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Thiazide diuretics are used to treat hypertension; however, compensatory processes in the kidney can limit antihypertensive responses to this class of drugs. Here, we evaluated compensatory pathways in SPAK kinase–deficient mice, which are unable to activate the thiazide-sensitive sodium chloride cotransporter NCC (encoded by Slc12a3). Global transcriptional profiling, combined with biochemical, cell biological, and physiological phenotyping, identified the gene expression signature of the response and revealed how it establishes an adaptive physiology. Salt reabsorption pathways were created by the coordinate induction of a multigene transport system, involving solute carriers (encoded by Slc26a4, Slc4a8, and Slc4a9), carbonic anhydrase isoforms, and V-type H⁺-ATPase subunits in pendrin-positive intercalated cells (PP-ICs) and ENaC subunits in principal cells (PCs). A distal nephron remodeling process and induction of jα1 NOTCH signaling, which expands the cortical connecting tubule with PCs and replaces acid-secreting α-ICs with PP-ICs, were partly responsible for the compensation. Salt reabsorption was also activated by induction of an α-ketoglutarate (α-KG) paracrine signaling system. Coordinate regulation of a multigene α-KG synthesis and transport pathway resulted in α-KG secretion into pro-urine, as the α-KG–activated GPCR (Oxgr1) increased on the PP-IC apical surface, allowing paracrine delivery of α-KG to stimulate salt transport. Identification of the integrated compensatory NaCl reabsorption mechanisms provides insight into thiazide diuretic efficacy.

Introduction
The sodium chloride cotransporter (NCC) of the SLC12A3 gene (1, 2) is the principal determinant of NaCl reabsorption in the distal convoluted tubule (DCT) (3) and the key target of thiazide diuretics (4, 5). Recent human genetic and epidemiological studies highlight the importance of NCC in electrolyte balance, blood pressure control, and treatment of hypertension. Indeed, loss-of-function mutations in NCC cause Gitelman syndrome (Online Mendelian Inheritance in Man [OMIM] entry no. 263800), a recessive disorder of urinary salt wasting, variable degrees of hypotension, and a constellation of secondary electrolyte abnormalities including hypokalemia, metabolic alkalosis, hypocalcuria, and hypomagnesemia (6). Genome sequencing efforts in the Framingham cohort revealed that rare loss-of-function mutations in NCC also contribute to blood pressure variation in the general population and reduce hypertension susceptibility (7). As exemplified by the Antihypertensive and Lipid-Lowering treatment to prevent Heart Attack Trial (ALLHAT), the human genetics of NCC have an important therapeutic corollary. This landmark trial demonstrated remarkable benefits of thiazides in the treatment of essential hypertension (8), establishing NCC as a key target of the most cost-effective and medically beneficial antihypertensive agents.

Significant individual differences in the therapeutic response to thiazides are observed, however (9). Moreover, genetic loss of NCC function in humans (6, 10) and mice (3, 11, 12) causes a relatively mild salt-wasting phenotype, especially considering the significant contribution the thiazide-sensitive NCC makes to overall renal NaCl absorption. These observations suggest that strong but variable compensatory mechanisms blunt the blood pressure response and prevent deleterious changes in fluid and electrolyte balance when NCC is compromised. The nature of this compensation is poorly understood. In NCC KO mice, activation of pendrin-mediated chloride absorption (13), ENaC (11, 14), and NHE3-mediated sodium reabsorption (14) have been shown to contribute, but an integrated molecular understanding of adaptive responses to NCC inhibition has been wanting. To develop a better appreciation of the response, we sought to identify the gene networks in the kidney that are activated to drive NaCl reabsorption when NCC phosphorylation is compromised, as occurs in the absence of the STE20/SPS1-related proline-alanine–rich protein kinase (15, 16).

A serine/threonine kinase–signaling cascade involving SPAK and the with-no-lysine kinases WNK1 and WNK4 has recently emerged as a key pathway for maintaining and activating NCC on demand (17). This signaling cascade is stimulated by intravascular volume depletion or dietary sodium restriction through angiotensin II (18–21) and/or aldosterone (22–24) and adrenergic

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factors (25, 26). Once stimulated, WNK kinases bind to and phosphorylate SPAK (27). Phosphorylated SPAK can then interact with and directly phosphoactivate NCC, making it the key terminal kinase in the cascade (27). In familial hyperkalemic hypertension (aka PHAII; OMIM 145260), mutations in WNK1 and WNK4 (28) aberrantly activate the signaling pathway, causing hyperphosphorylation of SPAK and NCC (29), hypertrophy of the DCT (30), excessive urinary sodium retention, and hypertension. Likewise, a common variant in SPAK, which has gain-of-function properties, has been identified as a hypertension-susceptibility allele (31).

A prevailing function of SPAK is to activate NCC via phosphorylation (32–34). The full-length, active form of SPAK predominantly colocalizes with NCC within the DCT apical membrane (31). Moreover, SPAK KO mice display a phenotype that is identical to that of NCC-deficient mice and humans with Gitelman syndrome (32–34). While SPAK can potentially phosphoregulate the loop diuretic-sensitive sodium-potassium-chloride cotransporter NKCC2 in the thick ascending limb (TAL), NKCC2 becomes hyperphosphorylated in SPAK KO mice (32–34). Activation of NKCC2 has been suggested to play a major role in the maintenance of salt balance in SPAK KO mice, but the glomerular-tubular feedback system, which would reduce glomerular filtration rate (GFR), may blunt these effects. Compensatory salt reabsorption is more likely to have major effects at segments distal to the macula densa. Because the salt-losing phenotypes of Gitelman syndrome, SPAK KO mice, and NCC KO mice are similar and relatively mild compared with the predicted effects of loss of NCC function, we hypothesized that multiple common compensatory mechanisms blunt natriuresis and chloriuresis and protect against a major drop in blood pressure.

The present study was designed to identify and characterize the gene networks and physiological pathways that underpin the compensatory response. Our results identify renal salt transport and signaling systems that are induced to maintain salt balance and blood pressure in the absence of NCC phosphorylation.

