Mechanisms of progression of chronic kidney disease (CKD), a major health care burden, are poorly understood. EGFR stimulates CKD progression, but the molecular networks that mediate its biological effects remain unknown. We recently showed that the severity of renal lesions after nephron reduction varied substantially among mouse strains and required activation of EGFR. Here, we utilized two mouse strains that react differently to nephron reduction — FVB/N mice, which develop severe renal lesions, and B6D2F1 mice, which are resistant to early deterioration — coupled with genome-wide expression to elucidate the molecular nature of CKD progression. Our results showed that lipocalin 2 (Lcn2, also known as neutrophil gelatinase–associated lipocalin [NGAL]), the most highly upregulated gene in the FVB/N strain, was not simply a marker of renal lesions, but an active player in disease progression. In fact, the severity of renal lesions was dramatically reduced in Lcn2−/− mice. We discovered that Lcn2 expression increased upon EGFR activation and that Lcn2 mediated its mitogenic effect during renal deterioration. EGFR inhibition prevented Lcn2 upregulation and lesion development in mice expressing a dominant negative EGFR isof orm, and hypoxia-inducible factor 1α (Hif-1α) was crucially required for EGFR-induced Lcn2 overexpression. Consistent with this, cell proliferation was dramatically reduced in Lcn2−/− mice. These data are relevant to human CKD, as we found that LCN2 was increased particularly in patients who rapidly progressed to end-stage renal failure. Together our results uncover what we believe to be a novel function for Lcn2 and a critical pathway leading to progressive renal failure and cystogenesis.

Introduction
Regardless of the initial insult, human chronic kidney disease (CKD) is characterized by progressive destruction of the renal parenchyma and the loss of functional nephrons, which ultimately lead to end-stage renal failure (ESRF). CKD represents a worldwide concern: in the United States, 102,567 patients began dialysis in 2003 (341 patients/million per year) (1), and similar rates were found in developing countries and in particular ethnic groups (2). However, these numbers are a small fraction of the millions of patients who are thought to have some degree of renal impairment. In the United States, the prevalence of chronically reduced kidney function is 11% of adults (3). Understanding the pathophysiology of CKD progression is, therefore, a key challenge for medical planning.

The mechanisms of CKD progression are poorly understood. It has been shown that reduction of the number of functional nephrons triggers molecular and cellular events promoting compensatory growth of the remaining ones (4). In some cases, this compensatory process becomes pathological, with the development of renal lesions and ESRF. Although the pathophysiology of compensation and progression is complex, unregulated proliferation of glomerular, tubular, and interstitial cells may promote the development of glomerulosclerosis, tubular cysts, and interstitial fibrosis (5–7). The molecular programs that control this cascade of events are largely unknown.

Attempts to dissect the molecular basis of CKD have been facilitated by the development of several experimental models of renal deterioration. Among these, the remnant kidney model is a mainstay, since nephron reduction characterizes the evolution of most human CKD. Consequently, this model recapitulates many features of human CKD, including hypertension, proteinuria, and glomerular and tubulointerstitial lesions. Over the last 50 years, this model has led to the discovery of critical pathways and, more importantly, to the design of therapeutic strategies to slow the progression of CKD, such as the widely clinically used renin-angiotensin inhibitors (8).

More recently, studies in various mouse strains have highlighted the importance of genetic factors in the evolution of experimental nephron reduction (9–11). We previously showed that the course and extent of renal lesions following nephron reduction vary significantly between two mouse strains: whereas FVB/N mice develop severe lesions, (C57BL/6 × DBA2)F1 (hereafter referred to as B6D2F1) mice undergo compensation alone (12). Moreover, we observed that the development of renal lesions paralleled the extent of cell proliferation (12). In fact, once the compensatory growth is achieved, a second wave of cell proliferation occurs only in the FVB/N strain. Hence, this model offers a powerful tool to unravel the transcrip tional programs and the critical mediators that are selectively activated long after nephron reduction to drive deterioration of the remaining nephrons.

