Chronic Metabolic Acidosis Enhances NHE-3 Protein Abundance and Transport Activity in the Rat Thick Ascending Limb by Increasing NHE-3 mRNA

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Abstract

Chronic metabolic acidosis (CMA) is associated with an adaptive increase in the bicarbonate absorptive capacity of the rat medullary thick ascending limb (MTAL). To specify whether NHE-3, the apical MTAL Na/H exchanger, is involved in this adaptation, NHE-3 mRNA was quantified by a competitive RT-PCR using an internal standard which differed from the wild-type NHE-3 mRNA by an 80-bp deletion. CMA increased NHE-3 mRNA from 0.025±0.003 to 0.042 ± 0.009 amol/ng total RNA (*P* < 0.005). NHE-3 transport activity was measured as the initial proton flux rate calculated from the Na-dependent cell pH recovery of Nadepleted acidified MTAL cells in the presence of 50 μ M HOE694 which specifically blocks NHE-1, the basolateral MTAL NHE isoform. CMA caused a 68% increase in NHE-3 transport activity (P < 0.001). In addition, CMA was associated with a 71% increase in NHE-3 protein abundance (P <0.05) as determined by Western blot analysis on MTAL membranes using a polyclonal antiserum directed against a cytoplasmic epitope of rat NHE-3. Thus, NHE-3 adapts to CMA in the rat MTAL via an increase in the mRNA transcript that enhances NHE-3 protein abundance and transport activity. (J. Clin. Invest. 1997. 99:24-30.) Key words: Na/H exchange • NHE-3 • acidosis • kidney

Introduction

Metabolic acidosis is associated with a number of adaptive changes in renal tubules that contribute to increased urinary net acid excretion. Chronic acid loading produces a large increase in the bicarbonate absorptive capacity of the proximal tubule (1). This adaptation occurs via parallel increases in V_{max}

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J. Clin. Invest. © The American Society for Clinical Investigation, Inc. 0021-9738/97/01/0024/07 \$2.00 Volume 99, Number 1, January 1997, 24–30 of the apical Na⁺/H⁺ antiporter (2–6) and vacuolar H⁺-ATPase (7), and basolateral Na⁺: $3HCO_3^-$ cotransporter (2, 3). Current evidence strongly suggests that NHE-3, which encodes an amiloride-resistant Na⁺/H⁺ exchanger (NHE),¹ is an apical isoform of the NHE family and plays a significant role in NaHCO₃ and NaCl absorption in the proximal tubule (8–10). Studies performed in a cell line derived from proximal tubules of the opossum kidney, the OKP cells, showed that both amiloride-resistant Na⁺/H⁺ exchange activity and NHE-3 mRNA adapt to acid incubation (11). Recent in vivo studies have shown that chronic metabolic acidosis (CMA) increases NHE-3 protein abundance in cortical brush border vesicles (12, 13). However, no change in cortical NHE-3 mRNA abundance was observed when assayed by Northern blot analysis (12). Thus, although NHE-3 is probably involved in the acidinduced apical antiport activation, the mechanisms of this adaptation, related to increased levels of mRNA or to posttranslational regulation, remain unclear.

CMA is also associated with an adaptive increase in bicarbonate absorptive capacity in the rat medullary thick ascending limb (MTAL) (14). Bicarbonate absorption is mediated mainly via an apical Na⁺/H⁺ antiporter in MTALs from both control and acidotic rats (15–17). We have shown recently, in microdissected rat MTAL, that NHE-3 mRNA is strongly expressed and NHE-1 mRNA is also detected, whereas NHE-2 and NHE-4 mRNAs are not detected (18). Furthermore, recent data have demonstrated that NHE-1 and NHE-3 are localized to the basolateral and apical plasma membrane of MTAL cells, respectively (9, 19–21).

The purpose of this study was to determine whether NHE-3 is directly involved in the in vivo adaptation of the MTAL bicarbonate absorption observed during CMA, and which mechanisms are involved. Thus, we developed a very sensitive competitive reverse transcription-PCR (RT-PCR) which allowed us to quantify NHE-3 mRNA in MTAL cells freshly isolated from NH₄Cl-induced acidotic rats. NHE-3 transport activity and NHE-3 protein abundance were also determined in the same preparations. The results show for the first time, in MTAL tubular cells, that CMA increases the level of mRNA transcript for NHE-3, associated with an increase in abundance and functional activity of the protein.

