Nuclear Factor-κB1 (p50) Limits the Inflammatory and Fibrogenic Responses to Chronic Injury

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In this study we addressed the role of the nuclear factor (NF)-κB1/p50 subunit in chronic injury of the liver by determining the inflammatory and fibrotic responses of nfd1-null mice in an experimental model that mimics chronic liver disease. Mice received repeated hepatic injuries throughout 12 weeks by intraperitoneal injection of the hepatotoxin carbon tetrachloride. In response to nfd1<sup>−/−</sup> mice developed more severe neutrophilic inflammation and fibrosis compared to nfd1<sup>+/+</sup> mice. This phenotype was associated with elevated heparin binding and dimerization motifs as well as the nuclear localization signal. The prototypic NF-κB consists of a dimer of p50 and p65 and is classical held in an inhibitory complex with IκBα. In response to stimulation, IκBα is phosphorylated via activation of the IKK complex and is subsequently polyubiquitinated and targeted for degradation by the 26S proteosome. This leads to release of the p50/p65 dimer that translocates to the nucleus to activate transcription of genes containing their DNA-binding motif, known as a κB-binding site.

This classical mechanism of activation and operation of NF-κB is however complicated by the fact there are at least 10 potential homodimeric and heterodimeric combinations of Rel factors. An additional level of complication is that there is emerging evidence from both in vitro and in vivo studies that the individual Rel proteins have distinct biological activities, however the precise physiological role of each protein is not fully understood. Of particular interest, transcription repression functions have been attributed to homodimers of p50 as well as to NF-κB heterodimers containing RelB. These repressive properties may serve to fine tune NF-κB-directed immune and inflammatory responses. Studies using mice carrying targeted disruptions of the Rel family genes are providing important evidence for distinct biological func-

Nuclear factor (NF)-κB is a cardinal regulator of immune and inflammatory responses, controlling the expression of genes encoding cytokines, chemokines, adhesion molecules, and regulators of the cell cycle and apoptosis. Prolonged activation of NF-κB leads to perpetuated inflammatory responses and as such the transcription factor is a target for a range of anti-inflammatory drugs used to treat diseases including rheumatoid arthritis, asthma, and inflammatory bowel disease. NF-κB is a dimeric transcription factor generated from combinations of Rel (c-Rel), p65 (RelA), RelB, p50/p105 (NFκB1), and p52/p100 (NFκB2) all of which contain an evolutionary conserved Rel homology domain that includes the DNA-binding and dimerization motifs as well as the nuclear localization signal. The prototypic NF-κB consists of a dimer of p50 and p65 and is classical held in an inhibitory complex with IκBα. In response to stimulation, IκBα is phosphorylated via activation of the IKK complex and is subsequently polyubiquitinated and targeted for degradation by the 26S proteosome. This leads to release of the p50/p65 dimer that translocates to the nucleus to activate transcription of genes containing their DNA-binding motif, known as a κB-binding site.

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tions for these factors. Nfkβ1−/− mice effectively lack two proteins, a 105-kd non-DNA-binding cytoplasmic protein (p105) and the p50 NF-κB subunit that is generated from p105 by proteolytic processing. Nfkβ1−/− mice develop normally but have multisite inflammatory responses and B cell function. Mice deficient in relB have a variety of pathological conditions including multiorgan inflammatory cell infiltrates, myeloid hyperplasia, and a variety of pathological conditions including multiorgan inflammatory cell infiltrates, myeloid hyperplasia, and splenomegaly because of extramedullary hyperplasia. However, mice lacking both relB and nfkβ1 spontaneously develop severe multiorgan inflammation, which suggests that these factors may operate in concert to limit inflammatory events.

In this study we have investigated the role played by nfkβ1 in the injury/wound-healing response by using a well-established model of iterative liver injury that mimics a repetitive inflammation-driven injury and wound-healing process. Importantly, this model of chronic liver disease significantly differs from the acute injury models that have previously been used to conclude lack of a role for p50 in liver disease. By comparing the response of nfkβ1−/− and nfkβ1−/− to iterative injury with the hepatotoxic carbon tetrachloride (CCL4), we show that lack of p50 results in an exacerbated neutrophilic inflammatory response and development of severe fibrosis. We describe a mechanism underlying this previously undescribed phenotype of nfkβ1 mice that involves the absence of a p50-dependent transcriptional repression of tumor necrosis factor (TNF)-α expression by the histone deacetylase HDAC1.

Materials and Methods

Nfkβ1−/−/Nfkβ1+/+ Mice and 12-Week CCL4 Injury Model

Experiments were performed on C57Bl/6;129PF2/J nfkβ1−/− and F2 hybrid nfkβ1+/+ mice. Because F2 hybrids carry random combinations of alleles derived from both the C57Bl/6 and 129 genetic backgrounds they are more appropriate physiological controls than inbred mice of either parental strain (http://jaxmice.jax.org/info/hybrid.html). Breeding pairs were purchased from Jackson Laboratories (Bar Harbor, ME) and were housed in pathogen-free conditions. CCL4 and olive oil were purchased from Sigma Chemical Co., Dorset, UK. Age-matched male nfkβ1−/− and nfkβ1+/+ mice were treated with CCL4 at 1 μl [CCL4:olive oil (1:3 {v/v})/g body weight by intraperitoneal injection twice weekly for 12 weeks]. At days 1, 3, 5, 7, and 14 after the last CCL4 injection animals were killed by cervical dislocation and livers harvested. To determine age-associated differences in liver pathology mice were maintained in pathogen-free conditions for 12 months before harvesting of livers.

Cell Culture

Hepatic stellate cells (HSCs) were isolated from normal livers of nfkβ1−/− and nfkβ1+/+ male mice. Livers were washed in Hanks’ buffered saline minus calcium and then cut into small pieces and digested with collagenase and pronase. Enzymatic digestion was followed by discontinuous density centrifugation in 11.5% OptiPrep (Life Technologies, UK). HSCs were cultured on plastic in Dulbecco’s modified Eagle’s medium supplemented with 100 U/ml penicillin, 100 μg/ml streptomycin, 2 mmol/L L-glutamine, and 16% fetal calf serum and maintained at 37°C at an atmosphere of 5% CO2. All experiments using HSC-derived myofibroblasts were performed on cells passaged between two and four times.

