Impaired JAK-STAT signal transduction contributes to growth hormone resistance in chronic uremia

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Received for publication December 1, 2000, and accepted in revised form June 22, 2001.

Chronic renal failure (CRF) is associated with resistance to the growth-promoting and anabolic actions of growth hormone (GH). In rats with CRF induced by partial renal ablation, 7 days of GH treatment had a diminished effect on weight gain and hepatic IGF-1 and IGFBP-1 mRNA levels, compared with sham-operated pair-fed controls. To assess whether GH resistance might be due to altered signal transduction, activation of the JAK-STAT pathway was studied 10 or 15 minutes after intravenous injection of 5 mg/kg GH or vehicle. Hepatic GH receptor (GHR) mRNA levels were significantly decreased in CRF, but GHR protein abundance and GH binding to microsomal and plasma membranes was unaltered, JAK2, STAT1, STAT3, and STAT5 protein abundance was also unchanged. However, GH-induced tyrosine phosphorylation of JAK2, STAT5, and STAT3 was 75% lower in the CRF animals. Phosphorylated STAT5 and STAT3 were also diminished in nuclear extracts. The expression of the suppressor of cytokine signaling-2 (SOCS-2) was increased twofold in GH-treated CRF animals, and SOCS-3 mRNA levels were elevated by 60% in CRF, independent of GH treatment. In conclusion, CRF causes a postreceptor defect in GH signal transduction characterized by impaired phosphorylation and nuclear translocation of GH-activated STAT proteins, which is possibly mediated, at least in part, by overexpression of SOCS proteins.


Introduction

Growth failure is a major complication of chronic renal failure (CRF) (1). Children with CRF are partially resistant to the growth-promoting effect of growth hormone (GH), and the response to endogenous GH and pharmacological GH treatment are impaired in proportion to the degree of renal failure. Partial GH resistance has been reproduced in the uremic rat (2), where hepatic IGF-1 gene expression fails to increase in response to GH (3).

Several mechanisms of GH resistance in uremia have been suggested, including reduced GH receptor (GHR) gene expression (4, 5), increased IGF-1 binding to accumulated serum-binding proteins (6), and a postreceptor defect in IGF-1 action (7), though the latter is not a uniform finding (8). Moreover, since GHR gene expression is reduced less than IGF-1 gene expression (5) and GHR binding is similar to pair-fed controls in uremic rats (9), the failure of GH to stimulate IGF-1 gene expression suggests the possibility that a postreceptor defect in GH signaling may also be present.

Recently, the intracellular signaling pathways activated by GH have been elucidated (10). Binding of GH to its membrane receptor induces dimerization of the receptor, resulting in tyrosine phosphorylation of Janus-associated kinase 2 (JAK2), a tyrosine kinase associated with the intracellular domain of the receptor. Phosphorylation induces the kinase activity of JAK2, which in turn phosphorylates a group of molecules known as signal transducers and activators of transcription (STATs). The GHR-JAK2 complex activates mainly STAT5 and to a lesser degree STAT3 and STAT1. Upon phosphorylation, the STATs form dimers that translocate into the nucleus, bind to specific promoter sequences of GH-dependent genes, and transactivate or repress their transcription. Recently, a family of proteins has been identified that bind to cytokine receptor-JAK2 complexes and inhibit JAK2 kinase activity (11). Some of these suppressors of cytokine signaling (SOCS), namely SOCS-1, -2, -3, and CIS, are induced by GH via the JAK-STAT signaling pathway and inhibit the GH-induced activation of the GHR-JAK2 complex (11–13), constituting an intracellular feedback loop regulating GH sensitivity.

The importance of the JAK-STAT signaling pathway for postnatal growth has been demonstrated in selective gene-knockout models. Male mice with STAT5b deficiency and female mice with a combined deletion of STAT5a and STAT5b isofoms are severely growth retarded (14, 15), whereas SOCS-2–knockout animals develop gigantism (16). It is therefore conceivable that alterations of the JAK-STAT signaling pathway might be involved in the postulated postreceptor defect of GH action in CRF. In this study, we tested this postulate by evaluating hepatic GH signal transduction in CRF rats with evidence of GH resistance.
Methods

Experimental animals and protocols

Pituitary-intact male SD rats weighing 100–120 g were allocated to CRF or pair-fed (PF) control groups. CRF was induced by a two-stage five-sixth nephrectomy procedure. In the first intervention, approximately two-thirds of the left kidney was excised through a flank incision. Seven days later, the right kidney was removed, subsequently resulting in a state of stable CRF. In control animals, a sham procedure (exposure of kidney) was performed. Control animals received the amount of food consumed by the CRF animals on the previous day (pair feeding).