**Results**

To identify genes that are regulated to maintain salt balance when phosphorylation of NCC is compromised, genome-wide analysis of

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**Figure 1. SPAK KO transcript profile.** (A) Heatmap rendering of the transcript expression profile of SPAK KO and WT mice (n = 4 mice per group). Differentially expressed genes in kidney cortex are shown. Each individual block represents the relative transcript level of an individual gene. Genes are arranged by row according to the FDR (from high [top] to low [bottom], with a lower-limit FDR <0.05). Each column represents the profile of an individual mouse. (B) GO classification of genes by biological function revealed that the SPAK KO profile was significantly enriched in 5 major functional classes. Level of significance (–log of the P value) is plotted for each significant GO term. Individual GO terms are arranged according to major functional class. (C) Number of SPAK–upregulated (red) and -downregulated (green) profile genes that were exclusively expressed in individual nephron segments. Glom, glomerulus; cTAL, cortical TAL.
transcript levels was performed on kidney cortex samples from WT and SPAK KO mice using the Affymetrix whole-genome GeneChip Mouse Gene 1.0 ST Array, which represents all mouse genes and contains more than 28,000 total transcripts. A distinct profile of 279 upregulated and 176 downregulated transcripts was identified in the SPAK KO mice (n = 4 WT versus n = 4 KO) using standard statistical analyses and expression levels as filtering thresholds (FDR <0.05 and relative transcript changes ± 120%) (Figure 1A). All identified genes that were selected for further analysis (n = 53 genes, see below) were verified as regulated transcripts by quantitative PCR (qPCR), providing strong support for the analytical approach. Gene ontological (GO) analysis (Figure 1B) using the Database for Annotation, Visualization and Integrated Discovery (DAVID) (35) indicated that the SPAK KO array was especially enriched in molecules that function in transport, signal transduction, gene regulation, kidney development, and metabolism.

To understand how these genes might be coregulated into functional networks, nephron segment–specific transcript profiles were created by selecting profile genes that normally have a highly restricted pattern of expression along the nephron (36) and then sorting them into groups according to their sites of expression (see Methods). Ninety-one of the upregulated genes and 32 of the downregulated genes could be classified in this way. More than one-third of the upregulated genes were expressed in PP-ICs (highlighted in red), and the majority of these are involved in ion transport. qPCR survey of IC transcripts, including SLC-type transporter family members (B), V-ATPase subunits (C), and carbonic anhydrase (CA) genes (D), confirmed the results of microarray and identified several other IC genes as being differentially expressed in SPAK KO mice. Data represent the mean ± SEM. n = 6 animals per genotype. *P < 0.05 by 2-tailed t test for WT versus KO.

Figure 2. CNT/CCD transcript profile reveals induction of an IC transport pathway in SPAK KO mice. (A) Heatmap of CNT/CCD-specific genes that are upregulated in SPAK KO mice. More than one-third of the upregulated genes were expressed in PP-ICs (highlighted in red), and the majority of these are involved in ion transport. qPCR survey of IC transcripts, including SLC-type transporter family members (B), V-ATPase subunits (C), and carbonic anhydrase (CA) genes (D), confirmed the results of microarray and identified several other IC genes as being differentially expressed in SPAK KO mice. Data represent the mean ± SEM. n = 6 animals per genotype. *P < 0.05 by 2-tailed t test for WT versus KO.

Activation of the electroneutral NaCl reabsorption transporter network. A salient feature of the CNT/CCD transcript profile is the presence of transport genes that have been implicated in mediating NaCl reabsorption by β- and non-α/non-β intercalated cells (ICs) (Figure 2A). Because both β- and non-α/non-β ICs express the Cl/HCO₃ exchanger pendrin (39–41), we refer to them collectively herein as pendrin-positive ICs (PP-ICs) for simplicity. In addition to verifying the transcript abundance increase of pendrin (Slc26a4), which is activated in NCC KO mice (13, 42), the profile revealed an increase in AE4 (Slc4a9), numerous V-type proton ATPase subunits (V0 subunit d2 and the V1 subunits B1, C2, and G3), and 2 different isoforms of carbonic anhydrase (CA2 and CA15). A comprehensive qPCR survey of IC transporters (Fig-
ure 2B), as well as proton ATPase subunits (Figure 2C) and carbonic anhydrase genes (Figure 2D), confirmed the array findings and revealed that the sodium-dependent chloride bicarbonate cotransporter NDCBE (Slc4a8) (43) was also increased. In striking contrast, the key basolateral $\text{Cl}^-/\text{HCO}_3^-$ exchanger of the $\alpha$-IC AE1 (Slc4a1) (44, 45) was absent in the SPAK KO transcript profile, and no changes in its transcript abundance were detected in the SPAK KO kidney by qPCR (Figure 2B).

Further experiments explored the effect of SPAK gene ablation on the protein abundance of IC transporters for which antibodies are available. Immunoblot analysis (Figure 3, A and B) confirmed an increased abundance of the PP-IC proteins pendrin, NDCBE, AE4, B1-VATPase, and CA15. As a negative control, no change was detected in the abundance of the $\alpha$-IC protein AE1.

Collectively, these observations identify a highly organized regulatory response in PP-ICs that orchestrates the simultaneous activation of a multitransporter network. Recent studies provide strong evidence that each of the upregulated transport molecules functions as an integral component of an electroneutral NaCl reabsorption system (Figure 3C and see Discussion). Its activation provides a means to increase NaCl reabsorption and compensate for the loss of NCC function.

Enhanced apical expression of pendrin and chloride reabsorption. Because apical plasma membrane pendrin abundance is strongly regulated through membrane-trafficking processes (46–49), we examined the effect of SPAK gene ablation on pendrin localization within $\beta$-type PP-ICs. In contrast to the diffuse cytoplasmic and weak apical membrane pendrin labeling that was observed in PP-ICs from WT mice, intense pendrin labeling was observed in the apical regions of PP-ICs from \(\text{SPAK}^-\)KO mice (Figure 4A).