Plasmid DNA, Cell Transfections, and Reporter Assays

All plasmid DNA was prepared using a commercial DNA extraction and isolation kit (Maxiprep; Qiagen). Expression vectors for HDAC1 and p50 were gifts from Dr. Neil Perkins (University of Dundee, Dundee, Scotland) and Professor Ron Hay (University of St. Andrews, Fife, Scotland), respectively. Reporter constructs ppx-TNF-α-luc (referred to here as pTNF-α-Luc) was kindly provided by Dr. Peter Johnson (NCI, Frederick, MD). Culture-activated nfkβ1−/− and nfkβ1+/+ mouse HSCs were transfected using the nonliposomal Effectene protocol (Qiagen) with 1 μg of reporter plasmid DNA, ±2 μg of expression plasmid/control vector, and 10 ng of control Renilla plasmid pRLTK according to the manufacturer’s instructions. For experiments using trichostatin A (TSA), 24 hours after transfection, nfkβ1−/− and nfkβ1+/+ mouse HSCs were treated with TSA (500 nmol/L) for 24 hours and reporter gene activity assay performed. Luciferase assays were performed using a dual luciferase kit (Promega) according to the manufacturer’s instructions. Firefly luciferase activities were normalized for differences in transfection efficiency by measurement of the activity of co-transfected pRLTK.

Sodium Dodecyl Sulfate-Polyacrylamide Gel Electrophoresis and Immunoblotting

Whole cell extracts were prepared, and protein concentration of samples was determined using a Bradford DC assay kit (Bio-Rad). Whole cell extracts (30 or 20 μg) from activated nfkβ1−/− and nfkβ1+/+ mouse HSCs, respectively, were then fractionated by electrophoresis through a 9% sodium dodecyl sulfate-polyacrylamide gel. Gels were run at a 100 V for 1.5 hours before transfer onto nitrocellulose. After blockade of nonspecific protein binding, nitrocellulose blots were incubated for 1 hour with primary antibodies diluted in Tris-buffered saline (TBS)/TWEEN 20 (0.075%) containing 3% Marvel. Mouse monoclonal horseradish peroxidase-conjugated antibody recognizing HDAC1 (Upstate Biotechnology, Lake Placid, NY) was used at 1 μg/ml. Mouse monoclonal antibodies directed against α-smooth muscle actin (SMA) (1:2000), desmin (1:2000), and β-actin (1:1000) were obtained from Sigma. Rabbit polyclonal directed against gial fibrillary acidic protein (1:1000) was obtained from Sigma. Rabbit polyclonal directed against p75 (1:2000) was ob-
tained from Promega. After incubation with primary antibodies, blots were washed three times in TBS/Tween 20 before incubation for 1 hour in goat anti-mouse or mouse anti-rabbit horseradish peroxidase conjugate antibody at 1:2000 dilution in TBS/Tween 20 (0.075%) containing 3% Marvel. After extensive washing in TBS/Tween 20, the blots were processed with distilled water for detection of antigen using the enhanced chemiluminescence system (Amersham Biosciences).

**Semiquantitative Reverse Transcriptase-Polymerase Chain Reaction (RT-PCR)**

Total RNA was purified from isolated cells or frozen livers using the Total RNA purification kit (Qiagen, UK) following the manufacturer’s instructions and was used to generate first strand cDNA using a random hexamer primer [p(dN)6] and MMLV reverse transcriptase. Oligonucleotide primers for PCR amplification of murine β-actin, interleukin (IL)-6, p53, p52, RelB, and c-Rel have been previously published. Primers for mouse TNF-α 5′-acg gca gca tgt atc tca-3′ (sense) and 5′-tac tgt gcc aga tgt acc-3′ (antisense), for mouse ICAM 1, 5′-acc cca agg acc cca agg aga-3′ (sense) and 5′-aga gcc gca gag cca aag aag-g3′ (antisense). Primers for mouse HDAC1 were 5′-cga gac ggg att gag gcc -3′ (sense) and 5′-tct tgt tgt cgg tag tga-3′ (antisense). PCR reactions comprised of 1 μl of cDNA, 100 ng each of sense and antisense oligonucleotide primers, 2.5 μl of optimized TaqPCR buffer (Promega), 0.4 mmol/L dNTP mixture, and 2 U of Taq polymerase in a total reaction volume of 25 μl. After an initial 5-minute incubation at 94°C, PCRs were performed using a 1-minute annealing step (TNF-α at 55°C, ICAM1 at 59.3°C, HDAC1 at 51.6°C), followed by a 2-minute elongation step at 72°C and a 45-second denaturation step at 94°C. Thirty-five PCR cycles were performed for amplification of all cDNAs, followed by a final elongation for 10 minutes at 72.0°C. PCR products were separated by electrophoresis through a 1% agarose gel and detected by ethidium bromide staining.

**Quantitative TaqMan RT-PCR**

18s rRNA TaqMan primers and probe were purchased from Applied Biosystems (UK) and the reaction was conducted according to the manufacturer’s instructions using 10 ng of cDNA for each reaction. Proccollagen I mRNA expression was determined using forward primer 5′-tgt ttt acc tac agc agc ctt ggt-3′, reverse primer 5′-cat gac tgt ctt gcc cca agt t-3′ and probe 5′-atg gct gcc gca gtc aca-3′ quencher, TAMRA and fluorophore, FAM. For α-SMA forward primer 5′-tct aca gac ctt ggg ttg tct ggt-3′, reverse primer 5′-aaa aac cac gag taa cta atc a-3′ and probe 5′-tcc aca ttc tcc cca a-3′ quencher, TAMRA and fluorophore, FAM. For TIMP-1, forward primer 5′-gga tgg aca ttt att ctc cag tgt-3′, reverse primer 5′-tcg tgt tct ggt tag tca-3′ (sense) and 5′-tct cta gga gcc ccc atg tgc-3′ (antisense), for mouse ICAM 1, 5′-acc cca agg acc cca agg aga-3′ (sense) and 5′-aga gcc gca gag cca aag aag-g3′ (antisense). Primers for mouse HDAC1 were 5′-cga gac ggg att gag gcc -3′ (sense) and 5′-tct tgt tgt cgg tag tga-3′ (antisense). PCR reactions comprised of 1 μl of cDNA, 100 ng each of sense and antisense oligonucleotide primers, 2.5 μl of optimized TaqPCR buffer (Promega), 0.4 mmol/L dNTP mixture, and 2 U of Taq polymerase in a total reaction volume of 25 μl. After an initial 5-minute incubation at 94°C, PCRs were performed using a 1-minute annealing step (TNF-α at 55°C, ICAM1 at 59.3°C, HDAC1 at 51.6°C), followed by a 2-minute elongation step at 72°C and a 45-second denaturation step at 94°C. Thirty-five PCR cycles were performed for amplification of all cDNAs, followed by a final elongation for 10 minutes at 72.0°C. PCR products were separated by electrophoresis through a 1% agarose gel and detected by ethidium bromide staining.