GH treatment study. Two weeks after the second surgical intervention, CRF and PF control animals were subdivided into two groups receiving either twice daily subcutaneous injections of recombinant human GH (rGH) (Genentech Inc., South San Francisco, California, USA) at a dose of 10 mg/kg/day or V (vehicle) for 7 days. This dose was selected based on pilot studies in which lower doses of GH, 1 and 5 mg/kg/day, failed to stimulate body growth and reflects the relative insensitivity of the pituitary-intact rat to exogenous GH compared with hypophysectomized rats (17). On the 7th day, the rats were sacrificed 5–6 hours after the last GH or V injection. Blood was collected by aortic puncture, and blood gas analysis was performed immediately. Sera were separated and kept at −20°C. The liver was quickly removed, snap-frozen in liquid nitrogen, and stored at −80°C.

GH-signaling study. CRF and PF control animals were studied 2 weeks after establishing CRF. The animals were anaesthetized, and for these signaling studies, a bolus of recombinant bovine GH (bGH) (5 mg/kg) (Monsanto Corp., St. Louis, Missouri, USA) or V was injected intravenously. In one set of experiments, a small piece of liver was ligated and removed for basal analysis immediately before, and the remaining liver was collected 15 minutes after the GH injection. In another series of experiments in which the rats were fasted overnight, the initial liver biopsy was omitted, and whole liver was collected 10 minutes after the intravenous administration of GH or V, snap-frozen, and stored at −80°C. A pharmacological dose of GH was used based on the pilot studies noted earlier. Indeed, pharmacological doses are commonly used in studies of hormone resistance because this ensures that differences between animals are not due to impaired hormone delivery (18). Finally, high-dose GH therapy more closely reflects the clinical situation where children with CRF are treated with pharmacological doses of GH (1).

In pilot studies with rats fed ad libitum, the levels of phosphorylated STAT5 in animals not treated with GH were variable and often elevated, reflecting the pulsatile nature of GH secretion. However, in CRF rats and their PF controls, basal phospho-STAT5 was undetectable or very low, reflecting the known suppressive effect of reduced food intake on endogenous GH secretion in rats with and without CRF (19). Accordingly, the study was restricted to CRF rats and PF controls.

Biochemical measurements and IGF-1 radioimmunoassay

Serum creatinine was measured with a Creatinine Analyzer (Beckman Coulter Inc., Palo Alto, California, USA) and tissue protein content by the Bradford method (Protein Assay Kit; Bio-Rad Laboratories Inc., Hercules, California, USA). Serum IGF-I concentrations were determined in acid-ethanol extracts using a radioimmunoassay kit (Nichols Institute Reference Laboratories, San Juan Capistrano, California, USA).

Preparation of liver membranes

Crude and purified liver plasma membranes were prepared by modifications of published protocols (20, 21). Briefly, crude membranes were obtained by homogenizing 0.5 g of liver in 0.25 M ice-cold sucrose at 20,000 rpm using an Ultra-Turrax homogenizer (IKA-LABortechnik, Staufen, Germany). After centrifugation at 1,000 g for 10 minutes, the supernatant was centrifuged at 100,000 g for 60 minutes at 4°C. The pellet containing crude liver membranes was resuspended with 0.25 M sucrose. For purification of plasma membranes, 2–3 g of liver were homogenized in cold buffer (ST buffer) containing 0.25 M sucrose and 10 mM Tris-HCl (pH 7.5). The homogenate was diluted, centrifuged at 1,500 g for 15 minutes, and resuspended in a self-forming Percoll-sucrose gradient. Centrifugation for 20 minutes at 35,000 g resulted in a viscous layer containing plasma membranes and DNA. This layer was collected, rediluted, and centrifuged at 45,000 g for 30 minutes in a gradient containing Percoll, sucrose, and CaCl2. The plasma membrane layer was collected, rediluted in ST buffer, pelleted by centrifugation at 1,500 g for 30 minutes, and resuspended in ST buffer. The activity of 5′-α-nucleotidase, a plasma membrane marker enzyme used to determine plasma membrane enrichment, was assayed in the homogenates and crude and purified plasma membrane preparations as described previously (22).

GHR binding assay

GHR binding was measured both in crude and plasma membranes by a receptor binding assay as before (23). The bGH was radiolabeled by the Iodo-Gen method (Pierce Chemical Co., Rockford, Illinois, USA). Briefly, membrane aliquots were preincubated with 3 M MgCl2 for 5 minutes to remove endogenous prebound GH and then incubated with 10−11 M [125I]-GH at 4°C overnight. Free and liver-membrane receptor-bound [125I]-GH were separated by centrifugation at 15,000 g for 10 minutes and quantitated in a gamma counter. Specific [125I]-GH binding was determined by displacement with 10−6 M unlabeled bGH.