Methods

Animal treatment. Experiments were performed on male Sprague-Dawley rats (200–300 grams body wt). The animals were allowed free

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^{1.} *Abbreviations used in this paper:* CMA, chronic metabolic acidosis; dNTP, deoxyribonucleotide; MTAL, medullary thick ascending limb; NHE, Na⁺/H⁺ exchanger; RT, reverse transcription.

access to food and drinking solution up to the time of the experiments. In each series, a group of experimental animals was compared directly with controls that were obtained from the same shipment and studied during the same period of time. CMA was obtained as follows: rats with CMA drank 0.28 M NH₄Cl for 6 d; control rats drank distilled H₂O. Both groups received standard rat chow ad libitum.

On the day of the experiment, rats were anesthetized with pentobarbital sodium. In most animals, blood was collected by aortic puncture for analysis of plasma electrolytes and the kidneys were rapidly removed, weighed, and immersed into ice-cold Hanks' solution containing (mM): 137 NaCl; 5.4 KCl; 25 NaHCO₃⁻; 0.3 Na₂HPO₄; 0.4 KH₂PO₄; 0.5 MgCl₂; 10 Hepes; 5 glucose; 1 leucine; and 1 mg/ml BSA.

Isolation of MTAL tubules. The method used has been described previously in detail (17, 22, 23). Briefly, small tissue pieces of inner stripe of outer medulla were subjected at 37° C to successive 7-min periods of collagenase digestion (0.75 grams/liter), which minimized the exposure time of the tubules to collagenase. The MTAL tubules were harvested by sieving the supernatants through a 75-µm opening nylon mesh to separate MTAL fragments from isolated cells and small fragments of other medullary tissues and resuspended in an appropriate volume of the desired medium. The suspensions obtained from control and acidotic rats as observed by light microscopy were similarly composed almost exclusively of MTAL tubular fragments (> 95%).

Measurement of pHi. MTAL cells were loaded with the pH-sensitive probe BCECF by incubation for 15 min at 37°C in a CO₂-free medium containing (mM): 140 Na; 3 KCl; 1 CaCl₂; 1 MgCl₂; 1 K₂HPO₄; 10 Hepes; 5 glucose; 1 leucine; 0.1 mg/ml BSA, pH 7.4, bubbled with 100% O₂ and containing 5 µM BCECF acetoxymethyl ester (AM; dissolved in DMSO and stored at -20°C). The BCECF-loaded tubules were then washed by gentle centrifugation to remove the extracellular dye and resuspended in the same medium. Just before each run, aliquots of the MTAL tubules were sodium depleted and acidified by three washings and gentle centrifugations in a CO₂-free and sodium-free medium containing (mM): 140 N-methyl-D-glucamine chloride (NMDG-Cl); 3 KCl; 1 CaCl₂; 1 MgCl₂; 1 K₂HPO₄; 10 Hepes; 5 glucose; 1 leucine; 0.1 mg/ml BSA, pH 7.4. The MTAL cells were then diluted into cuvettes containing 2 ml of the experimental medium to reach a final cytocrit of \sim 0.1 vol%. BCECF fluorescence was monitored by use of a spectrofluorometer (RF-5000; Shimadzu Corp., Tokyo, Japan) equipped with water-jacketed temperaturecontrolled cuvette holder and magnetic stirrer. Fluorescence intensity was recorded at one emission wavelength, 530 nm, whereas the excitation wavelengths alternated automatically at 2-s intervals between two wavelengths, 500 and 450 nm. The values of the 500-450-nm fluorescence ratio were converted into pHi values with use of calibration curves that were established after each experimental run: cells were permeabilized with Triton X-100 (0.025 g%), and the relationship between extracellular BCECF fluorescence and medium pH was established.

Calculations of H⁺ *fluxes.* To quantitate Na⁺/H⁺ exchange activity, proton flux rates $(J_{H}^{+}, \text{pmol/min/mm of tubule length})$ were calculated as $J_{H}^{+} = dpHi/dt \times \beta i \times V$, where dpHi/dt is the sodium-dependent initial rate of pHi recovery (pH units/min), βi is intrinsic intracellular buffering power (mmol/liter \cdot pH unit), and V is cell volume (nl/mm tubule length).

To initiate a sodium-dependent pHi recovery, sodium-depleted acidified MTAL cells were abruptly added into an isosmotic CO₂-free medium containing 30 mM Na (30 mM NMDG-Cl replaced with 30 mM NaCl). The first 10 s of time course of this sodium-dependent pHi recovery was fitted to a linear equation to calculate the initial rate of change of pHi (dpHi/dt); correlation coefficients for these linear fits were > 0.96.