In the present study, we performed an unbiased profiling of gene expression in the kidneys of the FVB/N and B6D2F1 mouse strains, 2 months after nephron reduction, when renal lesions develop and the second wave of cell proliferation is ongoing. We identified a critical mediator of progressive renal failure, namely the carrier protein lipocalin 2 (Lcn2, or neutrophil gelatinase-associated lipocalin [NGAL], siderocalin, 24p3, urocalin), and uncovered what we believe to be a novel function of Lcn2. Moreover, we elucidated a unique molecular
Research article

**Results**

**Gene profiling.** To elucidate the molecular pathways of CKD progression, we performed unbiased profiling of gene expression in remnant kidneys of two mouse strains that react differently to nephron reduction. Using microarrays containing 5,579 cDNAs, we found 70 genes whose expression levels differed significantly 2 months after nephron reduction (\(P < 0.05\)). Among these transcripts, 44 were upregulated (Supplemental Table 1; supplemental material available online with this article; doi:10.1172/JCI42004DS1) and 26 were downregulated (Supplemental Table 2) in damaged FVB/N kidneys as compared with well-preserved kidneys from B6D2F1 mice. Grouping these results by gene ontology category, we observed a range of functions for the 70 transcripts, although many of the downregulated mRNAs (38%) regulated metabolic processes (Supplemental Figure 1). The gene undergoing maximal transcriptional induction (9.95-fold change, \(P = 0.008\)) in the FVB/N lesion-prone strain was Lcn2.

**Lcn2 correlates with lesion progression in mouse and human with CKD.** Lcn2 is a member of the lipocalin superfamily (13), a family of proteins that transport hydrophobic molecules such as retinoids, fatty acids, and organic chelators of iron (14). Real-time RT-PCR confirmed that Lcn2 mRNA increased 10-fold 2 months after nephron reduction in FVB/N but not in B6D2F1 mice, while it was almost undetectable in control animals (Supplemental Figure 2). In situ hybridization and immunohistochemistry corroborated these observations and showed a marked increase in Lcn2 mRNA and protein in damaged kidneys of FVB/N mice (Figure 1, A and B). Lcn2 was predominantly found in proximal tubules and in a few ascending limbs of Henle’s loops and collecting ducts (Supplemental Figure 3A). High magnification revealed that Lcn2 was mainly located in cytoplasmic granules at the subapical zone (Supplemental Figure 3B).

By combining in situ hybridization and immunohistochemistry on serial sections, we found that a proportion of Lcn2 must have derived from the glomerular filtrate, since in some proximal tubules Lcn2 mRNA staining was negative while anti-Lcn2 staining was markedly positive (in situ and antibody). On the other hand, the majority of proximal epithelia that had undergone dilation and cystic transformation displayed both Lcn2 message and antibody staining (in situ and antibody), indicating not only endocytosis of filtered protein but ongoing local synthesis and secretion of Lcn2 (Supplemental Figure 3B). High magnification revealed that Lcn2 was mainly located in cytoplasmic granules at the subapical zone (Supplemental Figure 3B).

Renal Lcn2 mRNA (Figure 1C) and protein (Figure 1D) levels correlated with the intensity of tubular damage (\(r = 0.87, P < 0.001\) and \(r = 0.74, P < 0.01\) for mRNA and protein, respectively). In addition, we observed that renal Lcn2 protein content significant-
ly correlated with Lcn2 excretion ($r = 0.99, P < 0.01$) (Figure 1E), suggesting that the kidney is the major source of urinary Lcn2.

A time course analysis of Lcn2 expression and renal morphology revealed that the increase in both Lcn2 mRNA and protein levels preceded the development of renal lesions 4 weeks after nephron reduction (Supplemental Figure 4). Moreover, we confirmed that Lcn2 upregulation was associated with the progressive development of tubular dilations in another experimental model of CKD, the juvenile cystic kidney (jck) mice (Figure 2, A and B). Of note, these mice develop a form of polycystic kidney disease similar to human autosomal dominant polycystic kidney disease (ADPKD) (15). Last, in patients with ADPKD, who are similar to our model in displaying severe and progressive tubular dilations, LCN2 immunoreactivity was markedly increased, particularly in cysts (Figure 2C).

Urinary LCN2 was most prominent in fast progressors toward ESRF rather than in slow progressors (see Methods) ($496 \pm 146$ vs. $152 \pm 52$ ng/mg creatinine, $P < 0.01$) (Figure 2D), and it inversely correlated with residual estimated glomerular filtration rate (eGFR) ($r = -0.77, P < 0.0001$) (Figure 2E) and microalbuminuria ($r = 0.72, P < 0.0001$). Interestingly, LCN2 expression was also increased in renal tubules of kidneys from patients with either congenital nephron deficit, a pathological condition very similar to nephron reduction, or IgA nephropathy, the most common primary form of CKD (Supplemental Figure 5). Our findings in mice and humans together with recent works (16–18) suggested that Lcn2 might participate in the pathogenesis of cysts and CKD.