Northern blot analysis

The cDNA probes were prepared for the analysis of GHR, IGFBP-1, CIS, and SOCS-2 mRNA as before (23). The GHR probe was prepared from a 2.2-kb full-length mouse cDNA kindly provided by F. Talamantes (University of California Santa Cruz, Santa Cruz, California, USA).
USA) (24). This cDNA encodes a sequence encompassing
the extra- and intracellular domain of the receptor.
The IGFBP-1 probe consisted of 407 bp corresponding
to nucleotide positions 486 to 892 in the rat IGFBP-1
sequence (25). The cDNA probes for CIS (770 bp) and
SOCS-2 (794 bp) were kindly provided by A. Yoshimu-
ra (Karume University, Karume, Japan) (11). The cDNA
probes were labeled with 50 µCi [32P]-dCTP by a ran-
dom primer method (Multiprime DNA labeling system,
RNA was isolated from liver as described previously
(23). Samples containing 20 µg denatured RNA were
loaded and electrophoresed on a denaturing agarose gel,
transferred, and fixed by ultraviolet cross-linking to
nitrocellulose filters, and prehybridized at 42°C for 4
hours as before (23). The mRNA levels of interest were
detected by hybridizing 32P-labeled probes to mRNA at
42°C overnight. The filters were washed and the signal
either measured directly in a PhosphorImager device
using Image Quant software (Molecular Dynamics, Sun-
dyvale, California, USA), or after exposure of filters to
Kodak XAR film (Eastman Kodak Co., Rochester, New
York, USA) by densitometric analysis (Fluor-S device and
MultiAnalyst Software; Bio-Rad Laboratories Inc.).
Solution hybridization RNase protection assay
Transcript-specific 32P-CTP labeled cRNA probes were
transcribed in vitro from respective cDNA vector con-
structs. The IGF-1 probe, a 376-bp Sau3A-EcoRI frag-
ment, was designed to detect both Ea and Eb IGF-1
mRNAs (26). The 18S riboprobe contained 80 bases
complementary to a highly conserved region of the
human RNA (27). Rat-specific CIS, SOCS-2, and
SOCS-3 cDNA was amplified by PCR, using primer
pairs published previously (13). The PCR product was
cloned into a TOPO TA plasmid (Invitrogen Corp., San
Diego, California, USA) and in vitro transcribed and
radiolabeled as described above. Twenty micrograms
total RNA was hybridized with the radiolabeled anti-
sense riboprobes at 42°C overnight. The mixture was
then incubated with an RNase digestion buffer fol-
lowed by the addition of protease K. Ethanol-precip-
itated protected hybrids were separated on a polyacry-
lamide/urea denaturing gel. Autoradiography was
accomplished by enhanced chemiluminescence (ECL; Amer-
sham Biotech, Inc.) for 1 hour at room temperature and then
quantitated densitometrically with a Fluor-S digital image
analyzer and Multianalyst software (Bio-Rad Laboratories Inc.).
Immunoprecipitation and Western immunoblot
analysis
Ab’s against STAT1, STAT3, phospho-STAT1, and
phospho-STAT3 were obtained from New England
Biolabs Inc. (Beverly, Massachusetts, USA). The anti-
STAT5 Ab that detects STAT5a and STAT5b, the anti-
phosphotyrosine (α-PY20) Ab, and protein A agarose
were from Santa Cruz Biotechnology Inc. (Santa Cruz,
California, USA). For the detection of phospho-STAT5,
an Ab from Upstate Biotechnology Inc. (Lake Placid,
New York, USA) was used. This polyclonal Ab detects
the tyrosine-phosphorylated forms of both STAT5A
and STAT5B. The GHR Ab mab263, raised against
purified rat GH receptor (28), was purchased from Bio-
genesis Inc. (Kingston, New Hampshire, USA).
Frozen liver, 100 mg, was homogenized on ice with a
Polytron (Brinkmann Instruments Inc., Westbury, New
York, USA) at maximum speed for 30 seconds in 7 vol-
umes of lysis buffer (100 mM Tris, pH 7.4, 1% Triton X-
100) containing 100 mM sodium pyrophosphate, 100
mM NaF, 10 mM EDTA, 10 mM Na3VO4, 2 mM PMSF,
and 0.1 mg/ml aprotinin. After 30 minutes in a shaker at
4°C, extracts were centrifuged for 30 minutes at 45,000 g
and the supernatants collected for further analysis. For
the detection of liver GHR, tissue was homogenized in a
hot lysis buffer containing 1% SDS and 10 mM Tris (pH
7.5) with added 0.2 mM PMSF and 0.2 mM Na3VO4.
Homogenates were boiled for 10 minutes and then cen-
trifuged as described above. Nuclear extracts were
obtained from 100 mg frozen tissue aliquots using the
NE-PER extraction kit (Pierce Chemical Co.). For JAK2
immunoprecipitation (IP), 5 µg of protein was diluted
to 1 ml in IP buffer (containing 10 mM Tris, pH 7.4, 1%
Triton X, 150 mM NaCl, 1 mM EDTA, 1 mM EGTA, 0.2
mM Na3VO4, 0.2 mM PMSF, and 0.5% Nonidet P-40)
and incubated with 2 µg JAK2 Ab’s overnight at 4°C.
The immune complexes were then adsorbed to protein
A agarose beads for 2 hours at 4°C, washed 4 times by
centrifugation, and resuspended in IP buffer.
The JAK2 immunoprecipitates or 50 µg of liver lysate
(for direct assay of all other proteins) were heated in
Laemmli buffer at 100°C for 5 minutes, electrophoresed
on a 7.5% SDS polyacrylamide gel, and electroblotted
onto nitrocellulose membranes. Blots were blocked for
1 hour in TBST buffer (10 mM Tris, pH 7.4, 138 mM
NaCl, 0.05% Tween-20 containing 3–5% nonfat dehy-
drated milk or 1% BSA. Subsequently, the blots of the
JAK-2 IP were incubated overnight at 4°C with Ab’s
against phosphotyrosine residues (α-PY20). The blots of
the liver lysates were incubated with Ab’s directed against
GHR, JAK2, STAT1, STAT3, STAT5, phospho-STAT1,
phospho-STAT3, or phospho-STAT5. After washing
times for 15 minutes in TBST, the blots were incu-
bated with secondary anti-rabbit or anti-mouse Ab con-
jugated to horseradish peroxidase (Santa Cruz Biotech-
nology Inc.) for 1 hour at room temperature and then
washed again three times. The signal on the filter was
detected by enhanced chemiluminescence (ECL; Amer-
sham Life Sciences Inc.) and exposure to Kodak XAR
film (Eastman Kodak Co.). Protein expression was quanti-
titated densitometrically with a Fluor-S digital image
analyzer and Multianalyst software (Bio-Rad Laboratories Inc.). Relative density units refer to mean pixel den-
sity with local background subtraction.
Data analysis and statistics
PhosphorImager and autoradiographic readings of spe-
cific mRNAs were normalized for 18S rRNA signal
strength and expressed as transcript/18S ratios unless
indicated otherwise. Similarly, as indicated later, phos-
phorylated protein readings were normalized for the