**Histology and Immunohistochemistry**

Sirius Red and hematoxylin and eosin staining was performed as previously described. For immunohistochemical staining for SMA-α, slides were dewaxed in xylene and dehydrated in alcohol. Antigen retrieval was achieved by microwaving in citric saline for 15 minutes. Endogenous peroxidase activity was blocked by hydrogen peroxide pretreatment for 10 minutes then further blocked using Vector Laboratories avidin/biotin blocking kit, 3 drops/section, with TBS washes between each stage. Slides were incubated with complete culture medium for 20 minutes, then in mouse anti-human α-SMA-fluorescein isothiocyanate (Sigma) diluted 1:30,000 in TBS, and applied to the slides and incubated overnight at 4°C. Slides were washed and rabbit anti-fluorescein isothiocyanate antibody (DAKO) diluted 1:4000 and incubated at room temperature for 30 minutes. Slides were then washed and incubated at room temperature for 30 minutes with a 1:200 dilution of biotin swine anti-rabbit (DAKO). Slides were washed in TBS and a streptavidin biotin-peroxidase complex at 1:200 dilution (DAKO) was incubated at room temperature for 30 minutes. Cells positive for α-SMA were visualized by diaminobenzidine (DAB) staining. Slides were counterstained with Mayer’s hematoxylin for 30 seconds, dehydrated, cleared in xylene, and mounted in DPX. Immunohistochemical staining for neutrophils was performed using the neutrophil antibody clone 7/4 (Serotec). Slides were initially processed in the manner described for α-SMA immunostaining. The primary antibody (Serotec, UK) was diluted 1:100 and incubated overnight at 4°C, secondary and anti-IgG horseradish peroxidase-conjugated tertiary antibodies were incubated for 20 minutes (Vector Laboratories, UK) and neutrophil recruitment was visualized by DAB staining. Immunohistochemical staining for TNF-α was performed using the goat anti-mouse TNF-α antibody (Serotec). Dewaxed and microwave-treated slides were initially blocked for endogenous peroxidase activity by hydrogen peroxide pretreatment for 10 minutes then further blocked using Vector Laboratories avidin/biotin blocking kit, 3 drops/section, with phosphate-buffered saline (PBS) washes between each stage. The primary
antibody (Serotec, UK) was diluted 1:500 and incubated overnight at 4°C, secondary and anti-IgG horseradish peroxidase-conjugated tertiary antibodies were incubated for 20 minutes (Vector Laboratories, UK) and TNF-α was visualized by DAB staining. Slides were counterstained with Mayer’s hematoxylin for 30 seconds, dehydrated, and mounted in DPX. For TNF-α and α-SMA dual immunostaining the methods are as described above for the individual antibody stains with the exception of an additional serum-blocking step between the DAB visualization of α-SMA and the incubation of goat anti-mouse TNF-α antibody. To distinguish between TNF-α and α-SMA DAB plus nickel (black) was used to visualize TNF-α.

**Cell Counts and Pathology Scoring**

All cell counts were performed in a blinded manner and were representative of an average number of positive cells/field counted in 20 high-power fields from four to five mice per time point of recovery. Statistical differences between nfkβ1−/− and nfkβ1+/+ cell counts were calculated using a two-tailed unpaired t-test, assuming equal variation in SD between the two populations using Graph Pad Prism computer software. Histopathological analysis and grading of liver sections for degree of fibrosis and inflammation was performed by an expert pathologist (H.M.-S.) in a blinded manner.

**TNF-α Enzyme-Linked Immunosorbent Assay (ELISA)**

Livers or HSCs were lysed in Dignam buffer A supplemented with a cocktail of protease and phosphatase inhibitors [300 mmol/L, 4-(2-aminoethyl) benzenesulfonyl fluoride, 0.2 mmol/L ethylenediaminetetraacetic acid, 250 μg/ml leupeptin, 250 μg/ml pepstatin A, 10 mmol/L apro- tinin, 10 mmol/L NaV, and 10 mmol/L NaF]. Lysates were passed through a 19- and 21-gauge needle and then through a Qiashredder that was centrifuged for 2 minute at 13,000 rpm. Whole cell extracts were transferred to fresh Eppendorf tubes, and their protein content was determined using the Bradford DC assay kit (Bio-Rad). A 96-well ELISA plate was coated with capture antibody; rat monoclonal anti-mouse TNF-α (Biosource) at 4 μg/ml in diluted in PBS and incubated overnight at 4°C. Wells were washed three times in wash buffer (0.5%A Tween in PBS), then incubated at room temperature for 1 hour with blocking buffer (1% fetal calf serum, 5% sucrose in PBS). Wells were washed as before and 100 μl of standard (0 to 250 ng/ml of recombinant mouse TNF-α, Cambridge Bioscience) or sample (10 to 25 mg whole cell or whole liver extract) was added to each well and incubated at room temperature for 2 hours. Wells were washed as before and 100 μl of biotinylated rabbit anti-mouse TNF-α polyclonal antibody (Peprotech EC Ltd.) at 100 ng/ml was added and incubated at room temperature for 2 hours followed by further washes and incubation at room temperature for 20 minutes with 0.2 μg/ml of streptavidin-horseradish peroxidase (Biosource). After final washing, 100 μl of substrate solution (1:1 mix H2O2 and 40 μmol/L tetramethylbenzidine diluted in 0.1 mol/L sodium citrate buffer) was added to each well and incubated for 20 minutes at room temperature. Fifty μl of stop solution (1 mol/L sulfuric acid) and was then added to each well and the optical density of reactions was measured at 405 nm and the amount of TNF-α in samples was determined using the standard curve.