**Lcn2 gene inactivation prevents lesion development and cyst formation.** To determine the role of Lcn2 in progressive CKD, we performed 75% nephron reduction (Nx) in *Lcn2*−/− mice (19). To this end, we first introduced the *Lcn2* mutated allele in the lesion-prone (FVB/N) background. The *Lcn2*−/− FVB/N mice reproduced normally and had no apparent phenotype under physiological conditions (data not shown). As expected, 2 months after nephron reduction, wild-type mice developed severe renal lesions, mainly consisting of glomerulosclerosis, tubular atrophy, and cystic dilation, mild interstitial fibrosis, and multifocal mononuclear cell infiltration (Figure 3A).

However, the frequency and severity of renal lesions were dramatically reduced in *Lcn2*−/− mice. Quantification showed that Lcn2−/− mice had considerably fewer glomerular, tubular, and interstitial lesions as compared with wild-type littermates (Figure 3A). Notably, there were fewer tubular dilations and no cysts in *Lcn2*−/− mice. Consistent with these findings, renal function was better preserved in *Lcn2*−/− mice as compared with wild-type littermates, 2 months after nephron reduction. Serum creatinine and blood urea nitrogen were $5 \pm 0.5, 18 \pm 2.6$, and $11 \pm 0.6 \mu$mol/l ($P < 0.01$) and $29 \pm 1, 109 \pm 15$, and $65 \pm 4$ mg/dl ($P < 0.01$) in control, Nx *Lcn2*−/−, and Nx *Lcn2*−/− mice, respectively (Supplemental Figure 6, A and B). As expected, mean arterial blood pressure significantly increased in wild-type mice as compared with control animals ($135 \pm 7.5$ and $116 \pm 3.4$ mmHg, $P < 0.05$) 2 months after nephron reduction. The increase was of the same magnitude in *Lcn2*−/− mice ($143 \pm 2.2$ mmHg). Development of renal lesions was accompanied by severe proteinuria in wild-type mice ($6.16 \pm 1.21$ vs. $0.003 \pm 0.001$ mg/d in Nx and control mice, respectively, $P < 0.001$), whereas proteinuria was substantially decreased in *Lcn2*−/− animals ($3.30 \pm 1.03$ mg/d, $P < 0.05$) (Supplemental Figure 6C). Of note, Lcn2 inactivation did not change the course of nephron reduction in lesion-resistant C57BL/6 mice (Figure 4).
To confirm the beneficial effect of Lcn2 gene inactivation in renal deterioration and cyst formation, we bred Lcn2−/− mice with the jck mice. Notably, the severity of renal lesions was substantially reduced in double mutant jck/Lcn2−/− mice (Figure 3B). Quantification showed that the tubular dilation score was significantly lower in double mutant mice as compared with jck littermates 3 weeks after birth (Figure 3B). Collectively, these results demonstrated that Lcn2 is an effector of renal damage during CKD progression.

Iron accumulation does not account for progressive renal dysfunction. We next aimed at elucidating the mechanisms underlying the lesion-promoting effect of Lcn2. Lcn2 might act through iron mobilization (20). In fact, abnormal levels of iron accumulate in kidneys during CKD, where it may participate in the deterioration process (21, 22). Perls staining confirmed that iron content increased in damaged tubules 2 months after nephron reduction. However, iron accumulation was similar in remnant kidneys of Lcn2−/− mice as compared with wild-type littermates (Supplemental Figure 7). More importantly,
chelation of iron by desferrioxamine (DFO) (Figure 5A) unexpectedly worsened renal disease in FVB/N mice (Figure 5B). In particular, tubular dilations were more severe and diffuse in mice treated with DFO 2 months after nephron reduction. Notably, Lcn2 mRNA and protein expression were dramatically increased in kidneys of DFO-treated animals as compared with vehicle-treated counterparts (Figure 5, C and D). Proliferation of tubular cells was also significantly enhanced 2 months after nephron reduction in DFO-treated mice (Figure 5E). Hence, whereas iron deposited in the proximal tubules does not account for renal deterioration in our model, the experiments with DFO clearly show that manipulating Lcn2 levels is tightly correlated with hyperproliferation and progressive damage.