The Journal of Clinical Investigation | August 2001 | Volume 108 | Number 3
Table 1
Growth response and biochemical characteristics in rats with CRF and PF controls treated with human GH or V for 7 days

<table>
<thead>
<tr>
<th></th>
<th>Vehicle</th>
<th>GH</th>
<th>CRF</th>
<th>Vehicle</th>
<th>GH</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weight at start of treatment (g)</td>
<td>179 ± 17</td>
<td>178 ± 17</td>
<td>178 ± 5</td>
<td>178 ± 6</td>
<td></td>
</tr>
<tr>
<td>Weight change during treatment (g)</td>
<td>25.4±4.4A</td>
<td>55±1.6B</td>
<td>25.1±11.2A</td>
<td>34.4±9.2A</td>
<td></td>
</tr>
<tr>
<td>Food intake (g)</td>
<td>25.4±4.4A</td>
<td>55±1.6B</td>
<td>25.1±11.2A</td>
<td>34.4±9.2A</td>
<td></td>
</tr>
<tr>
<td>Food conversion ratio (g/g)</td>
<td>96.9 ± 0A</td>
<td>108 ± 0B</td>
<td>96.9 ± 19.5A</td>
<td>108 ± 11B</td>
<td></td>
</tr>
<tr>
<td>Food conversion ratio (g/g)</td>
<td>0.26 ± 0.04A</td>
<td>0.51 ± 0.02B</td>
<td>0.26 ± 0.13A</td>
<td>0.32 ± 0.07B</td>
<td></td>
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<tr>
<td>Serum creatinine (mg/dl)</td>
<td>0.70 ± 0.07A</td>
<td>0.56 ± 0.21A</td>
<td>1.8 ± 0.27A</td>
<td>1.69 ± 0.47A</td>
<td></td>
</tr>
<tr>
<td>Arterial pH</td>
<td>7.39 ± 0.02A</td>
<td>7.38 ± 0.01A</td>
<td>7.42 ± 0.09A</td>
<td>7.42 ± 0.06A</td>
<td></td>
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<tr>
<td>Plasma bicarbonate (mmol/l)</td>
<td>29.5 ± 0.7A</td>
<td>28.5 ± 2.1A</td>
<td>31 ± 1.2A</td>
<td>31.8 ± 2.2A</td>
<td></td>
</tr>
<tr>
<td>Serum IGF-1 (ng/ml)</td>
<td>625 ± 31A</td>
<td>817 ± 113A</td>
<td>828 ± 62A</td>
<td>1104 ± 69A</td>
<td></td>
</tr>
<tr>
<td>Serum creatinine (mg/dl)</td>
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<tr>
<td>Food intake (g)</td>
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<td>108 ± 0B</td>
<td>96.9 ± 19.5A</td>
<td>108 ± 11B</td>
<td></td>
</tr>
</tbody>
</table>

Results are mean ± SEM of five control and seven CRF rats per group. Serum creatinine represents mean of 100%, and individual values are expressed relative to PF vehicle mean. *P < 0.05; **P < 0.005; ***P < 0.0005.