Quantitative measurements of pendrin pixel intensity across the apical membrane and cytoplasm in multiple PP-ICs (\(n = 50\) cells per animal, 5 animals per group) (Figure 4B) confirmed that the increase in pendrin abundance at the apical membrane in \(\text{SPAK}^-\)KO mice was robust and statistically significant (Figure 4C). As assessed by measurements of transepithelial chloride flux in isolated perfused CCDs, the change in pendrin was associated with an increase in tubular transport of chloride (Figure 4D). Indeed, we found that CCDs from \(\text{SPAK}^-\)KO mice had approximately 6-fold greater chloride reabsorption than did WT CCDs (13.3 ± 4.2 vs. 2.4 ± 1.7 pmol/mm/min; \(P < 0.05\), \(n = 4\)). Thus, augmentation of pendrin abundance and apical localization were paralleled by an expected increase in chloride reabsorption.

Remodeling of the distal nephron. The CNT/CCD profile is also enriched in factors that control nephrogenesis (Figure 1B), and some of these have been implicated in IC differentiation and IC subtype conversion. Accordingly, we considered the possibility that an IC remodeling process is a chief underpinning of PP-IC network activation. To explore this question, IC subtypes were identified with cell markers (PP-IC and $\alpha$-IC AE1-positive cells) and counted. In addition, ICs were colocalized with nephron segment–specific markers so that potential sites
NOTCH receptor ligand jagged 1 (Jag1) and downregulation of Jag2 were accompanied by an increase in a key NOTCH-activated transcriptional factor, hairy and enhancer of split 1 (Hes1), as well as iroquois homeobox molecules (Irx1 and Irx2) (Figure 6A). The signature is indicative of activation of NOTCH signaling in the adult SPAK KO mouse kidney cortex. Western blot analyses confirmed that HES1 and JAG1 protein abundance was increased in the SPAK KO mouse (Figure 6, B and C).

Further experiments were performed to determine where NOTCH signaling occurs. As observed by immunofluorescence confocal microscopy, nuclear localization of HES1, a reporter of NOTCH activation, was predominantly observed in nuclei of cells within the CNT (Figure 6D). Colabeling of HES1 with PC (AQP2) and IC markers revealed that HES1 was exclusively induced within PC nuclei (Figure 6E and see below). JAG1 labeling was also increased along the CNT, and immunodetectable signal was observed within both PCs and ICs, but it was not possible with IHC to distinguish between cells that expressed the full-length receptor on the cell surface and those that had JAG bound to NOTCH. Nevertheless, these observations suggest that JAG1 activates NOTCH signaling in PCs.

Together, these observations suggest that activation of the NOTCH signaling pathway in CNT PCs coordinates a program of PC differentiation and proliferation, allowing the CNT to replace the dystrophic DCT1. The expansion of the CNT and the increase of remodeling could be identified. As shown in Figure 5A, SPAK KO mice displayed an obvious and remarkable increase in the number of PP-ICs. This was paralleled by a reciprocal reduction in the number of α-ICs, without remarkable changes in the absolute number of ICs (1.1% increase in total IC numbers in SPAK KO mice relative to WT mice). The remodeling response was greatest in the CNT, weaker in the CCD, and not present in the late DCT (Figure 5B). Thus, activation of the PP-IC transporter network can be explained, at least in part, by a cellular remodeling process that replaces α-ICs with PP-ICs in the CNT and CCD. The response was most predominant in the CNT.

This IC remodeling process was accompanied by an increase in principal cells (PCs), especially in the CNT (21.5% increase in PC numbers) (Figure 5C). Stereological morphometric measurements of individual distal nephron segments revealed that the increase in PCs is accompanied by an increase in CNT length (Figure 5D), and this precisely offsets the decline in DCT1, which fails to develop or atrophies in SPAK KO mice (33, 34) and in other models of NCC dysfunction (11, 37).

We considered that the kidney development genes identified in the profile may contribute to the remodeling process. We focused on profile components that are involved in the NOTCH signaling cascade, because this pathway has been implicated in determining PC/IC fate (50). As identified by transcription profiling and confirmed by qPCR experiments, upregulation of the NOTCH receptor ligand jagged 1 (Jag1) and downregulation of Jag2 were accompanied by an increase in a key NOTCH-activated transcriptional factor, hairy and enhancer of split 1 (Hes1), as well as iroquois homeobox molecules (Irx1 and Irx2) (Figure 6A). The signature is indicative of activation of NOTCH signaling in the adult SPAK KO mouse kidney cortex. Western blot analyses confirmed that HES1 and JAG1 protein abundance was increased in the SPAK KO mouse.
in PCs augment the capacity for electrogenic NaCl reabsorption via the epithelial sodium channel ENaC.

Na⁺ absorption in PCs is determined by ENaC (25, 51), and the SPANKO transcript profile as well as qPCR identified the ENaC β and γ subunits as upregulated genes (Figure 7A). The serum glucocorticoid–induced kinase SGK1, which increases ENaC apical abundance (52, 53), and claudin 4, a determinant of a Cl⁻-selective paracellular pathway in the distal nephron (54), were also identified as increased on the array. Taken together, the transcript profile provides support for the idea that ENaC-mediated sodium reabsorption is activated in parallel with a paracellular Cl⁻ reabsorption pathway and a transcellular electroneutral NaCl transport system in PP-ICs.

To test ENaC function, we assessed the diuretic and natriuretic responses to benzamil, which is considered to be a specific ENaC inhibitor at the low dose used here (1.0 mg/kg). SPANKO mice exhibited significantly increased responses to benzamil compared with that in WT mice, verifying that a functional increase in ENaC-mediated sodium reabsorption accompanies PC expansion (Figure 7, B and C). Together with the increase in PP-ICs and activation of electroneutral salt reabsorption, the cellular remodeling response provides two powerful mechanisms to offset the loss of NCC activity.