**Lcn2 is a target of EGFR signaling.** It is known that cell proliferation contributes to the development of renal lesions, and particularly to cystogenesis (23). Previous studies have suggested that Lcn2 can be induced by a number of growth factors that stimulate tubular cell proliferation (24). Among these, EGFR is of particular interest, since it is critical in the evolution of CKD (25). We therefore hypothesized that Lcn2 could act downstream of EGFR and mediate its growth effects.

To investi...
ing cells (Supplemental 9). Thus, it appears that Lcn2 is an essential mediator of the mitogenic effect of EGF in renal tubular cells.

The dual effect of Lcn2 inactivation on apoptosis. Tubular growth reflects the balance between cell proliferation and cell loss by apoptosis. Both EGFR and Lcn2 have been implicated in the control of apoptosis (25, 29). TUNEL analysis revealed an increase in apoptosis in both tubules (Figure 9A) and glomeruli (Figure 9B) of wild-type mice as compared with control animals 2 months after nephron reduction. The number of TUNEL-positive cells was significantly reduced in Lcn2−/− mice in both glomerular and tubular structures (Figure 9, A and B). However, Lcn2 silencing did not significantly affect the number of apoptotic tubular mIMCD-3 cells, regardless of the presence of EGF (Figure 9B).

Discussion

Unbiased profiling analyses offer a powerful approach to uncover critical mediators and dissect novel molecular networks of complex biological processes such as CKD progression. By combining experimental models of CKD in mice from different genetic backgrounds with microarray analyses, we have established a pivotal role for Lcn2 in regulating the progression of CKD and cyst formation. Furthermore, we have defined an important pathophysiological mechanism by which Lcn2 mediates the mitogenic effect of EGFR, consistent with its role in cell proliferation in cystogenesis. Inhibition of this pathway by Lcn2 gene inactivation or by the expression of a dominant negative EGFR isoform prevented lesion development in the transgenic mice. Conversely, overexpres-
LCn2 is a transcriptional target of EGFR. LCn2 protein (A) and mRNA (B) expression in mIMCD-3 cells, 24 hours after EGF treatment. (C and D) Overexpression of a dominant negative EGFR isoform abolishes renal LCn2 synthesis and prevents lesion development after nephron reduction. (C) LCn2 mRNA expression visualized by in situ hybridization (original magnification, ×100) and (D) renal morphology (PAS, ×200) of kidneys from control and 75% Nx wild-type and EGFR-M mice, 2 months after surgery. Data are mean ± SEM; n = 2–3 and 6–10 for in vitro and in vivo experiments, respectively. Wilcoxon test: *P < 0.05, vehicle- versus EGF-treated cells.

The mechanism for this observation remains unknown. It may result from the perfusion and filtration of serum LCn2 that we found by immunostaining in the tubules. Alternatively, since injuries to tubular cells, i.e., proteinuria, result in the expression of tubular cytokines and growth factors that ultimately lead to mesangial cell proliferation and matrix synthesis (40), it is tempting to hypothesize a crosstalk between tubular and surrounding renal cells. Studies in transgenic mice strongly support this idea. For example, it has been observed that mice that overexpress VEGF selectively in tubules developed interstitial fibrosis and glomerular disease (41). And we have previously shown that the overexpression of a dominant negative isoform of EGFR in proximal tubules prevented the development of glomerular and interstitial lesions after nephron reduction (26). On the other hand, it has been shown that interstitial scarring resulted in the loss of microvessels, which, in turn, impacted the adjacent unaffected glomeruli (42). It is worthy of note that the synthesis of paracrine mediators may increase in proliferating tubular cells (43). Hence, we speculate that by inhibiting tubular cell proliferation, LCn2 might protect glomeruli and interstitium from lesions development.