 β i was determined from immediate alkalinization of sodiumdepleted acidified MTAL cells caused by addition of 15 mM NH₄Cl in the Na-free medium containing 10 mM barium and 0.5 mM amiloride to block NH₄⁺ MTAL carriers as previously described (23, 24).

Cell volume was determined by measuring tubule diameters under light microscopy as described (25, 26). *RNA extraction, RT, and PCR.* Total RNA was extracted from MTAL cells using the method of Chomczynski and Sacchi as previously described (27) and quantified by the measure of its optical density at 260 nm.

Primers used in this study for cDNA synthesis and PCR amplification were chosen from the published rat NHE-3 cDNA sequences (28) and are located near the 3' end of the coding region, where there are low sequence homologies between the four isoforms. cDNA was synthesized from RNA samples with 200 U Moloney murine leukemia virus reverse transcriptase (MMLV), 20 pmol of downstream primer, 4 μ g of yeast transfer RNA, 2.5 mM each deoxyribonucleotide (dNTP), 10 mM DTT, 2 U of RNase inhibitor in 22 μ l reaction buffer containing 50 mM Tris-HCl, 75 mM KCl, and 3 mM MgCl₂ at 37°C for 60 min. Each reaction was performed in parallel with an otherwise identical one that contained no reverse transcriptase.

For PCR, 10 μ l of the cDNA solution was supplemented with 5 μ l of 10× PCR buffer, 5 μ l of a 25 mM MgCl₂ solution, 10 pmol of each primer, 1 μ l of a 25 mM dNTP solution, and 1.25 U *Taq* polymerase in a final volume of 50 μ l. Samples were overlayered with mineral oil, denatured at 94°C for 4 min followed by 30 cycles consisting of denaturing at 94°C (1 min), annealing at 60°C (1 min), and extension at 72°C (1.5 min). PCR was completed by a final extension step of 10 min at 72°C. The PCR products were size fractionated on 1.5% agarose gels stained with ethidium bromide.

Internal standard preparation and in vitro transcription. 500 ng of total RNA extracted from MTAL suspension was submitted to RT-PCR using the following primers corresponding to bases 1,073–1,103 and 2,466–2,486 of the NHE-3 coding region giving a fragment of 1,414 bp. The sequences of the primers were: 5'-CCACCACCGT-GCGCTACACTA-3' (sense) and 5'-GAGTCAGGCGGCGGAA-GTTGC-3' (antisense). A third oligonucleotide which was used as an amplification product–specific probe encompassed bases 2,052–2,082. The sequence of this oligonucleotide probe was 5'-GCCGACT-TAAAGGACTCCAGGCGCTTCCG-3'.

The PCR product was analyzed by agarose gel electrophoresis, recovered from gel with Geneclean (Bio 101, La Jolla, CA), phosphorylated with T4 polynucleotide kinase, and the ends were filled in with T4 DNA polymerase.

The NHE-3 fragment (1,414 bp) was then subcloned into the EcoRV site of linearized pBluescript (BSKS+) (Stratagene Inc., La Jolla, CA). To further characterize the DNA template, digestions by a set of restriction enzymes followed by Southern blot analysis using a ³²P-labeled oligonucleotide probe specific for NHE-3 cDNA was performed and gave products of the expected size (data not shown). Subsequently, NHE-3 DNA plasmid was submitted to digestions with Bpu1101I and BspMI restriction enzymes (bp 2,082 and bp 2,160 of the NHE-3 cDNA sequence, respectively), blunt-ended, and ligated, resulting in a 80-bp deletion of the 1,414-bp NHE-3 DNA insert.

Sense RNA was transcribed from 4 μ g of EcoRI-cut DNA template using an RNA transcription kit (Stratagene Inc.). Briefly, transcription was performed in a 100- μ l reaction volume containing 40 mM Tris (pH 7.5), 8 mM MgCl₂, 2 mM spermidine, 50 mM NaCl, 0.4 mM rUTP, 0.4 mM rCTP, 0.4 mM rATP, 0.16 mM rGTP, 0.03 mM DTT, 0.3 mM cap analogue (5'7meGppp5'G), 10 μ Ci of [³²P]UTP, and 40 U of T3 RNA polymerase as appropriate. Incubation was carried out at 37°C for 30 min. DNA template was then digested at 37°C for 10 min with 40 U of RNase-free DNase. The mixture was then extracted with phenol-chloroform-isoamyl alcohol, precipitated with ethanol, and the pellet was resuspended in sterile water. The amount of cRNA synthesized was determined by the measure of its optical density at 260 nm and by liquid scintillation counting of the TCA-precipitated RNAs.