**Results**

**Severe Injury-Induced Hepatic Inflammation in Nfkβ1−/− Mice**

A histological comparison revealed mild to severe spontaneous inflammatory changes in the livers of nfkβ1−/− mice that were absent in nfkβ1+/− mice (Figure 1 and Table 1). However these changes were not progressive or associated with a disease state because the severity of inflammation was not altered with aging (comparing 12-week-old to 1-year-old mice) and we were unable to detect any signs of fibrosis. The discovery of a reproducible inflammatory phenotype in the liver of nfkβ1−/− mice suggested that the animals may be susceptible to a more severe response to chronic injury of the liver. To test this idea we compared the effects of chronic chemical injury of the liver between nfkβ1−/− and nfkβ1+/+ mice. Chronic injury of the rodent liver by repeated administration of CCl₄ causes an inflammation-driven wound-healing response and fibrosis.15 Cessation of injury leads to spontaneous recovery and resolution of fibrosis.15,16 The responses of nfkβ1−/− and nfkβ1+/+ mice to CCl₄ injury throughout 12 weeks and their subsequent spontaneous recovery were evaluated. Firstly, we confirmed a differential expression of p50 between nfkβ1−/− and nfkβ1+/+ livers and also determined the expression of RelB and p52 because these factors can have overlapping functions with p50.4,11 Both p50 and RelB transcripts were expressed in wild-type sham (olive oil)-injured animals, however p52 mRNA was undetectable (Figure 2). As expected p50 transcript was not detected in nfkβ1−/− livers, whereas levels of RelB and absence of p52 was similar to wild type. Interestingly, although injury and recovery had no effect on expression of p50, a dramatic loss of RelB expression and weak induction of p52 was observed in both nfkβ1−/− and nfkβ1+/+ livers. In the case of RelB, expression was replenished during recovery (days 5 and 7) while there appeared to be a consistent but low-level induction of p52. The main conclusion from these observations is that CCl₄-injured nfkβ1−/− livers lack expression of both p50 and RelB which have been described to have compensatory regulatory functions in inflammatory responses.11

We next compared histological parameters of liver injury and disease. The most obvious and dramatic difference was a severe periportal/venular inflammatory infiltrate present in the livers of injured nfkβ1−/− mice that was not seen in injured nfkβ1+/+ mice (Figure 3A). Histological analysis showed that the cellularity of the infiltrate in nfkβ1−/− livers was mainly neutrophilic. By count-
ing numbers of stained neutrophils we found that at peak injury (day 1) there were a total of 40 (± 11 SEM) neutrophils/field in nfkb1−/− livers compared with 7 (± 1 SEM) neutrophils/field in nfkb1+/+ livers (Figure 3B). Although numbers of recruited neutrophils dropped by day 3 after injury in both nfkb1+/+ and nfkb1−/− livers there remained an elevated neutrophil count in the latter until at

Table 1. Stable Inflammatory Changes Associated with Lack of nfkb1

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<th>Mouse genotype</th>
<th>Inflammatory change 12 weeks</th>
<th>Inflammatory change 1 year</th>
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<tr>
<td>nfkb1−/− 1</td>
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Five (1 to 5) nfkb1−/− and five (1 to 5) nfkb1+/+ mice were aged for 1 year and liver pathology (normal, minor, or major inflammatory changes) compared with 12-week-old mice.

Figure 1. Stable inflammatory changes in nfkb1−/− liver. H&E staining of liver sections from 12-week-old and 1-year-old nfkb1+/+ and nfkb1−/− mice (numbering refers to precise animal from which liver section was derived as given in Table 1). Black arrows denote minor or major inflammatory changes in nfkb1−/− liver. Photomicrographs are representative of five nfkb1+/+ and four 12-week-old or five 1-year-old nfkb1−/− mice. Original magnifications, ×200.

Figure 2. Expression of hepatic p50, RelB, and p52 during injury and recovery. RT-PCR for p50, RelB, and p52 transcripts was performed using RNA isolated from nfkb1+/+ and nfkb1−/− livers at days 1, 5, and 7 of recovery after 12 weeks of carbon tetrachloride injury or olive oil (sham). The gels shown are at least two experimental repeats from a minimum of three animals/treatment group/genotype. A 1-kb DNA ladder (not shown) was run alongside the PCR products to confirm correct sizes of the amplified cDNA fragments.
least day 14 after injury. By contrast, numbers of hepatic T cells and macrophages were not significantly different between the two genotypes (data not shown). These data indicate that a severe and persistent neutrophilic inflammatory response to injury occurs in the absence of nfkb1<sup>+/−</sup>. In addition, we observed higher numbers of ballooned hepatocytes in nfkb1<sup>+/−</sup>/H9260<sub>b1</sub>/H11002<sub>/H11002</sub>-injured livers indicative of increased hepatocyte damage (Figure 3A). Since a previous study reported no significant differences in the acute response to CCl<sub>4</sub>-induced liver injury, the greater degree of hepatocyte damage in nfkb1<sup>+/−</sup> livers cannot simply be explained by inherent differences in CCl<sub>4</sub> metabolism or the mechanism of injury between nfkb1<sup>+/−</sup> and nfkb1<sup>+/+</sup> animals.

**Figure 3.** Severe inflammation in chronically injured nfkb1<sup>+/−</sup> livers. A: H&E (top left and right, yellow arrows denote ballooned hepatocytes) and neutrophil immunostaining (bottom left and right, red arrows denote neutrophils) staining on nfkb1<sup>+/−</sup> and nfkb1<sup>+/+</sup> liver sections at day 1 of recovery after 12 weeks of CCl<sub>4</sub> injury. Photomicrographs are representative of five nfkb1<sup>+/−</sup> and four nfkb1<sup>+/+</sup> mice. B: Numbers of neutrophils were counted in 15 high-power fields and expressed as average number of neutrophils/high-power field ± SEM. Day 1, P = 0.0016; day 3, P = 0.0185; day 5, not significant, day 7, P = 0.0039; and day 14, P = 0.0117; P values calculated using a paired t-test. Original magnifications, ×100.