Activation of EGFR has been implicated in the evolution of CKD. Overexpression of an active EGFR form, the e-erb-B2 receptor, induc-
es tubular hyperplasia and the development of renal cysts in transgenic mice (44). Conversely, expression of a dominant negative EGFR isoform inhibits cell proliferation, leading to reduced tubular dilations after nephron reduction (26). Other genetic and pharmacological approaches have confirmed the key role of EGFR and cell proliferation in polycystic kidney diseases (45, 46), and overexpression and mislocalization of EGFR were observed in cystic epithelia of jck mice (15). On the other hand, we have previously established that EGFR acts as a central integrator of angiogenesis II pathway, a potent mediator of CKD (47). While the exact molecular networks that mediate the deleterious effect of EGFR during CKD have not been yet elucidated, our data point to Lcn2 as the crucial transcriptional target of EGFR during cyst formation and glomerulosclerosis. It is worthy of note that a very recent study showed that Lcn2 is also required for c-erb-B2 receptor signaling in breast cancer (48). In addition, our data show that Hif-1α is a critical intermediate between EGFR and Lcn2, consistent with the finding that Lcn2 is upregulated in most pathological conditions characterized by hypoxia, such as ischemia or cancers (24, 33). Whether Hif-1α is more extensively involved in the control of Lcn2 gene expression requires further investigations.

Clinical studies have suggested that urinary Lcn2 excretion might mark patients with the most severe clinical course (49), but whether Lcn2 is simply a marker of tubular damage or a key mediator of the deterioration process has been unknown. Our data now show a direct relationship between Lcn2 expression and disease progression and provide the first demonstration to our knowledge that Lcn2 is instrumental in CKD. CKD is a progressive disease, and there are many possible medical interventions over its course if the disease is recognized and treated in a timely manner. Current biomarkers of CKD progression, i.e., creatinine and albuminuria, have their limitations in terms of achieving this goal (50). An ideal biomarker should reflect tissue pathology, act as a critical component of disease, and be easily detectable by noninvasive approaches. By showing that Lcn2 unites these characteristics, we have provided strong evidence for the use of this molecule as a candidate biomarker of CKD progression.

In conclusion, we have uncovered what we believe to be a novel function of Lcn2 and highlighted its crucial role in the pathogenesis of progressive CKD. This is the first in vivo demonstration to our knowledge that Lcn2 acts as a growth regulator by mediating the mitogenic effect of EGFR signaling. Moreover, we have identified Lcn2 as one of the key effectors of renal damage and cystogenesis and one of the most promising biomarkers of CKD progression, worthy of study in large patient cohorts. We suspect that our findings will be critical in other pathological conditions that are also characterized by aberrant growth, such as cancers that demonstrate both EGFR activation and intensive Lcn2 expression (51, 52).

Methods

Animals. Mice used for these studies were FVB/N, C57BL/6, and B6D2F1 (Charles River); mutant jck bearing a Nek8 mutation (The Jackson Laboratory); transgenic EGFR-M expressing a dominant negative isoform of EGFR under the control of kidney-specific type 1 γ-glutamyl transpeptidase promoter (26); and Lcn2+/– mice (19). Lcn2−/− mice on the FVB/N genetic background were obtained using a marker-assisted speed congenic strategy. Ninety-three microsatellite markers spanning each autosomal chromosome (average distance of 14.2 cM) were used to discriminate C57BL/6 and FVB/N alleles. Heterozygous C57BL/6 Lcn2−/– mice were bred with heterozygous jck mice to obtain double-homozygous transgenic jck/Lcn2−/– mice. All experiments were performed on 9-week-old females, except for jck mice that were studied 3 weeks after birth. Animals were fed ad libitum and housed at constant ambient temperature in a 12-hour light cycle. Animal procedures were approved by the Departmental Director of “Services Vétérinaires de la Préfecture de Police de Paris” and by the ethical committee of Université Paris Descartes.

Mice were subjected to 75% Nx or sham operation (controls), as previously described (26). After surgery, mice were fed a defined diet containing 30% casein and 0.5% sodium. Several groups of mice were investigated in complementary studies. For microarray studies, 6 and 9 mice from each strain were subjected to either sham operation or Nx, respectively. For Lcn2 time course analysis, 5–6 sham-operated and 4–8 Nx mice were studied at each
time point. Transgenic studies employed EGFR-M or Lcn2−/− mice and wild-type littermates; for each group, 4–6 mice were subjected to sham operation and 10–16 mice to nephron reduction. For iron chelation experiments, 5 sham-operated and 6Nx mice were injected with 100 mg/kg DFO (Sigma-Aldrich) by subcutaneous osmotic minipumps (2004, Alzet) for 2 months. For hypoxyprobe experiments, 6 sham-operated and 6Nx mice were injected intraperitoneally with 60 mg/kg pimonidazole (Chemicon) 2 hours before sacrifice. Postischemic kidneys (2 hours renal pedicle clamping) were used as positive hypoxic controls.