Quantification of NHE-3 mRNA. NHE-3 mRNA abundance was quantified by a competitive RT-PCR method, using the internal standard of cRNA which differed from the wild-type NHE-3 mRNA by an 80-bp deletion. Both cRNA and wild-type NHE-3 mRNA were reverse transcribed and amplified using primers 5'-GGAACAGAG-GCGGAGGAGCAT-3' (sense, bp 1,885–2,085) and 5'-GAAGT- TGTGTGCCAGATTCT-3' (antisense, bp 2,186–2,206). These NHE-3 primers yielded a product of 321 bases (18) and 241 bases for wild-type and cRNA, respectively.

For each determination, seven to eight competitive RT-PCR reaction points were performed and resolved on 1.5% agarose gels stained with ethidium bromide. As illustrated in Fig. 1 A, addition of decreasing amounts, from 168 to 0.84 amol, of the competitive template to replicate reactions containing identical amounts of MTAL total RNA resulted in a progressive decrease of the competitive template PCR product (241 bp) and a corresponding progressive increase of the wild-type template PCR product (321 bp). Photography of the gel was numerized and quantification of the fluorescence intensity of each PCR product, which is proportional to the amount of DNA, was performed on NIH Image 1.51. NHE-3 mRNA abundance was calculated as follows (Fig. 1 B): a log-log scale plot of the ratio of fluorescence intensity of the wild-type to competitive PCR products versus the known amount of cRNA added in the replicate reaction yielded a linear plot. To correct for differences in molecular weight, fluorescence data obtained from competitive template were multiplied by 321/241. At the point where wild-type and competitive PCR products are in equivalence [i.e., log(ratio) = 1], the amount of wild-type (NHE-3 mRNA) present in the initial sample is equal to the known starting amount of competitive template. Results are expressed in attomoles of NHE-3 mRNA per nanogram of total RNA.

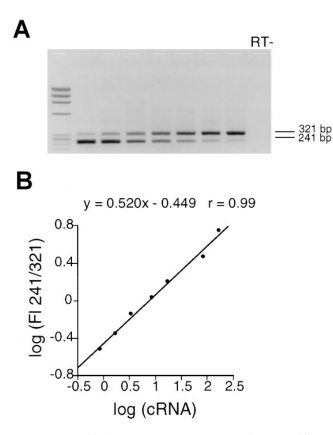


Figure 1. Quantification of MTAL NHE-3 mRNA by competitive RT-PCR. (*A*) Ethidium bromide–stained gel for competitive RT-PCR analysis of 288 ng MTAL total RNA. The amount of cRNA added in each reaction was 168, 84, 17, 8.4, 3.4, 1.68, 0.84 amol, respectively, resulting in a progressive competition between PCR products derived from wild-type NHE-3 mRNA (*upper band*, 321 bp) and from cRNA (*lower band*, 241 bp). RT–, reaction performed in the absence of reverse transcriptase (to rule out contaminating DNA). (*B*) Log–log plot of the ratio of quantitative fluorescence data versus initial amount of cRNA (attomoles).

Dot-blot analysis. Serial dilutions from 130 to 12.5 ng of MTAL total RNA were denatured in 2.2 M formaldehyde, 15× SSC (1× SSC is 0.15 M NaCl and 0.015 sodium citrate, pH 7.0) and dot-blotted onto Biodyne nylon membranes using a commercial vacuum-driven dot-blot (GIBCO-BRL, Gaithersburg, MD) apparatus and fixed by ultraviolet cross-linking. Filters were prehybridized in 4× SSC, 50 mM NaPO₄, 1× Denhardt's solution for 2 h at 42°C, hybridized in the same solution containing 25×10^6 cpm of ³²P-radiolabeled oligo-dT at 42°C overnight, and washed once in 2× SSC with 0.1% SDS at room temperature for 10 min, and twice in the same solution at 37°C for 20 min. Filters were exposed to film for 1 h at -80° C, and labeling was quantified by densitometry.