**Greater Numbers of Activated Hepatic Stellate Cells and Fibrosis in Injured Nfkb1<sup>−/−</sup> Mice**

CCl<sub>4</sub>-induced chronic liver injury leads to a fibrogenic response characterized by trans-differentiation (or activation) of quiescent hepatic stellate cells (HSCs) into α-SMA-posi-
tive, proliferating myofibroblast-like cells. The increased synthesis and secretion of collagen I/III and tissue inhibitor of metalloproteinase 1 (TIMP-1) by these cells leads to a net deposition of fibril-forming collagens and eventually liver fibrosis and cirrhosis. Examination of the extent of fibrosis in \( \text{nfb1}^{-/-} \) and \( \text{nfb1}^{+/+} \) livers of 12-week \( \text{CCl}_4 \)-injured mice was performed by histopathological analysis and grading of liver sections stained with Sirius Red that stains collagen I/III containing fibers (Figure 4A). This analysis showed a significantly worse fibrotic response in \( \text{nfb1}^{-/-} \) mice evident at both days 1 and 7 after injury (Figure 4, A and B). Underlying the more severe fibrosis in \( \text{nfb1}^{-/-} \) mice was elevated hepatic expression of transcripts for procollagen \( \alpha(1)(I) \) and TIMP-1 (Figure 4, C and D). Furthermore, numbers of profibrogenic \( \alpha \)-SMA-positive cells were dramatically elevated in \( \text{nfb1}^{-/-} \) livers both at peak injury and throughout the recovery process consistent with increased fibrosis in these animals (Figure 5, A and B). In support of these findings, we observed a trend toward elevated levels of transcript for \( \alpha \)-SMA in \( \text{nfb1}^{-/-} \) livers at peak injury and up to day 7 of recovery although this effect did not reach statistical significance (Figure 5C).

**Selective Elevated Expression of TNF-\( \alpha \) by Activated \( \text{nfb1}^{-/-} \) HSCs**

The more pronounced inflammatory and fibrogenic responses of \( \text{nfb1}^{-/-} \) mice to chronic injury raised the possibility that the phenotype of HSCs may be attenuated in these animals. HSCs were therefore isolated from uninjured mouse livers and then activated to a myofibroblast-like phenotype by culturing on plastic, this being a widely accepted and used model of HSC activation. Expression of the p75 low-affinity nerve growth factor receptor and high levels of GFAP expression in the cultured myofibroblasts confirmed that the cells originally isolated were of HSC origin. We did not observe any significant differences in the rate of culture growth or expression of classic markers (\( \alpha \)-SMA and desmin) of activation between \( \text{nfb1}^{-/-} \) and \( \text{nfb1}^{+/+} \) HSCs (Figure 6A). A weak, although not statistically significant, increase in procollagen \( \alpha(1)(I) \) mRNA expression was observed in \( \text{nfb1}^{-/-} \) HSCs (Figure 6B). HSC activation is accompanied by reprogramming of their constitutive NF-\( \kappa \)-B activity to persistently elevated levels by a mechanism involving CBF-1-mediated repression of IkB-\( \alpha \) that results in the induction of NF-\( \kappa \)-B-dependent genes such as ICAM-1 and IL-6. As shown in Figure 7A, the

**Figure 4.** Liver fibrosis is worse in \( \text{nfb1}^{-/-} \) mice. A: Sirius red staining of liver sections at days 1 and 7 of recovery after 12 weeks of \( \text{CCl}_4 \) injury. Black arrows denote collagen fibers and are representative of five \( \text{nfb1}^{+/+} \) and four \( \text{nfb1}^{-/-} \) mice. B: Fibrosis scores expressed as average grade \( \pm \) SEM of five \( \text{nfb1}^{+/+} \) and four \( \text{nfb1}^{-/-} \) mice at days 1 and 7 of recovery. C: TaqMan analysis of procollagen 1 mRNA measured in 20 ng of cDNA from whole liver extracts at days 1, 3, 5, 7, and 14 after 12 weeks of \( \text{CCl}_4 \) injury. The relative level of transcriptional difference was calculated and expressed as an average \( \pm \) SEM from four \( \text{nfb1}^{-/-} \) and five \( \text{nfb1}^{+/+} \) animals. Average cycle numbers were: day 1, 24.60 and 27.21; day 3, 25.63 and 27.32; day 5, 26.53 and 27.10; day 7, 27.0 and 27.87; and day 14, 31.87 and 29.88 for \( \text{nfb1}^{-/-} \) and \( \text{nfb1}^{+/+} \), respectively. The differences were statistically significant, \( P = 0.0329 \), and were calculated using a paired \( t \)-test. D: TaqMan analysis of TIMP1 mRNA measured in 20 ng of cDNA from whole liver extracts. The relative level of transcriptional difference was calculated and expressed as an average \( \pm \) SEM from four \( \text{nfb1}^{-/-} \) and five \( \text{nfb1}^{+/+} \) animals. Average cycle numbers were: day 1, 31.42 and 33.47; day 3, 34.09 and 35.15; day 5, 34.19 and 35.14; day 7, 34.27 and 34.23; and day 14, 36.68 and 36.09 for \( \text{nfb1}^{-/-} \) and \( \text{nfb1}^{+/+} \), respectively. The differences were statistically significant, \( P = 0.0302 \), and were calculated using a paired \( t \)-test. Original magnifications, \( \times 100 \).
expression of ICAM-1 and IL-6 was similar between nfkb1−/− and nfkb1+/+ HSCs indicating no general attenuation of NF-κB-directed gene expression. By contrast, levels of transcript for TNF-α were dramatically higher in nfkb1−/− HSCs. To confirm that nfkb1−/− HSCs produced elevated levels of TNF-α we additionally showed by ELISA that these cells produce 10-fold higher amounts of the cytokine than nfkb1+/+ HSCs (Figure 7B). If this elevated production of TNF-α by HSCs results in a general increase in hepatic expression of the cytokine it would be of relevance to the pathology observed in nfkb1−/− mice. We therefore compared TNF-α expression and found both the level of mRNA (Figure 8A) and protein (Figure 8B) to be elevated in injured nfkb1−/− compared to nfkb1+/+ livers. As shown in Figure 8C, immunohistochemical analysis for TNF-α in nfkb1−/− livers demonstrated that expression was mainly located in cells surrounding the inflammatory infiltrate and was of a similar pattern to that for α-SMA staining (Figure 5A). Dual staining for α-SMA and TNF-α confirmed in situ expression of TNF-α by nfkb1−/− myofibroblasts (Figure 8D). This supports the conclusion of the in vitro studies that nfkb1−/− hepatic myofibroblasts are one source of the elevated levels of TNF-α. These are highly relevant findings because TNF-α is proinflammatory, functions as a recruitment factor for neutrophils to the liver, and has hepatotoxic properties.23,24