Results
Biochemical characteristics and GH effect on body weight gain and food consumption. As seen in Table 1, the subtotal nephrectomy produced a threefold increase in serum creatinine levels, which were stable during the observation period (serum creatinine at 2 weeks, 1.66 ± 0.18 mg/dl; at 3 weeks, 1.74 ± 0.37 mg/dl). Resistance to GH therapy was evident in the CRF rats; average weight gain during GH treatment was diminished by 69% and food use (i.e., weight gain per unit weight of food consumed) by 75% compared with the PF control group. The relative weight gain in the GH-treated CRF animals was inversely correlated with serum creatinine levels (r = −0.75, P < 0.05). Total serum IGF-1 levels were lower in both CRF and PF control animals than in ad libitum–fed controls (data not shown), but did not differ between CRF and PF controls. Serum IGF-1 levels were slightly increased by GH treatment both in PF (P = 0.14) and in CRF animals (P = 0.12).

Effect of GH on IGF-I and IGFBP-1 gene transcription. Resistance to GH in CRF was also evident with respect to the expression of its hepatic target genes IGF-I and IGFBP-1. The results of the measurements of the IGF-I Ea (276 bp) transcript and the IGFBP-1 transcript are expressed in relative arbitrary units and are shown in Figure 1. Basal IGF-I mRNA levels in the V-treated group did not differ between control (IGF-I Ea [276 bp], 100 ± 9; IGF-I Ea [100 bp], 100 ± 7; IGF-I Eb, 100 ± 23 relative arbitrary units) and CRF animals (IGF-I Ea [276 bp], 103 ± 15; IGF-I Ea [100 bp], 91 ± 8; IGF-Eb, 129 ± 38, not significant [NS]). In the controls GH treatment increased IGF-I Ea (276 bp) mRNA by 79% ± 19% (P < 0.05), IGF-I Ea (100 bp) by 67% ± 17% (P < 0.05), and IGF-Eb mRNA by 158% ± 36% (P < 0.01). In contrast, GH had no significant effect in the CRF animals (IGF-I Ea [276 bp], 2% ± 38%; IGF-I Ea [100 bp], +21% ± 29%; IGF-I Eb, +22% ± 58%) (Figure 1a). The effect of GH on IGF-I mRNA levels differed significantly for the IGF-I Ea (276 bp) and the IGF-I Eb transcript (P < 0.05). Basal IGFBP-1 mRNA levels were similar in the V-treated control and CRF animals. However, the response to GH differed quantitatively. In the controls GH treatment suppressed IGFBP-1 gene expression by 86% ± 4% (P < 0.05), while in the CRF group by...
GHR gene expression, protein abundance, and binding capacity. Hepatic GHR mRNA levels were significantly reduced in the V-treated CRF animals of the 7-day GH treatment study (Figure 2a, \( P < 0.05 \)). Similarly, in the animals used for the GH signaling experiments, GHR mRNA levels were decreased by 61\% \pm 4\% compared with PF controls (\( P < 0.001 \)). Chronic GH treatment did not change GHR gene expression in either the PF (117\% \pm 20\% of PF-V), NS) or CRF group (113\% \pm 11\% of CRF-V, NS). In contrast to the difference in GHR mRNA levels, Western immunoblotting of the GHR protein showed a band around 130 kDa that was equally abundant in the CRF and in the control animals in both studies (Figure 2b).

In addition to the analysis of GHR protein expression, measurements of specific GH binding to liver membranes were also performed. In the crude microsomal isolates, \( \alpha \)-S'-nucleotidase activity was only slightly higher (166 \pm 6 nmol/mg protein/min) than in native liver homogenates (105 \pm 4 mmol/mg/min). In contrast, the mean \( \alpha \)-S'-nucleotidase activity of the membranes obtained by Percoll gradient centrifugation was 1,451 \pm 134 mmol/mg protein/min vs. 77 \pm 19 mmol/mg protein/min in homogenates, indicating a 22-fold enrichment of plasma membranes by this procedure. Consistent with the results of Western immunoblot analysis, specific GH binding to the crude microsomes or the purified plasma membranes did not differ between CRF animals and controls (Figure 2c).

**JAK-STAT signal transduction.** The protein levels of JAK2, STAT1, STAT3, and STAT5 in liver lysates were not different between CRF and control animals (Figure 3a). In contrast, differences in the level of phosphorylated proteins were evident after GH treatment (Figure 3b). In the basal state (V-treated animals), little phosphorylation of JAK2, STAT1, and STAT3, and no phosphorylation of STAT5, was detected in either group. JAK2 tyrosine phosphorylation increased 10 minutes after GH injection, but this response was blunted in the CRF animals (Figure 3, b and c). Similarly, GH-stimulated phosphorylation of STAT5 was 75\% lower in CRF animals than in controls, as determined both by absolute signal strength (\( P < 0.0001 \)) and when expressed relative to the corresponding level of STAT5 protein (\( P < 0.0001 \)). In these experiments the V- and the GH-treated tissue was obtained from different animals. In another experiment (six animals per group), where tissue was obtained immediately before and 15 minutes after GH in the same animal, a similar reduction of STAT5 phosphorylation was observed (PF, 100 \pm 13, vs. CRF, 31 \pm 6 arbitrary units; \( P < 0.0005 \)). A significant positive relationship between the relative phosphorylation of JAK2 and STAT5 after GH administration was observed (\( r = 0.87; P < 0.001 \)) (Figure 3d). Regression analysis showed that relative STAT5 and JAK2 phosphorylation levels were inversely correlated with serum creatinine levels (\( r = -0.91 \), and \( r = -0.89 \), respectively, \( P < 0.0001 \)).