**Paracrine activation of NaCl reabsorption in PP-ICs by α-ketoglutarate.** One mechanism to activate pendrin is revealed by the identification of Oxgr1 as an upregulated gene in the CNT/CCD transcript profile. Oxgr1 encodes a novel GPCR for α-ketoglutarate (α-KG) (55), and recent studies demonstrated that this receptor is responsible for activating pendrin-dependent chloride reabsorption (56). As shown in Figure 8A, qPCR confirmed Oxgr1 as an upregulated transcript in SPANKO mice. As assessed by immunoblot analysis with specific OXGR1 antibodies (Supplemental Figure 1; supplemental material available online with this article; doi:10.1172/JCI78558DS1), the increase in Oxgr1 transcripts was paralleled by an increase in protein abundance. OXGR1 protein migrates at the predicted size (38 kDa) and in higher molecular weight forms (~75 and 100 kDa), consistent with glycosyla-
tion and/or receptor oligomerization (Figure 8B); all forms were increased in SPARKO mice relative to WT mice (Figure 8C). Immunofluorescence confocal microscopy revealed that OXGR1 was exclusively expressed in PP-ICs at the apical membrane and was more abundant in SPARKO mice than in WT mice (Figure 8, D and E). Thus, OXGR1 is poised to hyperactivate pendrin, so long as there is sufficient α-KG in the pro-urine.

The PT transcript profile revealed a mechanism to enhance the release of α-KG into the tubular fluid for paracrine delivery to OXGR1 (Figures 9 and 10). In addition to its key role as an intermediate of the Krebs (TCA) cycle, α-KG is also produced in the PT as an intermediate of ammoniagenesis, a process that is stimulated by acidosis or potassium depletion through the activation of glutamine and glutamate deamination enzymes and uptake transporters (57–59). Remarkably, the SPARKO PT profile (Figure 9A) contained many of the key transporters (NaDC1, OAT10, SAT2, SNAT3) and enzymes (glutamate oxaloacetate transaminase) that facilitate ammoniagenesis and could potentially enhance α-KG production. The observation prompted a comprehensive qPCR survey of the ammoniagenesis pathway and measurements of urinary ammonium and α-KG excretion.

Two of the four known sodium-dependent glutamine uptake proteins (60), the apically localized SAT2 (Slc38a2) and the basolateral SNAT3 (Slc38a3) (61), were identified as upregulated genes in the array and verified by qPCR (Figure 9B). Additional qPCR screening identified increased levels of the apically expressed sodium-dependent glutamine transporter B OAT1 (Slc6a19) (62) and the basolateral glutamate transporter AGT1 (Slc7a13) (63), but not ASCT2 (Slc1a5), in the SPARKO kidney. Genes encoding the dicarboxylate transporter (64) Slc13a2 (NaDC1) and organic anion exchangers (Slc22a13, OAT10; ref. 65) were also identified as upregulated on the array (Figure 9A) and verified using qPCR (Figure 9C), along with Slc13a3 (NaDC3) and the sodium-hydrogen exchanger NHE3 (Slc9a3). By contrast, the α-KG transporter OAT1 (Slc22a6), which mediates α-KG efflux across the basolateral membrane (66, 67) and would limit α-KG secretion, was found on the downregulated transcript profile and verified by qPCR. The transport properties of these transporters
predict that their regulation would increase α-KG and ammonia secretion (see Figure 9D).

Glutaminase (GLS) and glutamate dehydrogenase (GLUD1), the key regulated enzymes that deaminate glutamine and glutamate to produce ammonia and α-KG (Figure 10A), were also detected as upregulated transcripts by qPCR (Figure 10B). By contrast, enzymes that drive the reverse reactions to produce glutamine from α-KG were downregulated. Phosphoenolpyruvate carboxykinase (PEPCK) of the PCK1 gene), which is activated for ammoniagenesis in states of acidosis to convert excess α-KG into glucose and generate bicarbonate (68, 69), was not on the SPAK transcript profile, and no change in transcript abundance could be detected by qPCR. These data indicate that the PT is poised to enhance the production of α-KG.

Plasma and urine α-KG were evaluated to test whether such a mechanism results in increased α-KG in the tubular fluid. As shown in Figure 10, C and D, urine α-KG levels normalized by urinary creatinine increased nearly 3-fold in SPAK KO mice, with no detectable change in the plasma concentration, consistent with increased secretion of α-KG into the tubular fluid. The response was matched by an increase in urinary ammonia excretion (Figure 10E), consistent with the predicted parallel activation of ammoniagenesis and α-KG secretion. A slight alkalization of the urine (Figure 10F) was also observed, consistent with parallel activation of pendrin-mediated bicarbonate secretion.

Taken together, the data identify a paracrine signaling system that connects PT ammoniagenesis and α-KG production and secretion to the activation of pendrin-mediated salt transport in the distal nephron.

Discussion

Our data unveil a multifaceted and integrated compensatory mechanism that preserves salt balance when the activity of the thiazide-sensitive NaCl cotransporter NCC is compromised, as occurs in the absence of the SPAK kinase. We found that a distal nephron remodeling process and the activation of a paracrine signaling system act together to induce salt-reabsorbing transport pathways in the CNT and CCD.

Induction of the PP-IC transport network. The compensatory response highlights PP-ICs, which have recently emerged as an important avenue of regulated salt transport in the kidney (70, 71). Extending the recent discovery that increased expression of pendrin is critical for maintaining salt reabsorption in NCC KO mice (13), our observations reveal that the induction of pendrin is part of a highly organized regulatory response involving the simultaneous activation of a multitransporter network. On the basis of physiological studies in different transporter KO mice, Eladari and coworkers proposed that each of the upregulated transport molecules, identified here, acts as an integral component of an electro-neutral NaCl reabsorption machine in PP-ICs (Figure 3C and ref. 71). According to their model, the concerted activity of pendrin-mediated Cl/HCO3 exchange and NDCBE-mediated Na/HCO3/C exchange at the apical membrane allows the uptake of sodium and chloride from the pro-urine (43, 72), while the activity of the H+-ATPase, the AE4 Na/HCO3 cotransporter (71), and an undefined Cl channel facilitate NaCl efflux across the basolateral membrane into the interstitium. In this scheme, the H+-ATPase powers the system, creating a favorable outwardly directed HCO3 gradient for the bicarbonate-dependent transporters pendrin and AE4 to drive electroneutral salt reabsorption (73). It is not presently clear what driving forces are available for NDCBE to mediate sodium uptake (70), but it seems likely that enhanced sodium delivery in the absence of NCC provides a sufficient chemical gradient to drive inward sodium movement across the apical membrane and increase sodium reabsorption in parallel with pendrin-dependent chloride uptake. Our observation that the entire

![Figure 7. ENaC expression and function are enhanced in SPAK KO mice.](image-url)
transport system behaves as a coregulated unit offers further support for the NaCl-reabsorbing transport model.