Preparation of MTAL membranes and detection of NHE-3 protein abundance by Western blot analysis. MTAL tubule suspensions were homogenized in membrane buffer (150 mM NaCl, 50 mM Tris-HCl, 5 mM EDTA containing in µg/ml: 50 AEBSF, 2 leupeptin, 2 aprotinin) and centrifuged at 10,000 g for 15 min at 4°C. The supernatant was then centrifuged at 100,000 g for 20 min (Optima TLX Ultracentrifuge; Beckman Instruments Inc., Fullerton, CA). Pellets were resuspended in membrane buffer and protein content was assessed by Bradford assay. 30 μ g of protein was diluted 1:5 in 5× SDS loading buffer (5 mM Tris HCl, pH 6.8, 1% SDS, 10% glycerol, 1% 2-mercaptoethanol), boiled for 10 min, size fractionated by SDS-PAGE on 7.5% gels, and electrophoretically transferred to nitrocellulose. After transfer, equal loading of the lanes was controlled first by Ponceau red staining of the lower third of the membrane (which was cut off), and second by Coomassie blue staining of the remaining membrane. In all cases, protein loading of the lanes was equal between control and experimental groups. After blocking with 5% nonfat milk and 0.05% Tween 20 in PBS for 1 h, blots were probed in the same buffer for 1 h with a polyclonal anti-rat NHE-3 antibody directed against amino acids 809-822 of the COOH-terminal cytoplasmic domain of rat NHE-3 (antiserum 1568) (9), at a dilution of 1:100. Blots were washed in 1% Tween 20 in PBS for 15 min once, and for 5 min twice, incubated with a 1:10,000 dilution of peroxidase-labeled sheep antirabbit IgG in 5% nonfat milk and 0.025% Tween 20 in PBS for 1 h, washed as above, and then visualized by enhanced chemiluminescence (Amersham Corp., Arlington Heights, IL). NHE-3 protein abundance was quantified by densitometry.

Statistics. Results were expressed as mean \pm SEM. Differences between means were evaluated using the paired or unpaired *t* test as appropriate. *P* < 0.05 was considered statistically significant.

Materials. Collagenase CH grade was obtained from Boehringer Mannheim France (Meylan, France), BCECF-AM was from Molecular Probes (Eugene, OR). NMG-Cl, amiloride, bumetanide, DMSO, and all chemicals were obtained from Sigma-Chimie (La Verpillère, France). Guanidium thiocyanate, phenol, *Taq* polymerase, RNase inhibitor, MMLV reverse transcriptase, and restriction enzymes were obtained from GIBCO-BRL. [³²P]γATP was obtained from Amersham. dNTP and yeast tRNA were obtained from Stratagene.

Results

Systemic measurements made at the end of the treatment period are shown in Table I. Animals tolerated each of the treatment regimens well, and all animals gained weight throughout the treatment period. Final body weight was significantly lower in the acidosis group. As expected, plasma pH and bicarbonate concentration was significantly reduced in the NH₄Cl-treated rats.

Quantification of NHE-3 mRNA in MTAL cells. In the inner stripe of the outer medulla, NHE-3 is exclusively expressed in the TAL (9, 29), which allowed us to quantify NHE-3 mRNA by a very sensitive quantitative RT-PCR method from small amounts of MTAL suspension.

Table I. Body Weight, Kidney Weight, and Plasma Electrolytes

	Initial body wt	Final body wt	Kidney wt	pH	pCO_2	HCO_3^-	Na^+	\mathbf{K}^+
	grams	grams	grams	pH units	mmHg	тM	mМ	mM
Control	279±5	336±5	1.23 ± 0.04	7.42 ± 0.01	37.3±0.7	23.9±0.4	142.6 ± 0.34	4.4 ± 0.1
	(20)	(20)	(7)	(20)	(20)	(20)	(11)	(11)
Acidosis	278±5	304±5*	1.23 ± 0.02	$7.31 \pm 0.01*$	33.8±0.5*	$17.1 \pm 0.3*$	$144.3 \pm 0.4*$	$3.9 \pm 0.1 *$
	(20)	(20)	(7)	(20)	(20)	(20)	(11)	(11)

Values are means \pm SEM for the number of rats given in parentheses. *P < 0.05 acidosis vs. control (unpaired t test).

To test the efficiency and the reproducibility of NHE-3 mRNA quantification, we performed different competitive RT-PCR reactions as described in Methods with increasing amounts of MTAL total RNA extracted from three different preparations of MTAL tubules. As shown in Fig. 2, an excellent linearity was obtained when NHE-3 mRNA was plotted against the amount of initial MTAL total RNA (from 100 to 600 ng). These results show that the competitive RT-PCR reaction, performed from a fixed amount of MTAL total RNA enclosed in the linear range obtained above, can therefore estimate the MTAL NHE-3 mRNA abundance in optimal conditions.