Analogous differences between CRF and PF control animals were found with respect to STAT3 and STAT1 tyrosine phosphorylation. Absolute and relative phosphorylation of STAT3 were decreased in the CRF animals by 68\% and 75\%, respectively (Figure 3, b and c). The GH-induced STAT1 phosphorylation was very weak and more variable than the phospho-STAT3 and phospho-STAT5 signals, but also tended to be lower in the CRF animals (Figure 3, b and c).

To see whether the low whole-cell phospho-STAT levels were reflected in the nucleus, STAT phosphorylation was also measured in nuclear extracts 15 minutes after GH injection (Figure 4a). As in the whole cell, nuclear phospho-STAT5 was low in the CRF animals compared with the controls (38 \pm 22 vs. 100 \pm 14 arbitrary units, \( P < 0.05 \)). Nuclear phospho-STAT3 levels were on average lower in the CRF group, but this did not reach statistical significance (52 \pm 20 vs. 100 \pm 19 arbitrary units, \( P = 0.09 \)). Phospho-STAT1 was undetectable in the nuclear extracts both in CRF and control animals.

**SOCS-2, CIS, and SOCS 3 gene expression.** SOCS-2 mRNA was detectable by Northern blot analysis only in animals that received prolonged GH treatment (Figure 4b). The 18S-corrected SOCS-2 mRNA levels were increased 2.5-fold in the GH-treated CRF animals, compared with the GH-treated PF controls (250 \pm 41 vs. 100 \pm 10 arbitrary units, \( P < 0.005 \)). As shown in Table 2, the increased

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**Figure 2**

Hepatic GHR mRNA levels(a), but not protein abundance(b) or GH membrane binding(c), are reduced in CRF. Hepatic GHR mRNA was measured by Northern blot analysis and GHR protein by Western immunoblot analysis. Specific binding of radiolabeled bGH to receptors in purified liver plasma membranes and microsomes was measured by displacement with an excess of unlabeled bGH. All measurements were made in livers from the same animals. Data are expressed relative to PF vehicle-treated control mean value, assigned a value of 100. Bars indicate mean \pm SEM. n = 4–6 rats/group. *\( P < 0.05 \).
SOCS-2 gene expression in GH-treated CRF animals was confirmed by RNase protection assay (RPA) (909 ± 137 vs. 482 ± 17 arbitrary units, $P < 0.01$). This sensitive assay also detected the SOCS-2 mRNA in the V-treated animals. Although on average higher in the CRF group, the levels were not significantly different from the PF control values because of the variability of basal expression. However, bivariate ANOVA showed a significant GH-independent stimulatory effect of CRF on SOCS-2 expression ($P < 0.05$). CIS mRNA expression, detectable only by RPA, was increased by GH treatment in the CRF and control animals ($P < 0.0001$). CIS expression tended to be higher in the CRF groups, but significance was not reached.

SOCS-3 mRNA levels, also only detectable by RPA, were not elevated in either group when measured 5–6 hours after the last GH injection, compared with V injection, a finding consistent with its rapid and transient treatment response (13). Accordingly the V- and GH-treatment values were regarded as reflecting basal values and were pooled for analysis. This showed that the SOCS-3 mRNA levels were significantly higher in the CRF animals ($P < 0.05$; Table 2).

**Discussion**

In this study we set out to determine whether impaired GH-mediated signal transduction could contribute to the GH resistance that develops in CRF, and our results show that a defect in the JAK-STAT signal transduction pathway is indeed present in uremia. To evaluate GH signaling we used a rat model of nonacidotic CRF, severe enough to cause GH resistance. In these animals hepatic GHR binding and protein levels were similar to PF, sham-operated controls. Despite this, GH treatment failed to stimulate IGF-I gene expression, and, compared with the controls, suppression of IGFBP-1 gene expression and body weight gain were severely blunted. The key alterations in signal transduction uncovered were a diminution in GH-induced tyrosine phosphorylation of the GH receptor–associated tyrosine kinase JAK2 and reduced tyrosine phosphorylation and nuclear translocation of GH-activated STAT proteins. We also found increased mRNA levels of the intracellular feedback regulatory proteins SOCS-2 and SOCS-3 in CRF rats that we propose may contribute to the impaired JAK-STAT signaling.