Repression of TNF-α Gene Transcription by p50 and HDAC1

We next investigated the mechanism by which absence of the nfkb1 gene leads to elevation of TNF-α expression by HSC-derived myofibroblasts and wanted to confirm that absence of p50 is mechanistically the critical factor. A murine TNF-α promoter-luciferase construct (pTNF-α-Luc) was transfected into nfkb1−/− and nfkb1+/+ activated HSCs either alone or together with an expression vector for p50. The TNF-α promoter was 10-fold more active in nfkb1−/− than nfkb1+/+ HSCs (Figure 9A) and activity of the promoter in both cell types was suppressed by co-transfection of p50 (Figure 9B). Hence, p50 functions in wild-type cells to suppress TNF-α gene transcription, we therefore examined the mechanism underlying this function. It has recently emerged that homodimeric p50 can associate with the transcriptional co-repressor histone deacetylase 1 (HDAC1) and that this complex is stable when p50 homodimers bind to NFκB DNA-binding sites.25 It is proposed that recruitment of p50-HDAC1 complexes to NFκB-responsive genes results in deacetylation of histones and modifies local chromatin structure into a condensed and transcriptionally repressed state.25 To determine whether there is a role for deacetylases in p50-mediated repression of TNF-α promoter activity we initially treated pTNF-α-Luc-transfected nfkb1−/− and nfkb1+/+ HSCs with the broad specific HDAC inhibitor TSA. Figure 10A shows that TSA stimulated TNF-α promoter activity by threefold in nfkb1+/+ HSCs but had no effect on activity of the TNF-α promoter in nfkb1−/− HSCs. However, nfkb1−/− HSCs transfected with a p50 expression vector responded to TSA treatment with a 10-fold stimulation of TNF-α promoter activity. Ability of the TNF-α promoter to respond to TSA therefore critically depends on expression of p50, this observation implicates p50 as an essential component of HDAC-mediated repression of TNF-α expression. We next investigated a specific role for HDAC1. Firstly we showed that both nfkb1−/− and nfkb1+/+ HSCs express HDAC1 (Figure 10B), hence the differences in TSA responses for the TNF-α promoter between the two genotypes cannot be simply explained by differences in HDAC1 expression.
Overexpression of HDAC1 in nfkβ1−/− and nfkβ1+/+ HSCs repressed TNF-α promoter activity, by contrast this effect was not observed in nfkβ1−/− HSCs (Figure 10C). However, transfection of p50 into nfkβ1−/− HSCs rescued the ability of exogenous HDAC1 to repress TNF-α promoter activity. We therefore conclude that HDAC1 is a repressor of TNF-α gene transcription that requires p50 to perform this function. We suggest that in wild-type cells p50 homodimers act to recruit HDAC1 to the TNF-α promoter and that absence of p50 in nfkβ1−/− cells disables the repressive function of HDAC1 resulting in aberrant overexpression of TNF-α. This provides a molecular mechanism to explain the elevated expression of TNF-α and severe neutrophilic inflammation in injured nfkβ1−/− livers.

Figure 6. Lack of differences in markers of activation between nfkβ1−/− and nfkβ1+/+ HSCs. A: Whole cell extracts were isolated from culture-activated HSCs and 25 μg were used to immunoblot for α-SMA, desmin, p75, glial fibrillary acid protein (GFAP), and β-actin. All gels were representative of at least two independent experiments. B: TaqMan analysis of procollagen I mRNA measured in 20 ng of cDNA from activated HSCs. The relative level of transcriptional difference was calculated and expressed as an average ± SEM from three independent cell preparations. Average cycle numbers were 18.45 and 20.55 for collagen and 18.68 and 20.45 for 18s RNA in nfkβ1−/− and nfkβ1+/+, respectively. Use of paired t-test revealed difference not to be significant (P = 0.4).

Figure 7. Selective overexpression of TNF-α by nfkβ1−/− HSCs. A: Three nfkβ1−/− and three nfkβ1+/+ mice were used to generate six independent lines of activated HSCs (1 to 3 and 1 to 3). Total RNA was isolated from all six lines and used as a template in RT-PCR reactions for detection of ICAM-1, IL-6, TNF-α, and β-actin transcripts using protocols described in the Materials and Methods. The gels shown are presented as three sets of genotype pairs and are representative of two repeat RT-PCRs for each RNA sample. A 1-kb DNA ladder was run alongside the PCR products to confirm correct sizes of the amplified cDNA fragments (not shown). B: Whole cell extracts were isolated from nfkβ1−/− and nfkβ1+/+ activated HSCs and sandwich ELISA was used to quantify TNF-α levels as average ng of TNF-α/μg whole cell extract ± SEM, n = 3 nfkβ1−/− and nfkβ1+/+-independent isolations.
**Figure 8.** Elevated TNF-α in chronic injured nfkbia/− liver. A: Total RNA was isolated from the livers of three nfkbia/− and three nfkbia/+ mice at day 1 after 12 weeks after injury and were used as templates in RT-PCR reactions for TNF-α and β-actin transcripts. The gels show RT-PCR products for the three livers/genotype and were representative of two repeat RT-PCRs for each RNA sample. B: Whole liver protein extracts were at day 1 after injury and sandwich ELISA was used to quantify TNF-α (as average ng of TNF-α/μg whole liver protein ± SEM, n = 4 nfkbia/− and 5 nfkbia/+). C: TNF-α immunostaining on liver sections at day 1 after injury, arrows denote TNF-α-expressing cells mainly localized to regions surrounding inflammatory infiltrates in nfkbia/− livers. Photomicrographs are representative of four nfkbia/− and five nfkbia/+ animals. D: TNF-α and α-SMA dual immunostaining on nfkbia/− liver sections at day 1 after injury, yellow arrows denote TNF-α and α-SMA co-localization. Original magnifications: ×100 (C, top), ×200 (C, bottom), ×1000 (D).