Effect of CMA on NHE-3 mRNA. For each experiment, experimental and control rats were studied in parallel. The effect of CMA on the MTAL NHE-3 mRNA abundance is shown in Fig. 3. Results showed a 69% increase in the NHE-3 mRNA abundance in acidotic versus control groups (0.0418±0.0091 vs. 0.0248 ± 0.003 amol/ng total RNA, n = 7, P < 0.005). To appreciate the specificity of the acidosis-induced increase in NHE-3 mRNA, we quantified the amount of mRNAs present in MTAL total RNA samples by dot-blot analysis using radiolabeled oligo-dT as described in Methods. Serial dilutions of the same initial sample resulted in an equivalent decrease of labeling (Fig. 4). Results showed that, for a fixed amount of total RNA, the amount of mRNAs was not significantly affected by CMA (242±43 vs. 288±40 integrated OD units in acidosis, n = 7, NS). Thus, the increase in NHE-3 mRNA abundance observed during CMA cannot be explained by a global increase of mRNAs. The next step of this study was to determine the effects of CMA on NHE-3 transport activity and protein abundance.

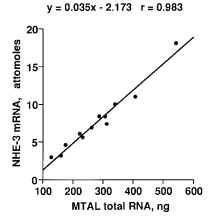
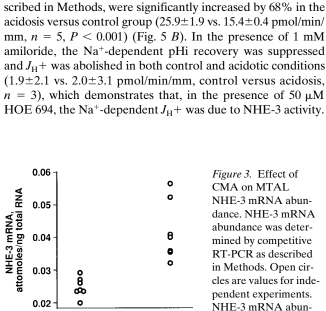


Figure 2. Separate competitive RT-PCR were performed as described in Methods, using varying amounts of MTAL total RNA ranging from 100 to 550 ng. Plot of NHE-3 mRNA abundance versus MTAL total RNA illustrates an excellent linearity over the MTAL total RNA range tested.



 0.0042 ± 0.003 amol/ng total RNA (n = 7; P < 0.005, unpaired t test), in control and acidosis, respectively.

Acidosis

Control

cell volume, and Na-dependent pHi recovery (dpHi/dt) were performed on the same MTAL tubule suspension of control and experimental rats. Bi was not different in the acidosis versus control rats (41.3±2.9 vs. 46.4±5.7 mmol H⁺/liter/pH unit, n = 5, NS) whereas the cell volume was significantly increased in the acidosis versus control group $(0.65\pm0.07 \text{ vs}, 0.43\pm0.02$ nl/mm tubule length, n = 5, P < 0.05). To specifically study NHE-3 transport activity, NHE-1 activity was inhibited by 50 μM HOE 694, a compound that blocks NHE-1 \sim 4,000 times more potently than NHE-3 (30); we have shown previously (31) that global NHE activity in the MTAL reached a maximum plateau inhibition of $\sim 35\%$ with 4–100 μ M HOE 694; 0.1 mM bumetanide and 1 µM amiloride (that had per se no significant effect on NHE-3 activity) were also present to block Na-K-2Cl cotransport and possible Na conductance, respectively, that might be altered by acidosis (23). CMA induced a significant increase in the dpHi/dt (0.985 ± 0.072 vs. $0.773\pm$ 0.039 pH unit/min, n = 5, P < 0.05) (Fig. 5 A), whereas initial acidification pHi values were not different between the two groups (6.86 \pm 0.04 vs. 6.89 \pm 0.03, acidosis vs. control, n = 5, NS). Na⁺-dependent proton flux rates $(J_{\rm H}+)$, calculated as described in Methods, were significantly increased by 68% in the

Effect of CMA on NHE-3 transport activity. For each ex-

periment, measurements of intracellular buffering power (βi),

dances were

0.0025±0.001 vs.