Previously, it has been suggested that GH resistance in CRF may be caused by a defect in GHR expression. Support for this has been provided by studies in uremic rats that showed reduced hepatic GHR mRNA levels. However, this reduction could in part be attributed to reduced food intake (4, 5). Indeed, others observed no difference in GHR mRNA levels between CRF and PF control animals (9, 29), questioning an independent
suppressing effect of uremia. Similarly, whereas an early study suggested reduced GH binding to crude rat liver membranes in CRF compared with ad libitum–fed control animals (30), a later study showed that specific GH binding in CRF was not different from PF controls (9). On the other hand, reduced GHR protein expression has been found using immunohistochemistry in uremic rat growth cartilage (31). Clinical studies using circulating GH-binding protein levels, largely derived by cleavage of the extracellular GHR domain, as a surrogate marker of tissue GHR abundance in children with CRF have yielded conflicting results; both low (32) and normal GHBP plasma levels have been reported (33).

In this study we observed a significant reduction of steady-state GHR mRNA levels by 20–50%, whereas GHR protein levels, measured by Western immunoblot analysis, were unchanged in the CRF rats compared with the PF controls. The reason for the disparity between the GHR mRNA and protein levels in the CRF animals was not examined, but appears to be due to a posttranscriptional event such as increased translational efficiency or decreased receptor protein degradation. To confirm our immunoblot findings and to exclude the possibility that resistance to GH is caused by a maldistribution of the plasma membranes, we measured GH-specific binding to crude microsomes and purified plasma membranes. Consistent with our measurements of GHR protein levels, receptor binding in both preparations was unaltered by CRF. Hence, the GH resistance in the CRF animals cannot be attributed to an alteration in receptor number, affinity, or distribution and suggests a more distal defect.

The levels of the key-signaling proteins of the JAK/STAT signal-

### Table 2

<table>
<thead>
<tr>
<th></th>
<th>PF control</th>
<th>GH</th>
<th>Vehicle</th>
<th>GHR</th>
<th>CRF</th>
<th>CRF effect</th>
<th>GH effect</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>100 ± 25</td>
<td>457 ± 85</td>
<td>219 ± 61</td>
<td>602 ± 126</td>
<td>NS</td>
<td>P &lt; 0.0001</td>
</tr>
<tr>
<td>CIS</td>
<td></td>
<td>100 ± 26</td>
<td>482 ± 17</td>
<td>222 ± 68</td>
<td>909 ± 137</td>
<td>P &lt; 0.05</td>
<td>P &lt; 0.0001</td>
</tr>
<tr>
<td>SOCS-2</td>
<td></td>
<td>100 ± 26</td>
<td>115 ± 18</td>
<td>179 ± 26</td>
<td>167 ± 45</td>
<td>NS</td>
<td>NS</td>
</tr>
<tr>
<td>SOCS-3</td>
<td></td>
<td>100 ± 26</td>
<td>115 ± 18</td>
<td>179 ± 26</td>
<td>167 ± 45</td>
<td>NS</td>
<td>NS</td>
</tr>
</tbody>
</table>

Tissue was obtained 5–6 hours after the last GH injection. The mRNA levels were determined by RPA. Values were corrected for 18S rRNA levels measured in the same assay. Data are expressed relative to PF vehicle mean value, which was assigned a value of 100. Results are mean ± SEM of three to seven rats per group. Superscript letters indicate significant differences between the groups within each row (Newman-Keuls test, P < 0.05); groups with common superscript letters do not differ from each other. Results of two-way ANOVA are given in the two last columns.
mice: circulating IGF-1 levels are decreased in males deficient for the STAT5b isoform and in females if STAT5a and STAT5b genes are mutated (14, 15). Whereas no direct DNA-binding site for STATs has been found to date, GH-dependent IGF-1 transcription may be mediated by the cooperative action of the JAK-STAT pathway and one or more nuclear cofactors (36, 37). Hepatic IGFBP-1 expression is rapidly suppressed by GH (34) by a mechanism involving c-Fos and c-Jun (38). GH stimulates hepatic c-Fos transcription by binding of STAT1 and STAT3 homo-and heterodimers to a Sis-inducible element in the c-Fos promoter (38, 39). In addition, GH activates c-Fos expression by stimulation of c-Raf and the MAP kinase pathway (10). The activation of both pathways requires phosphorylation of the GHR-associated kinase JAK2. Hence, the attenuated suppression of IGFBP-1 gene expression in GH-treated CRF rats, as well as the absent stimulation of IGF-1 expression in these animals, are compatible with diminished JAK2 and/or STAT activity in uremia.