Activation of the PP-IC transport system not only provides a means to limit urinary salt loss, but it can also compensate for the hypochloremic metabolic alkalosis that can develop with loss of NCC function. Although the precise stoichiometry of transporters in the PP-IC transport system is unknown, current evidence suggests that it allows PP-ICs to secrete bicarbonate in exchange for the reabsorption of NaCl. Indeed, pendrin/NCC double-KO mice develop much more severe metabolic alkalosis and salt wasting than do mice lacking NCC or pendrin alone (13). Likewise, thiazide diuretic treatment was reported to cause severe hypochloremic metabolic alkalosis and salt wasting in a child with pendrin syndrome (74).

**Distal tubule remodeling.** Activation of the PP-IC transporter pathway is explained in part by a remarkable distal nephron cellular remodeling process involving an increase in the number of PP-ICs. Because the response is matched by a decrease in α-ICs, it is tempting to speculate that IC remodeling is driven by the differentiation of α-ICs into PP-ICs, but further studies will be required to explore this idea and test other possible mechanisms. Convincing in vitro and in vivo data indicate that acidosis provokes the differentiation of α-ICs from β-ICs (75) in a process involving the induction of the cell adhesion factor hensin (76) and β1 integrin (77). Nevertheless, neither hensin nor β1 integrin was identified as a downregulated gene in the SPAK KO mouse profile, and it is not clear whether the IC conversion works in the opposite direction. In fact, PP-ICs have been reported to be progenitors of α-ICs and PCs in vitro (78), and, as far as we know, the differentiation of α-ICs into PP-ICs is unprecedented. It is also conceivable that the remodeling is shaped by differences in IC proliferation rates, as has been observed in response to loop diuretics and high dietary salt intake (79), although this does not profoundly alter the ratio of the different IC types in the manner observed here in our SPAK KO mice. During the early postnatal period, PP-ICs appear to arise from undifferentiated precursor cells (80), raising another possibility that the neonatal differentiation process is recapitulated and exaggerated in the adult kidney when NCC-mediated transport is inhibited and the DCT atrophies. The profile of nephrogenic factors identified here should provide a roadmap for future studies to elucidate the basis of the IC remodeling mechanism.

In fact, our identification and characterization of NOTCH signaling elements revealed that embryonic nephrogenic cell fate determinants can be reactivated in the adult kidney to shape the distal nephron remodeling process. In the developing collecting
The activities of ENaC and pendrin are often coupled together. For example, genetic ablation of pendrin causes a reduction in ENaC (83), and activation of pendrin stimulates ENaC by increasing luminal bicarbonate and alkalizing the tubular fluid (84). Other positive coupling mechanisms involving pendrin-dependent ATP release from PP-ICs and purinergic activation of ENaC have been reported (85, 86). By increasing the number of PP-ICs and PCs together, the distal nephron remodeling process is poised to amplify pendrin-ENaC coupling so as to activate the electroneutral and electrogenic salt transport pathways in parallel and optimally enhance NaCl reabsorption in distal nephrons.

α-KG/OXGR1 paracrine signaling. Our data reveal that salt reabsorption through the PP-IC pathway is also activated by induction of a newly discovered intrarenal α-KG paracrine signaling system. Remarkably, increased expression of OXGR1, the α-KG GPCR on the PP-IC apical membrane, is accompanied by an increase in α-KG within the tubular fluid (84). Other positive coupling mechanisms involving pendrin-dependent ATP release from PP-ICs and purinergic activation of ENaC have been reported (85, 86). By increasing the number of PP-ICs and PCs together, the distal nephron remodeling process is poised to amplify pendrin-ENaC coupling so as to activate the electroneutral and electrogentic salt transport pathways in parallel and optimally enhance NaCl reabsorption in distal nephrons.

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duct, NOTCH signaling is required to drive the differentiation of PCs (50). Our observations that NOTCH signaling was selectively activated in PCs of adult SPAK KO mice, concurrent with their proliferation, strongly suggest that this developmental process is recapitulated in the adult kidney to increase the number of PCs and expand the CNT. It will be interesting to learn the lineage of these NOTCH-activated PCs. Upon recovery from lithium-induced nephrogenic diabetes insipidus (81) or potassium deficiency (82), PCs have been reported to arise from the conversion of α-ICs, raising the possibility that NOTCH signaling in the SPAK KO mouse drives a similar process to expand PCs at the expense of α-ICs. Given that the morphological changes in the distal nephron of the SPAK KO mouse are remarkably similar to those in the NCC KO mouse (11), in which the CNT is also expanded to replace an atrophic distal tubule, it seems likely that NOTCH signaling provides a common mechanism to drive PC differentiation and proliferation when sodium reabsorption in the early distal tubule is compromised.