**Figure 9.** TNF-α promoter activity is increased in nfkbia/− HSCs. A: Activated HSCs isolated from nfkbia/− and nfkbia/+ mice were transfected with 10 ng of pRLTK (Renilla control) and 1 μg of pTNF-α-Luc. After 48 hours the luciferase activities were determined, normalized to pRLTK, and promoter activity in nfkbia/− cells expressed as promoter activity relative to the activity measured in nfkbia/+ cells (mean ± SE of triplicate transfections) B: HSCs were transfected with 10 ng of pRLTK, 1 μg of pTNF-α-luc, and 2 μg of pCDNA3-p50 (+p50) or pCDNA3 (−). After 48 hours, luciferase activities were determined, normalized to pRLTK activity, and expressed as promoter activity relative to the activity measured in pCDNA3-transfected cells (mean ± SE of triplicate transfections).
Discussion

Now that there is unequivocal and direct evidence that NF-κB drives inflammation,26 there is a need to understand the specific physiological role played by each individual Rel protein in the inflammatory response. In addition, the emerging function of transcriptional co-activators and co-repressors as regulators of Rel activities27 necessitate studies that determine how these factors interplay with each Rel protein to achieve appropriate physiological responses of tissues to stress, injury, and infections. The data presented in this study define a protective role for p50 that operates to limit the inflammatory and fibrogenic responses in the chronically injured liver. We further define a transcriptional mechanism to explain this anti-inflammatory property of p50, which involves HDAC-dependent repression of TNF-α gene transcription.

The nfkb1−/− mouse has previously been used in attempts to determine a function for p50 in acute liver injury and regeneration, however these studies concluded a lack of a significant role in either process.12,13 The fundamental difference between these earlier studies and the work presented here is our use of a model that addresses the potential for p50 to influence the physiological response to iterative injury. The chronic CCl4-injury model generates repeated and overlapping rounds of injury, inflammation, and wound-healing processes that mimic the pathological events underlying the development of fibrosis of the chronically diseased liver. Using this model we show that absence of nfkb1 is associated with more profound hepatic inflammation and development of a more severe fibrogenic response. Of note nfkb1−/− mice are also susceptible to more severe lung inflammation and damage when challenged by pulmonary infection with Escherichia coli.28 Taken together these observations indicate that p50 is an important negative regulator of tissue inflammation in multiple organs responding to injury and infections. It was therefore of interest to determine a mechanism through which p50 may exert its protective anti-inflammatory mode of action.

Figure 10. HDAC1-mediated repression of TNF-α transcription is p50-dependent. A (top): Activated HSCs were transfected with 10 ng of pRLTK and 1 μg of pTNF-α-Luc for 24 hours before 24 hours of treatment with or without 500 nmol/L TSA. Luciferase activities were determined, normalized to pRLTK activity, and expressed as promoter activities relative to activity measured in cells lacking TSA treatment. A (bottom): As above but with the inclusion of pCDNA-p50 in the transfection mixture. All transfections were performed in triplicate. B (top): Total RNA was isolated from nfkb1−/− and nfkb1+/+ HSCs and used as a template for RT-PCR detection of HDAC1. B (bottom): Whole cell extracts were isolated from nfkb1−/− and nfkb1+/+ activated HSCs and 25 μg of protein was used to immunoblot for HDAC1 and β-actin. C: Activated HSCs were transfected with 10 ng of pRLTK and 0.5 μg of pTNF-α-Luc alone (−) or together with 1 μg of pCDNA1 and 2 μg of pHDAC1 (HDAC1) or 1 μg of pCDNA-p50 and 2 μg of pHDAC1 (p50+HDAC1). After 48 hours, luciferase activities were determined, normalized to pRLTK activity, and expressed as promoter activities relative to activity in cells transfected with pTNF-α-Luc alone (mean ± SE of triplicate transfections).
The most dramatic difference between CCl₄-injured nfkb1⁻/⁻ and nfkb1⁺/+ mice was the massive accumulation of neutrophils in perportal/venular regions of nfkb1⁻/⁻ livers. Neutrophils accumulate in and adhere to sinusoids and postsinusoidal venules during hepatic inflammation.²⁹ Neutrophils have been strongly implicated in the development of liver damage and HSC activation via their production of reactive oxygen species and various hydrolytic enzymes.²⁴,²⁹,³⁰ Disease states in which neutrophil-induced liver damage is associated include hepatic ischemic-reperfusion, alcoholic hepatitis, and drug-induced liver injuries.³¹ Again of relevance to our findings the inflammation generated in the lungs of E. coli-infected nfkb1⁻/⁻ mice mainly consisted of neutrophils suggesting that absence of p50 renders tissues more susceptible to neutrophil infiltration.²⁸ One of the most potent mediators of the hepatic accumulation and activation of neutrophils is TNF-α.³² The livers of CCl₄-injured nfkb1⁻/⁻ mice expressed higher levels of TNF-α than injured nfkb1⁺/+ mice and immunohistochemical analysis showed that TNF-α expression was mainly associated with α-SMA-positive cells surrounding inflammatory infiltrates. There are currently believed to be at least two main sources of α-SMA-positive myofibroblasts in the injured liver, these being the perportal myofibroblast and the activated HSCs.¹⁸ Importantly, both cell types have a similar profibrogenic and proinflammatory phenotype and if their persistence in the chronically injured liver leads to progressive deposition of cross-linked collagen and ultimately hepatic fibrosis. By isolating HSCs and subsequently activating them in culture we were able to show that although there were no gross phenotypic differences between nfkb1⁻/⁻ and nfkb1⁺/+ HSCs, the former cell type produced highly elevated levels of TNF-α. There have been no previous reports of HSCs expressing TNF-α and we measured only low barely detectable levels of the cytokine in nfkb1⁺/+ HSCs. We therefore propose that aberrant production of TNF-α by myofibroblasts probably of both perportal and perisinusoidal (HSCs) origin provides an explanation for the severe neutrophilic inflammatory infiltrate observed in injured nfkb1⁻/⁻ mice.