Mice were sacrificed 2 months after surgery. In addition, for Lcn2 time course study, mice were also sacrificed at 4 and 6 weeks after surgery. One week before sacrifice, blood pressure was recorded in both sham-operated (n = 3) and subtotally nephrectomized (n = 6) awake Lcn2−/− and Lcn2−/− mice for 2 consecutive days, using tail-cuff plethysmography and PowerLab/4SP software (AD Instruments). Urine samples were also collected using metabolic cages from 6 mice of each experimental group over the course of 24 hours. At the time of sacrifice, the kidney was removed for morphological, protein, and mRNA studies.
The study was conducted on 87 subjects with ADPKD (40 male, 47 female; mean age 52.4 years, range 24.7–79.2 years). The mean serum creatinine level of patients was 252 ± 169.9 mmol/l, and the eGFR value (assessed using the MDRD formula; ref. 53) was 33 ± 20 ml/min/1.73 m².

Of the 87 patients, 76 were hypertensive under treatment. The decline in renal function was evaluated retrospectively over 6 years, then patients were divided into 2 groups: slow progressors (eGFR decline <4.5 ml/min/1.73 m² per year; mean, 2.4 ± 0.1; n = 52) or fast progressors (eGFR decline >4.5 ml/min/1.73 m² per year; mean, 6.0 ± 0.2; n = 35).

Kidneys from patients with ADPKD (n = 9), oligomeganephronia (n = 11), and IgA nephropathy (n = 12) were analyzed for Lcn2 expression. Normal kidneys not used for transplantation or tumor-free pole of kidneys removed for carcinoma were used as controls (n = 9).

This protocol was approved by the Hospital Plan for Clinical Research (PHRC) program of the French Ministry of Health. Informed consent was obtained from volunteers before enrollment in the study.

Cells. For siRNA transfection experiments, transient inactivation of Hif-1α expression in mIMCD-3 cells was achieved using siRNA SMARTpool from Dharmacon according to the manufacturer’s recommenda-
tions. Cells were transfected with siRNA (100 nM) using DharmaFECT4 siRNA Transfection Reagent (Thermo Fisher Scientific). Eight hours after transfection, cells were serum starved for 12 hours and then treated with 40 ng/ml EGF (R&D Systems) in serum-deprived medium for 48 hours.

For shRNA transfections, mIMCD-3 cells were stably transfected with pSuppressor Retro vector (Imgenex) containing an shRNA for Lcn2 or a scramble oligonucleotide (Dharmacon). The Lcn2 shRNA sequence contains the cloning nucleotides 5′-GGAAATATGCACAGGTATC-3′ or 5′-GCTACTGGATCAGAACATT-3′ followed by a 9-base loop and the inverted cloning sequence. In the scramble sequence, the cloning sequence is replaced by 5′-GAGCGTACCAGATTAAAGT-3′ or 5′-GATTCGACCAGATGTAT-3′. Cells stably transfected were maintained in DMEM/HamF12 medium containing 10% FBS.

For EGF experiments, cells were serum starved for 18 hours and then treated with 40 ng/ml EGF in serum-deprived medium for 24–96 hours. Cells were collected at 24 hours for Lcn2 assay and apoptosis experiments and at 24–96 hours for proliferation experiments.

cDNA microarray. RNAs were obtained from whole kidneys of 9 Nx mice from each strain using RNeasy Midi Kit (QIAGEN) according to the

Figure 9
Impact of Lcn2 inactivation on apoptosis. (A) TUNEL assay and quantification of TUNEL-positive tubular cells (white arrows) in kidneys from control, 75% Nx Lcn2+//, and Lcn2–/– mice, 2 months after surgery. Original magnification, ×400. (B) TUNEL assay and quantification of TUNEL-positive glomerular cells (white arrows) in kidneys from control, Nx Lcn2+//, and Lcn2–/– mice, 2 months after surgery. G, glomerulus. Original magnification, ×600. Because no differences were detected between wild-type and mutant control mice, results for only 1 group are shown. (C) Cell apoptosis quantification in scrambled shRNA and Lcn2 shRNA mIMCD-3 cells, 24 hours after EGF treatment. Data are mean ± SEM; n = 3 and 4 for in vitro and in vivo experiments, respectively. ANOVA followed by Tukey-Kramer test: **P < 0.01, ***P < 0.001, control versus Nx mice; ##P < 0.01, ###P < 0.001, Nx Lcn2+/+ versus Nx Lcn2–/– mice.
manufacturer’s protocol. RNAs were reverse transcribed and labeled with either cyanine Cy-3 or Cy-5. FVB/N Cy3-cDNAs and B6D2F1 Cy5-cDNAs (and, conversely, FVB/N Cy5- and B6D2F1 Cy3-cDNAs) were cohybridized on mouse cDNA microarrays containing 5,579 cDNAs expressing sequences that (Genopole). Preparations of RNAs and cDNAs and hybridization were performed according to the Genopole protocol, as previously reported (54). Six arrays were hybridized. For each array, the RNAs from 3 mice were pooled. Hybridized microarrays were scanned and images were analyzed using GenepixPro 4.0 software (Molecular Devices) by the Genopole microarray facility.