The collective distal nephron remodeling process ultimately allows pendrin- and ENaC-mediated transport processes to be efficiently coordinated. In recent years, it has become appreciate...
The upregulation of key enzymes necessary for ammoniagenesis and \( \alpha \)-KG production activate \( \alpha \)-KG secretion. (A) Reversible multienzyme ammoniagenesis pathway produces \( \alpha \)-KG from glutamine. (B) Summary of qPCR survey of key enzymes responsible for \( \alpha \)-KG anabolism (Gls and Glutd1) and catabolism (Glut1 and Got2). Data represent the mean ± SEM. \( n = 6 \) animals. *\( P < 0.05 \) by 2-tailed \( t \) test for WT versus KO. Summary data for plasma \( \alpha \)-KG (C), as well as for urinary excretion of \( \alpha \)-KG (D) and ammonia (\( \text{NH}_3 \) and \( \text{NH}_4^+ \)) (E) and urine pH (F). Data represent the mean ± SEM. \( n = 6 \) animals. *\( P < 0.05 \) by 2-tailed \( t \) test for WT versus KO.

aldosterone, angiotensin II, and alkalosis (46, 47), can be elevated in Gitelman syndrome, as well as in NCC- and SPAK KO mice, and likely work with \( \alpha \)-KG to maximize pendrin-mediated transport.

We found that enhanced production of \( \alpha \)-KG results from induction of the multigene ammoniagenesis pathway. The concerted upregulation of glutamine and glutamate uptake transporters (SAT2 [Slc38a2], SNAT3 [Slc38a3], B0AT1 [Slc6a19], and AGT1 [Slc7a13]), the upregulation of glutamine and glutamate deaminating enzymes (Gls and Glutd1), and the downregulation of \( \alpha \)-KG catabolic enzymes (Glut1 and Got2) cause the production of ammonia to increase in the SPAK KO mouse, similar to the response that is provoked by acidosis and potassium deficiency. But unlike acidosis, which also activates phosphoenolpyruvate carboxykinase (PEPCK) to consume \( \alpha \)-KG and generate glucose and bicarbonate (69, 87), PEPCK is not induced in SPAK KO mice, and, as a result, \( \alpha \)-KG levels increase rather than diminish.

Differential regulation of PT \( \alpha \)-KG transporters explains how enhanced \( \alpha \)-KG production is accompanied by an increase in \( \alpha \)-KG delivery to the tubular fluid. Net \( \alpha \)-KG uptake from the blood across the basolateral membrane is predicted to increase from the upregulation of NaDC3 (64, 88), which uses the inwardly directed electrochemical sodium gradient to transport \( \alpha \)-KG into the cell across the basolateral membrane, and from the downregulation of OAT1, which mediates basolateral \( \alpha \)-KG efflux (66, 89). The transporter(s) responsible for carrying \( \alpha \)-KG across the apical membrane into the tubular fluid have not been established, but our identification of Oat10 as an upregulated gene provides a clue. Because OAT10 is expressed on the PT apical membrane and has the capacity to exchange organic anions for the efflux of short-chain fatty acids (65), we propose that its activation leads to an increased efflux of \( \alpha \)-KG across the apical membrane.

Although further studies will be required to identify the factors responsible for activating the paracrine signaling process, there is reason to suspect that hypokalemia and metabolic alkalosis are involved. Both can develop as a consequence of NCC inhibition, and each have the capacity to either stimulate \( \alpha \)-KG production or enhance \( \alpha \)-KG secretion. Dietary alkaline loading, by itself, has recently been shown to enhance urinary \( \alpha \)-KG excretion (56). Because alkalosis usually suppresses ammoniagenesis, it is more likely to activate the secretory transport of \( \alpha \)-KG than to stimulate production. Differential regulation of NaDC3 and the OAT transporters observed here may offer an explanation. By contrast, potassium deficiency induces ammoniagenesis (59, 90) and therefore probably activates the production of \( \alpha \)-KG. By stimulating \( \alpha \)-KG through separate pathways, the combined effects of potassium deficiency and alkalosis are expected to have potentiating effects on the stimulation of \( \alpha \)-KG as a paracrine factor.

Such a system would be ideally suited to limit hypokalemic metabolic alkalosis. Ammonium excretion has been proposed to be stimulated in states of potassium deficiency as an adaptive mechanism to reduce urinary potassium loss (59, 91). If accompanied by an increase in \( \alpha \)-KG signaling, activation of bicarbonate
secretion by pendrin will limit the renal acid excretion that otherwise accompanies the increase in ammonium secretion and help compensate for the metabolic alkalosis.

Hormones that stimulate ammoniagenesis or enhance the expression of α-KG transporters should also be considered as candidate activators of the paracrine pathway. Observations that angiotensin II stimulates ammoniagenesis (92), for example, raise the intriguing possibility that it also activates α-KG signaling. Parallel activation of ammonium (acid) secretion and pendrin-mediated Cl/HCO₃ exchange would have the net effect of stimulating chlo-

Summary. In summary, gene profiling, combined with biochemical, cell biological, and physiological phenotyping, defined adaptive mechanisms that maintain salt balance in the absence of SPAK and NCC phosphorylation. It will be of interest to understand whether overactivation of these adaptive pathways contributes to the development of diuretic resistance.

Methods

Animals. The generation and phenotypic characterization of SPAK KO (SPAK−/−) mice have been described in detail previously (33, 34, 93). Heterozygous pairs, backcrossed more than 20 generations, were bred to generate WT and KO mice for experimentation. Male pups (~4 weeks of age) were screened by PCR genotyping of tail DNA to identify the KO allele. Because a PCR-based strategy cannot distinguish heterozygous and KO animals, the final determination of genotype was assessed using SPAK antibodies in Western blot analyses of kidney lysates. For these studies, mice were fed a control diet (1% K⁺, 0.32% Na⁺, 0.9% Cl⁻) (TD.88238; Teklad Lab Animal Diets, Harlan Laboratories) after weaning. Paired 8-week-old WT and KO littersmates were compared. Mice were maintained on a 12-hour light/12-hour dark cycle, with lights on at 6 a.m. Food and water were available ad libitum. SPAK KO animals used for these studies had the same phenotypic properties as those previously reported (see Supplemental Table 1).

