | 52 ± 8  |
| Urinary Na+ excretion (nmol/min/g bw) | 5.5 ± 1.1 | 2.9 ± 0.6a |
| Urinary K+ excretion (nmol/min/g bw) | 8.6 ± 1.1 | 6.7 ± 0.7 |
| Urinary Ca2+ excretion (pmol/min/g bw) | 29 ± 7  | 12 ± 3a  |

* P < 0.05.

Values are means ± SEM.

**Figure 8. Fractional delivery of fluid, Na+, K+, and Ca2+ to the last surface loop of proximal tubule (LPT), distal convolution (DC), or urine (U) in NCC+/+ and NCC−/− mice. Values are means ± SEM from seven mice (U), 29 to 34 nephrons (LPT), or six to nine nephrons (DC) per group. *P < 0.05 versus NCC+/+ values.**
sites with a low $K^+$ to $Na^+$ ratio (presumably late DCT), whereas little further $Ca^{2+}$ reabsorption occurs in DC sites with higher $Na^+$ versus $K^+$ exchange (presumably CNT; Figure 10). The $K^+$ to $Na^+$ ratio increases from the DCT to the end of the CNT as a result of ENaC activity. The observed ion transport profile is consistent with the relative abundance of $Ca^{2+}$-transporting proteins along the late DCT (very high) and CNT (lowered) and strongly suggests unaffected $Ca^{2+}$ transport rates in both segments of NCC $-/-$ mice. Thus, the hypocalciuria is unlikely to be related to altered $Ca^{2+}$ transport along the DC.

Extracellular volume contraction is a well-known stimulus for paracellular $Ca^{2+}$ reabsorption in the proximal tubules (53). Recent studies by Nijenhuis et al. (26) indicated that this mechanism may explain the hypocalciuria induced by chronic thiazide treatment. Thiazide-induced hypocalciuria, present despite drastically downregulated distal tubular $Ca^{2+}$ transporting proteins, was completely prevented by oral replacement of diuresis-related salt and fluid losses (26). NCC-deficient mice have elevated plasma aldosterone concentration, pointing to some degree of extracellular volume depletion. Dietary $Na^+$ restriction has been reported to aggravate the volume depletion and to augment the hypocalciuria in NCC-deficient mice (23).

In the present micropuncture study on mice, the observed hypocalciuria in NCC-deficient mice was the consequence of a reduced GFR and thus lower $Ca^{2+}$ filtration and of an increased fractional proximal reabsorption of $Ca^{2+}$ that was associated with an increased fractional $Na^+$ reabsorption at this site. The enhanced fractional proximal reabsorption may reflect a primary tubular stimulation but could also, at least in part, be secondary to the reduced GFR, consistent with imperfect glomerulotubular balance. Nevertheless, considering lower fractional $Ca^{2+}$ delivery to the late proximal tubule but similar fractional excretion in urine in NCC knockout than in wild-type mice, the absolute and fractional $Ca^{2+}$ reabsorption downstream of the late proximal tubule, i.e., in the distal nephron segment including the DCT, was actually reduced in mice deficient of NCC (54).

In summary, the genetic loss of NCC function leads to remarkable structural and functional changes in the kidney. Our data provide strong evidence that reduced glomerular filtration and enhanced fractional reabsorption of $Na^+$ upstream of the DCT and enhanced ENaC-mediated sodium transport downstream of the CNT plays an important role in compensating for the $Na^+$ transport defect in the DCT. Our data do not provide any support for enhanced $Ca^{2+}$ transport by the DCT but indicate that reduced glomerular filtration and possibly stimulation of proximal reabsorption are causative for the hypocalciuria in NCC-deficient mice and perhaps in patients with Gitelman’s syndrome.

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