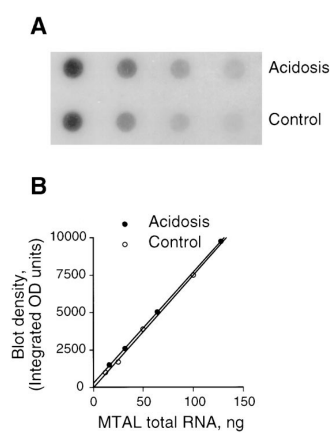


Figure 4. Dot-blot analysis of mRNAs during CMA. Serial twofold dilutions of MTAL total RNA were subjected to dot-blot analysis, hybridized with ³²P-labeled oligo-dT as described in Methods. After a 2-h exposure, autoradiograms were scanned, and band densities were quantified. (*A*) Representative autoradiograms of acidosis (*top*) and control groups (*bottom*). (*B*) Blot densities are plotted against the amount of total RNA loaded per lane, and data are fitted by linear analysis.

Effect of CMA on NHE-3 protein abundance. NHE-3 protein abundance was determined by Western blot analysis using a polyclonal anti-rat NHE-3 antibody. The specificity of this anti-NHE-3 antibody was previously demonstrated (9). As shown in Fig. 6, this antibody recognized an \sim 87-kD protein

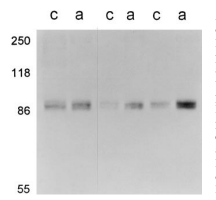


Figure 6. Immunoblot of membranes (30 μ g/ lane) from MTAL suspensions of control (c) and acidotic (a) rats obtained from three independent experiments. Membranes were probed with a polyclonal anti–rat NHE-3 antibody (1:100). Molecular weight standards are shown on the left.

in membranes isolated from MTAL cells which corresponded to the NHE-3 protein, as previously described in the rat outer medulla (9). Furthermore, NHE-3 protein amount was enhanced during acidosis (Fig. 6). Results of eight experiments showed a mean $71\pm32\%$ increase in NHE-3 protein abundance (P < 0.05) in rats fed with NH₄Cl compared with control animals.

Discussion

Bicarbonate absorption by the rat MTAL displays chronic and acute regulation, suggesting that this segment has an important role in the physiological and pathophysiological control of acid-base balance (32-34). CMA is associated with an adaptive increase in the ability of the rat MTAL to absorb bicarbonate (14), and apical Na⁺/H⁺ exchange has been shown to be responsible for bicarbonate absorption in MTALs from both control and acidotic rats (15). We have shown previously that, in the rat MTAL, NHE-1 and NHE-3 are expressed, but not NHE-2 and NHE-4 (18). Since NHE-3 is the apical Na⁺/H⁺ exchanger of the rat MTAL (9, 20), the present experiments were designed to determine if CMA is associated with intrinsic adaptation of NHE-3, and which mechanisms are involved. Our results show that NHE-3 mRNA, protein abundance, and transport activity are increased in MTAL cells freshly isolated from chronically acidotic rats.

In the inner stripe of the outer medulla, NHE-3 is almost exclusively expressed in the TAL (9, 29). We then used a very sensitive competitive RT-PCR method which allowed us to de-

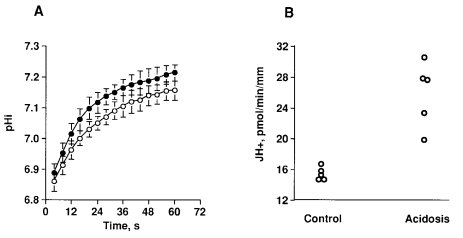


Figure 5. Effect of CMA on NHE-3 transport activity. (*A*) Effect of CMA on sodium-dependent initial rates of pHi recovery (dpHi/dt). Na-depleted acidified MTAL cells were abruptly added into isosmotic CO₂-free medium containing 30 mM Na. Points are means±SEM of five separate experiments in control (*open circles*) and acidosis (*filled circles*). (*B*) Effect of CMA on proton flux rates (J_H⁺). J_H⁺ was 15.4±0.4 vs. 25.9±1.9 pmol/min/mm (n = 5; P < 0.001, unpaired t test), in control and acidosis, respectively. tect and quantify absolute amounts of NHE-3 mRNA from small amounts of total RNA extracted from MTAL cells. Our results clearly show a 69% increase in NHE-3 mRNA abundance during CMA. This increase is specific for NHE-3 mRNA since the abundance of mRNAs relative to total RNA, assayed by dot-blot analysis in the same suspensions of MTAL cells, is not modified by CMA.