Real-time RT-PCR. Lcn2 mRNA was detected in mouse kidneys and mLincMD-3 cells by real-time RT-PCR using an ABI PRISM 7700 Sequence Detection system (Applied Biosystems). Primers (Eurogentec) were as follows: Lcn2 forward 5′-GGACCGGGCTGCTGACTT-3′ and reverse 5′-GGTG-GCCACCTGACATTGT-3′; Hif-1α (i) forward 5′-CCTGGAAACGAGT-GAAGGATTC-3′ and reverse 5′-GCTGCAATACCGAACGGTATTAA-3′; Hif-1α (ii) forward 5′-TCACCAGACAGACAGAAGAG-3′ and reverse 5′-GGCAAGACGTAAACCGCTTT-3′. Gapdh and Sdhα were used as the normalization controls in kidneys and cells, respectively.

Renal function and morphology. For mouse samples, proteinuria and blood urea nitrogen were measured using an Olympus multiparameter analyzer (Instrumentation Laboratory), while serum creatinine was evaluated by HPLC. For human samples, creatininuria and albuminuria were measured using a Hitachi 917 analyzer (Roche Diagnostics).

Kidneys were fixed in 4% paraformaldehyde and paraffin embedded, and 4-μm sections were stained with PAS, Masson’s trichrome, H&E, and picrosirius red. Ferric iron deposits were detected using Prussian blue staining according to Perls reaction. The degree of glomerular and interstitial lesions was evaluated using semiquantitative score methodology as previously described (7). The degree of tubular lesions was calculated as the number of TUNEL-positive nuclei per tubule in 20 randomly selected fields. The glomerular PI was calculated as the number of glomeruli with at least 1 PCNA-positive nuclear for the total number of glomerular nuclei. In vitro, proliferation was evaluated by counting the cell number or by using CellTiter 96 AQueous Cell Proliferation Reagent (Promega) according to the manufacturer’s instructions.

Apoptosis assay. Apoptosis was detected in 4-μm sections of paraffin-embedded kidneys by TUNEL assay using the In Situ Cell Death Detection Kit (Roche) according to the manufacturer’s protocol. The number of apoptotic cells was determined as the number of TUNEL-positive nuclei per tubule in 20 randomly selected fields. The glomerular apoptotic index was calculated as the number of glomeruli with at least 1 TUNEL-positive nuclear for the total number of glomeruli. In vitro, apoptotic cells were detected by DAPI staining, and the apoptotic index was calculated as the number of apoptotic-positive nuclei for the total number of nuclei in 10 randomly selected fields.

Measurement of urinary LCN2. Fresh urine was collected with protease inhibitors, centrifuged at 805 g at 4°C for 5 minutes, and the supernatant was removed and stored at −80°C. LCN2 was measured using ELISA (AntibodyShop). Specimens, standards, and reagents were prepared according to the manufacturer’s instructions. LCN2 levels were expressed as nanograms per milligram of creatinine. All experiments were performed in duplicate.

Statistical data. Data are expressed as mean ± SEM. Differences between the experimental groups were evaluated using ANOVA, followed when significant (P < 0.05) by the Tukey-Kramer test. When only 2 groups were compared, Mann-Whitney U or Wilcoxon tests were used. The Pearson’s correlation coefficient was used to test correlation between variables. For microarray experiments, results are expressed as a log2 of the ratio of Cy5 to Cy3. Genes with a false discovery rate (FDR) less than 0.05 (using the Benjamini-Hochberg procedure) and a fold change greater than 1.5 were considered significant. The statistical analysis was performed using GraphPad Prism software.

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