Recently, a family of intracellular proteins has been defined that serve as intracellular negative feedback regulators of the JAK-STAT pathway by binding to cytokine receptor-JAK signaling complexes (11). Among the eight members of this family identified to date, GH stimulates the expression of SOCS-1, -2, -3, and CIS (11–13). When overexpressed in vitro, these GH-inducible proteins partially or completely inhibit GH-dependent JAK2 activation (12). Whereas SOCS-1 and SOCS-3 appear to be stronger inhibitors than SOCS-2 and CIS in transfection models, in vivo evidence for a role of SOCS in growth regulation has been provided for SOCS-2 and CIS. Deletion of SOCS-2 causes IGF-1 hyperexpression and gigantism in mice (16), whereas CIS overexpression causes severe growth retardation (40). To test whether altered expression of one of these inhibitors could be implicated as a potential cause of the depressed JAK2 phospho-

rlation, we measured mRNA levels of SOCS-2, SOCS-3, and CIS. We observed a wide variability of basal hepatic SOCS-2 expression, with a trend toward higher levels in CRF. However, after 1 week of GH treatment, SOCS-2 was clearly hyperexpressed in CRF animals and was more than twofold higher than in GH-treated controls. Interestingly, in the septic rat, another model of acquired GH resistance, treatment with GH also increases SOCS-2 mRNA levels (41). Basal SOCS-3 mRNA levels were modestly elevated in CRF. Whether the protein levels of these hyperexpressed genes are also increased in uremia remains to be established.

The mechanism for the increase in SOCS mRNA levels in the presence of impaired GH-activated JAK2-STAT signaling in uremia is not readily apparent. One potential process could involve the action of other members of the cytokine family (11), since several cytokines induce signal transduction by activating other members of the Janus kinase family such as JAK1, JAK3, and Tyk2 (42). Accordingly, if there were selective inhibition of JAK2 activation in the uremic state, it is conceivable that the increase in SOCS gene expression reflects cytokine sig-

naling by way of a different Janus kinase. It is also possible that the increase in SOCS expression in uremia might be mediated by GH through a non–STAT-mediated pathway, for there is recent evidence, at least for SOCS-3, that nuclear STAT-binding sites are not essential for GH activation of the SOCS-3 promoter (43). Finally it should be kept in mind that the measured level of gene expression reflects the impact of uremia on several interacting processes including gene transcription and stability, cross-talk between signaling pathways, and the action of a variety of hormones and cytokines.

An acquired defect of GH-induced JAK-STAT signaling has been reported recently in rats with acute endotoxemia (18, 44). This manifests as a decrease in GH-induced JAK2 phosphorylation in the absence of a change in GHR abundance (18), diminished hepatic accumulation of nuclear STAT5P and STAT3P, and diminished binding of STAT5 to a GH-response element from the serine protease 2.1 inhibitor promoter (44). Interestingly, while the phosphorylation of the JAK2 protein was diminished, total JAK2P was unchanged because of an endotoxin-induced increase in total Jak2 protein levels. These changes were accompanied by an acute increase in SOCS-3 and CIS, and to a lesser degree SOCS-2 expression, findings confirmed by others (45). These studies are supported by in vitro data that indicates that proinflam-
matory cytokines such as IL-6 and IL-1β released during an endotoxin challenge may stimulate SOCS as part of an autoinhibitory intracellular feedback loop, causing concomitant GH resistance (43, 45, 46). This is another example where sensitivity to some cytokines is maintained, despite resistance to GH. These reports may be relevant to the situation in patients with end-stage renal disease, for chronic subclinical inflammation is common in dialysis patients and appears to be a major cause of malnutrition and increased cardiovascular morbidity (47). It is tempting to speculate that persistent subclinical inflammation with proinflammatory cytokine release may contribute to the malnutrition of uremia by worsening the GH resistance in these patients by increasing SOCS protein production.

Taken together, we have defined a novel postreceptor mechanism of GH resistance in CRF. We found that despite a lack of change in GHR abundance, tyrosine phosphorylation of the GHR-associated tyrosine kinase JAK2 is diminished in uremia. This was associated with impaired phosphorylation and nuclear translocation of the downstream STAT-signaling proteins. Of note, expression of SOCS-2 and -3 mRNA levels were increased in CRF, and we propose that if accompanied by an increase in protein expression, then by suppressing JAK2 activity this may be a cause of the signaling defect. However, since regulation of the JAK-STAT pathway involves more than just the SOCS proteins, it is likely that several molecular mechanisms operative in uremia contribute to the impairment in GH signal transduction. These might include increased protein tyrosine phosphatase activity with more rapid dephosphorylation of the GHR and its signaling proteins, altered GH receptor and signaling processes.
protein degradation, and abnormal expression of novel signal regulatory proteins (11, 48). In conclusion, it appears that resistance to GH in uremia is not caused by a change in GHR number, but rather is the result of a defect in GH signal transduction. This signaling defect, together with resistance to IGF-I, likely contributes to the retarded body growth of children and the wasting that occurs in adults with renal failure.

Acknowledgments

This study was supported by grants from the Department Veterans Affairs (Merit Review grant to R. Rabkin), the NIH (grant 5R21 DK-53613 to R. Rabkin), and the Max Kade Foundation and the NIKI Foundation (to F. Schaefer).