There is both clinical and experimental evidence in support for a pathophysiological role for TNF-α in liver fibrosis. Elevated serum TNF-α concentrations in patients with alcoholic hepatitis have been reported by several groups and the values correlate with severity of cirrhosis and mortality.³³ Similarly, there is a strong correlation between level of serum TNF-α and development of periportal fibrosis in human Schistosoma mansoni infection, with higher concentrations of TNF-α being associated with a higher risk of fibrosis.³⁴ Liver fibrosis resulting from chronic CCl₄ exposure is almost completely absent by histological analysis in TNF receptor-deficient mice and is accompanied by marked reductions in hepatic collagen and transforming growth factor-β expression.³⁵ Furthermore, neutralizing antibodies to TNF-α prevent the development of rodent liver fibrosis induced by concanavalin A.³⁶ However, it is not yet clear as to whether TNF-α has a direct influence on the fibrogenic process and given its highly pleiotropic function in the liver it is more likely that it exerts its influence indirectly via enhancing the inflammatory response to liver damage.

TNF-α expression is tightly regulated at the transcriptional level and is under the repressive influence of two NF-κB proteins, p50 and RelB.⁵⁻⁷ Knockout mouse studies show that p50 and RelB have overlapping compensatory functions with respect to inflammation.¹¹ Although RelB⁻/⁻ mice spontaneously develop a mild inflammatory infiltrate in several organs, a combined deficiency of p50 and RelB leads to a severe multiorgan inflammatory phenotype involving recruitment of T cells, macrophages, and neutrophils. Of note in the current study we observed a loss of hepatic RelB at peak injury that was replenished on recovery, the physiological significance of this injury-induced attenuation of RelB is currently not understood. However, it is significant that peak liver injury in nfkb1⁻/⁻ mice was effectively associated with a combined deficiency of p50 and RelB. RelB⁻/⁻ fibroblasts overexpress TNF-α in response to stimulation with LPS although the precise mechanism by which RelB negatively regulates TNF-α gene transcription remains poorly defined.⁶ Seven The repressive function of p50 on TNF-α is also incompletely described although it is known that the distal region of both the human and murine TNF-α promoters carry κB sites that preferentially associate with p50 homodimers.⁵,³⁷ Mouse macrophages have been shown to secrete a TNF-α-inhibitory factor that stimulates their enhanced expression of p50 and repression of TNF-α promoter activity.⁷ This effect and the ability of transfected p50 to repress TNF-α promoter activity was lost if three upstream κB sites with preference for p50 were removed from the promoter. Our study not only provides a physiological explanation for the requirement for tight inhibitory regulation of the TNF-α locus by p50 (ie, the protection of tissues against excessive inflammation and damage resulting from overexpression of TNF-α) but also describes how p50 mediates this repression. The murine TNF-α promoter not only displayed highly elevated activity in nfkb1⁻/⁻ HSCs, it was also unresponsive to the stimulatory effects of TSA and the repressive effects of HDAC1. Importantly we demonstrated that ectopic expression of p50 in nfkb1⁻/⁻ HSCs repressed activity of the TNF-α promoter and restored the sensitivity to TSA and HDAC1 observed in wild-type HSCs. Overexpression of TNF-α by nfkb1⁻/⁻ HSCs is therefore because of absence of p50 (rather than a function attributed to the precursor protein p105) and specifically arises as a consequence of the loss of p50-dependent HDAC-mediated transcriptional repression of the TNF-α gene.

Zhong and colleagues²⁵ described how a complex of homodimeric p50 and HDAC1 binds to κB sites and functions as a general repressor of NF-κB-dependent transcription in resting cells. On cell stimulation, the phosphorylated p65 subunit of newly formed nuclear p50/p65 heterodimers associates with CBP/p300 and this complex then replaces the repressive p50/p50-HDAC1 complex at κB sites to activate transcription. Interestingly we did not observe a general difference in the expression of NF-κB-responsive genes between nfkb1⁻/⁻ and nfkb1⁺/+ HSCs, including IL-6 that Zhong and colleagues²⁵ showed to be a target for p50/p50-HDAC1-
mediated repression in resting cells. The most obvious explanation for this difference between the two studies is that the HSC-derived myofibroblasts used in the present study are in an activated state in which the majority of NF-κB-responsive genes will be under the influence of p50/p65-CBP complexes. This idea is supported by the fact that p50/p65 dimers are the major κB-binding proteins detected by electrophoretic mobility shift assay in activated HSCs.20–22 However, genes such as TNF-α that are under the control of κB sites with high-binding affinity for p50 homodimers will preferentially associate with p50/p50-HDAC1 complexes over p50/p65-CBP complexes. This is supported by experimental evidence that TNF-α transcription is highly sensitive to the repressive effects of p50 even when in the presence of high levels of p65.5 Our findings therefore complement and advance those of Zhong and colleagues23 by showing that p50/p50-HDAC1 complexes remain functional in the stimulated cell to enable the selective repression of a subset of NF-κB-responsive genes. Additional and possibly signal- or cell-dependent regulatory mechanisms are presumably then required to overcome p50/p50-Bcl3 and p50/p50-HDAC1 repression for appropriate induction of TNF-α. The severity of tissue inflammation and damage observed in the livers of the CCl4-injured mice serves to illustrate how this complexity of transcriptional regulation operates to prevent the pathological consequences of inappropriate overexpression of the cytokine. An intriguing observation was that treatment of p50-transfected nfkb1−/− HSCs with TSA resulted in the powerful activation of the TNF-α promoter indicating that the deacetylase inhibitor converts p50 from a repressor to an activator of TNF-α transcription (Figure 10A). One way in which p50/p50 dimers can function as transcriptional activators is by associating with Bcl3 that itself binds to CBP/p300 and can interact with components of the basal transcription machinery.37 HSC activation is accompanied by induction of Bcl3.21 This raises the possibility that p50/p50-Bcl3 and p50/p50-HDAC1 complexes may co-exist in HSCs and compete for κB sites or alternatively that association of Bcl3 and HDAC1 with p50 may be mutually exclusive events. Regulation of these interactions will help dictate the level of TNF-α-driven inflammation in infected and injured tissues.

In summary, the p50 subunit of NF-κB is protective in the chronically injured liver and we have described a plausible mechanism to explain this effect involving repression of TNF-α expression by p50/p50-HDAC1 complexes. Further studies on the function of p50 in inflammatory states and the role played by its co-activators and co-repressors may lead to new anti-inflammatory drugs that specifically target p50 activities.

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References

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