The next step of this study was to determine if NHE-3 transport activity and NHE-3 protein abundance were also increased in MTAL cells during CMA. Since MTAL cells expressed only NHE-1, the amiloride-sensitive Na⁺/H⁺ exchanger, and NHE-3 which is amiloride resistant, the use of HOE 694, a compound that completely blocks NHE-1 without altering NHE-3 at the concentration of 50 µM (30, 31), allowed us to study specifically NHE-3 transport activity. Our results demonstrate that CMA induced a mean 68% increase in NHE-3-mediated $J_{\rm H}^{+}$, which is of similar magnitude to that observed for HCO₃ transport capacity in isolated, perfused MTALs of chronically acid-loaded rats (14). In the present study, NHE-3 protein abundance was assayed on MTAL membranes by immunoblot analysis using specific anti-NHE-3 antibodies which have been shown to specifically stain the luminal membrane of MTAL (9). Results showed that CMA also increased NHE-3 protein abundance by 71% in a membrane fraction obtained from MTAL tubules. This latter result is in agreement with recent results obtained by immunocytochemistry that show increased NHE-3 abundance in the TAL apical membranes of chronically acid-loaded rats (12).

Thus, this study demonstrates for the first time that in vivo CMA increases the level of mRNA transcript for NHE-3, which corresponds to quantitatively similar increases in NHE-3 protein abundance and transport activity in freshly isolated MTAL cells. These results thus strongly suggest that an increase in protein synthesis related to the increase in mRNA is responsible for the increase in NHE-3 transport activity we observed. In addition, these adaptations can explain the increase in HCO₃ transport capacity observed in in vitro microperfused MTAL harvested from chronically acidotic rats (14).

Recent in vitro studies performed in cell lines derived from the proximal tubule have shown that acid incubation increases NHE-3 mRNA and EIPA-resistant Na⁺/H⁺ antiporter activity in OKP cells (11), and NHE-3 protein abundance in LLC-PK₁ cells (35). These results suggest that an increased protein synthesis is, at least in part, involved in these adaptations. In recent studies, CMA has been shown to increase NHE-3 protein abundance in brush border membrane vesicles isolated from NH₄Cl-treated rats (12, 13). However, cortical NHE-3 mRNA abundance assayed by Northern blot analysis was not significantly modified by acidosis (12). One possible explanation for this absence of significant effect of acidosis on cortical NHE-3 mRNA could be the difference of sensitivity of the methods used, Northern blot versus quantitative RT-PCR. Although, in this study, increases in mRNA transcript for NHE-3 and in protein synthesis seem to be the main mechanism for antiporter activation, other additional mechanisms such as a decrease in protein degradation, exocytic insertion, or activation of an inactive subpopulation of transporters cannot be excluded.

The mediators of these adaptations of NHE-3 during CMA have not been identified in these experiments. In this study, CMA led to a 50% increase in cell volume in agreement with epithelial cell hypertrophy described previously (36). It should

be noted that in vitro acid incubation, which does not induce cell hypertrophy (37), increases NHE-3 mRNA abundance in OKP cells (11), suggesting that these adaptations can occur in the absence of cell hypertrophy. Furthermore, in vitro studies on OKP cells have shown that herbimycin A, a tyrosine kinase inhibitor, and csk overexpression, which inactivate the c-src nonreceptor protein tyrosine kinase, prevented the acid activation of the apical Na^+/H^+ exchanger (38). These data suggest that, in OKP cells, the c-src nonreceptor protein tyrosine kinase plays a key role in acid activation of NHE-3. Whether such mechanisms are involved in NHE-3 adaptation in the rat MTAL remains to be determined. At present, it is not known whether the increased NHE-3 mRNA abundance observed in our study results from increased transcription of the NHE-3 gene and/or decreased transcript degradation. Indeed, in vitro studies in LLC-PK1 cells have shown that both mechanisms could be involved in the acid-induced increase in mRNA abundance of glutaminase (decreased mRNA degradation) (39) and phosphoenolpyruvate carboxykinase (increased transcription) (40), two enzymes that contribute to renal acidification by increasing ammonia synthesis.

Acidosis-induced NHE-3 activation, by enhancing transcellular H⁺ secretion in the TAL, could contribute to the decreased luminal HCO₃⁻ concentration observed in the distal tubule (41). This process seems particularly important since the increased NH₄⁺ absorption by the MTAL during CMA, by generating luminal HCO₃⁻ (14), is susceptible to increase luminal HCO₃⁻ amounts in the loop of Henle during acidosis.

In summary, this study demonstrates that NHE-3 adapts to CMA in the rat MTAL. This adaptation involved an increase in mRNA transcript that can explain the increased NHE-3 protein abundance and activity.

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