Sample collection, preparation, and analysis. Animals were anesthetized by i.p. injection with ketamine/xylazine (100 mg/kg ketamine, 10 mg/kg xylazine), and the left kidney was removed, dissected into cortex and medulla samples, and stored in RNA later (QIAGEN) or flash-frozen in liquid nitrogen for later RNA and protein extraction. Blood samples were collected from the carotid artery and immediately spun down for measurements of plasma α-KG, creatinine, and electrolytes. Urine samples were collected from the carotid artery and immediately spun down for measurements of plasma α-KG, creatinine, and electrolytes. Urine samples were collected from the anesthetized mice by puncturing the bladder with a 24-gauge needle and drawing out urine with a syringe. Plasma and urine creatinine levels were measured using the Ammonia Reagent Set (Pointe Scientific Inc.).

RNA isolation, microarray, qPCR. RNA was extracted and purified from the cortex of 4 SPAK KO and 4 WT littermate mice using TRIzol reagent (Life Technologies). A microarray assay was performed using the Affymetrix GeneChip (Mouse Gene 1.0 ST), containing probes for more than 28,000 transcripts, and analyzed on an Affymetrix GeneChip 3000 system (Biopolymer-Genomics Core Facility at the University of Maryland School of Medicine). Quality control analysis, robust multiarray average (RMA) normalization, and linear model for microarray-based (LIMMA-based) differential expression analysis were subsequently performed. Array data were deposited in the NCBI’s Gene Expression Omnibus (GEO GSE60728) (94). qPCR was used to verify the altered expression of array-identified candidate genes and to assess differential gene expression of proteins involved in the compensatory response, using a Roche LightCycler 480 qPCR system and LightCycler 480 SYBER Green I Master reagents pathways. All reactions were performed in triplicate for each animal (n = 6 WT and n = 6 KO). Relative transcript abundance was calculated using the Pfaff equation (95), a derivation of the ∆∆Ct method. Unlike the latter, the Pfaff equation accounts for actual efficiency of doubling within the linear range of amplification, a value that can vary depending on template concentration. The values represented in the figures are normalized Pfaff values relative to WT.

Enrichment analysis of array genes. The DAVID database was used to sort array-identified genes by biological function and then statistically rank the most overrepresented (enriched) functional categories (35).

Nephron segment-specific transcript profiles. Serial analysis of gene expression (SAGE) data (36) from isolated mouse nephron segments was used as a reference to identify profile genes that are specifically expressed within individual nephron segments. Segment-specific genes were defined as genes that have at least 75% of their total transcript count within a single nephron segment. Profile genes that matched this criterion where sorted into groups, according to their sites of expression, to generate nephron segment–specific transcript profiles.

Sample preparation for Western blot analysis. Mouse cortex tissue was sonicated in HEEN buffer (20 mM Hepes [pH 7.6], 125 mM NaCl, 1 mM EDTA, 1 mM EGTA, 10% glycerol) containing 1% Triton and 0.5% SDS with protease and phosphatase inhibitors. Samples were sonicated for three 10-second pulses. The samples were rotated in 4°C for 1 hour at low speed (10 g) and then spun down at high speed (20 g) for 10 minutes to pellet-insoluble material. The supernatant was collected and quantified for protein yield using a bicinchoninic acid protein assay reagent kit (Pierce). Equal amounts of kidney protein were suspended in Laemmli buffer (at room temperature for 45 minutes) and resolved on 4% to 15% gradient SDS-PAGE gels, then transferred to Amersham Hybond ECL membranes (GE Healthcare) and blocked in Tris-buffered saline with 0.1% Tween-20 (TBS-T) containing 5% nonfat dry milk (NFDM) for 1 hour at room temperature. Membranes were then incubated in 5% NFDM containing primary antibody (4°C, overnight), washed in TBS-T, incubated in 5% NFDM containing HRP-conjugated secondary antibody, and then washed 3 times for 10 minutes each time in TBS-T. Bound antibodies were then revealed using an enhanced chemiluminescence reagent (Pierce) and fluorography. Protein quantification was done by scanning autofluorograms, followed by ImageJ analysis (NIH). Bands were measured in the linear range of the fluorographic signal. Duplicate gels were processed and developed in parallel. After detection of the target, membranes were stripped and reprobed for tubulin as a loading control. The full, uncropped gels are provided in the supplemental material.

Immunolocalization. Anesthetized mice were fixed by perfusion with 2% paraformaldehyde in PBS via the left ventricle for 5 minutes
at room temperature. The kidneys were then removed and fixed for an additional 24 hours at 4°C in the same fixative, rinsed in PBS, and embedded in paraffin. Cross-sections of 3 μm thickness, cut at the level of the papilla, were picked up on chrome-alum gelatin–coated glass coverslips and dried on a warming plate. The sections were then depa-

raffinized in 2 xylene baths and 2 absolute ethanol baths (5 minutes for each bath), and rehydrated in a graded ethanol series to distilled water.

For epitope retrieval, the coverslips were placed in a pH 8 aqueous solution of Tris (1 mM), EDTA (0.5 mM), and SDS (0.02%). The retrieval solution was heated to boiling in a microwave oven, transferred to a conventional boiling water bath for 15 minutes, and then allowed to cool to room temperature before the sections were thor-

oughly rinsed in distilled water to remove the SDS.

Sections were preincubated for 30 minutes with Image-iT blocking solution (Invitrogen), rinsed in PBS, and then preincubated for an additional 30 minutes in a solution of 2% BSA, 0.2% fish gelatin, 5% normal donkey serum, and 0.2% sodium azide in PBS. Tissues were thoroughly rinsed with TBS to remove the PBS. Incubations with specific antibodies (as described above) diluted in TBS containing 1% BSA, 0.2% fish gelatin, 0.1% Tween-20, 10 mM CaCl₂, and 0.2% sodium azide took place overnight in a humid chamber at 4°C. After thorough washing in a high-salt wash (incubation medium plus added NaCl at 0.5 M), Alexa Fluor secondary antibodies (Molecular Probes) were diluted 1:100 in incubation medium and applied to samples for 1 hour, followed by a high-salt wash, and then mounted onto slides with VECTASHIELD Mounting Media (Vector Laboratories).

Antibodies. The following antibodies were used for Western blot analysis and/or immunolocalization studies: mouse anti-calbindin D-28K (IHC 1:600; catalog c-9848; Sigma-Aldrich); chicken anti-AQP2 (from James Wade; IHC 1:100); rabbit anti-pendrin (from Susan Wall; Western blot 1:5,000; IHC 1:50,000); goat anti-pendrin (IHC 1:50; catalog sc-16894; Santa Cruz Biotechnology Inc.); goat anti-NDCBE (Western blot 1:100; catalog sc-169346; Santa Cruz Biotechnology Inc.); goat anti-B1-VATPase (Western blot 1:50; catalog sc-21206; Santa Cruz Biotechnology Inc.); goat anti-CXAV (Western blot 1:100; catalog sc-54772; Santa Cruz Biotechnology Inc.); rabbit anti-AE4 (Western blot 1:100; catalog AE41-A; Alpha Diagnostic International); rabbit anti-AE1 (Western blot 1:1,000, IHC 1:50); goat AE11-A; Alpha Diagnostic International); rabbit anti-tubulin (Western blot 1:2,000; catalog 2144s; Cell Signaling Technology); rabbit anti-HES1 (Western blot 1:1,000, IHC 1:3,000; catalog 2144s; Cell Signaling Technology); rabbit anti-OXGR1 (Western blot 1:100, IHC 1:25; catalog PA9-32960; Thermo Scientific); and goat anti-JAG1 (Western blot 1:200, IHC 1:100; catalog sc-6011; Santa Cruz Biotechnology Inc.).

Measurement of transepithelial Cl⁻ flux in perfused, isolated CCDs. Littermate SPAK KO and WT mice were fed the control diet described above and drank water ad libitum. CCDs were dissected from meso-

dular rays and perfused at flow rates of 2 to 3 nl/min in the presence of symmetric, HCO₃⁻-buffered physiological solutions containing 125 mM NaCl, 24 mM NaHCO₃, 2.5 mM K₃HPO₄, 2 mM CaCl₂, 1.2 mM MgSO₄, and 5.5 mM glucose. Tubules were equilibrated at 37°C for 15 minutes before starting collections. Cl⁻ concentration was measured in the perfusate and collected samples using a continuous-flow fluorimeter and the Cl⁻-sensitive fluorophore 6 methoxy-N-(3-sulfopropyl) quinolinium (SPQ) (Molecular Probes, Life Technologies), as described previously (96–98). Transepithelial Cl⁻ flux, \( \dot{J}_{\text{Cl}} \), was calculated according to the following equation: \( \dot{J}_{\text{Cl}} = (C_e - C_i)Q/L \), where \( C_e \) and \( C_i \) are the perfu-
sate and collected fluid Cl⁻ concentrations, respectively. \( Q \) is the flow rate in nl/min, and \( L \) is the tubule length. Net fluid transport was taken to be zero, since net fluid flux has not been observed in CCDs when perfused in vitro in the presence of symmetric solutions and in the absence of vasopressin (99, 100). \( \dot{J}_{\text{Cl}} \) is expressed in pmol/mm/min.

Quantification of nephron segment length in kidney sections. Low-power images of the kidney cortex were acquired using a Zeiss LSM 510 confocal microscope with a ×10 objective lens and appropriate filters. The DCT1, DCT2, CNT, and CCD could be identified for quantification using antibodies against the segment-specific proteins parvalbumin (parva), calbindin D28 (calbn), and aquaporin 2 (AQP2): DCT1 = parva⁺ calbn⁻AQP2⁺; DCT2 = parva⁺ calbn⁻AQP2⁻; CNT = parva calbn⁻AQP2⁻; and 3 fluorophores. Every cortical distal nephron segment in each tissue section was classified using this system.

Total DCT1, DCT2, CNT, and CCD lengths were determined with a curvilinear stereometric system developed by Merz (101) and previously published by our group (33). For these measurements, a curve line-

ter test grid composed of evenly spaced curved lines was superimposed over the images of the cortex. Tubule length, \( L \), was determined by \( L = I \times D \), where \( I \) is the number of intersections between the curvilinear test grid and the tubule, and \( D \) is the distance between corresponding points on either the same line or the adjacent line, which was 21 μm for our test grid. The length of each identifiable DCT1, DCT2, CNT, and CCD was measured and normalized to the area of the cortex in the image. Sections from 5 animals of each genotype were evaluated in this manner, and the figures represent the average of each genotype (\( n = 5 \)).

Quantification of distal nephron cell types in kidney sections. Quan-
tification of pendrin- and AE1-expressing ICs in the DCT2, CNT, and CCD was performed by labeling sections with antibodies against pendrin (goat), AE1 (rabbit), and the segment-specific markers calbindin D28 (mouse) and aquaporin 2 (chicken), as described above. The total number of each cell type (PCs = calbindin D28⁺ and/or aquaporin 2⁺; α-ICs = AE1⁺; non-α/non-β and β-ICs = pendrin⁺) in each segment was counted within the entire cortex of 4 WT and 4 SPAK KO animals. Similar methods were used to localize and subsequently quantify HES1 expression within the cortical distal nephron.

Quantification of pendrin and OXGR1 intracellular localization in kidney sections. Evaluation of pendrin and OXGR1 intracellular localization was performed by measuring the pixel intensity over a 6-μm distance from the tubule lumen into the cytoplasm at 0.4 μm–increm-

ent resolution using Velocity 3D Image Analysis Software (Perkin-

Elmer). Random selection of 50 cells from each of 5 WT and 5 SPAK KO mice (\( n ≥ 250 \) cells per group) was performed, and the cells were measured and compared.

Statistics. Data are presented as the mean ± SEM. Statistical anal-

ysis was performed using GraphPad Prism, version 6 (GraphPad Soft-

ware). Statistical significance was determined by a 2-tailed \( t \) test when comparing 2 groups and by 1-way randomized ANOVA, followed by a Tukey’s multiple comparisons test when comparing multiple groups. A \( P \) value of less than 0.05 was considered significant.

Study approval. The IACUCs of the University of Maryland School of Medicine and Emory University School of Medicine approved all experimental procedures.

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