Glucocorticoid receptors are downregulated in hepatic T lymphocytes in rats with experimental cholangitis

K Tjandra, T Le, M G Swain

Background and aims: Primary sclerosing cholangitis is a Th1 cytokine driven disease with a poor clinical responsiveness to glucocorticoid therapy. We have previously documented elevated circulating glucocorticoid levels in cholestatic rats and in addition have noted increased hepatic expression of the Th1 cytokine interferon-γ (IFN-γ) in a rat model of cholangitis. Therefore, we examined the relationship between circulating glucocorticoid levels, hepatic IFN-γ expression, and hepatic T cell glucocorticoid receptor (GR) expression in a rat model of cholangitis to provide insight into the possible mechanism underlying hepatic T cell glucocorticoid resistance in cholangitis disease.

Methods: Cholangitis was induced in male Sprague-Dawley rats by oral administration of low dose α-naphthylisothiocyanate (ANIT). On day 14, ANIT fed and control rats were sacrificed, serum collected, and hepatic, splenic, and peripheral blood T lymphocytes isolated for GR expression, as determined by reverse transcription-polymerase chain reaction and western blotting.

Results: Circulating glucocorticoid levels were markedly elevated in ANIT fed rats. Hepatic T lymphocyte GR mRNA and protein levels were significantly reduced in ANIT treated rats compared with controls. In contrast, GR mRNA and protein expression in splenic and circulating T lymphocytes was similar in both groups. Furthermore, reduced hepatic T cell GR expression in ANIT fed rats was associated with reduced hepatic CD4+ T cell sensitivity to dexamethasone inhibitory effects (that is, inhibition of interleukin 2 receptor expression).

Conclusion: We conclude that hepatic T lymphocyte resistance to elevated endogenous glucocorticoid levels in rats with experimental cholangitis appears, in part, to be mediated by decreased GR expression.

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Material and methods

Animal model of chronic cholangitis

Experimental cholangitis was induced in male Sprague-Dawley rats (200–250 g; Charles River, St Constant, Quebec, Canada) by chronic oral administration of low dose α-naphthylisothiocyanate (ANIT 1 mg/kg powdered rat chow; Sigma Chemicals, St Louis, Missouri, USA) as previously described. Both control and ANIT treated rats were kept in a light controlled room (12 hour light/12 hour dark cycles) and treated in accordance with the Canadian Council of Animal Care guidelines. Rats were handled twice daily to avoid immune mediated diseases. It is not clear however whether the poor response to steroid treatment in these patients is due specifically to glucocorticoid resistance.

The objectives of this study were to determine whether circulating endogenous glucocorticoid levels are increased in rats with experimental cholangitis, and if they are, to determine whether the development of a hepatic Th1 cytokine profile is due to changes in hepatic T lymphocytes in the setting of cholangitis which may render them less responsive to glucocorticoids.

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Abbreviations: PSC, primary sclerosing cholangitis; IL, interleukin; IL-2, IL-2 receptor; IFN, interferon; DHEA, dehydroepiandrosterone; IBD, inflammatory bowel disease; GR, glucocorticoid receptor; PBC, primary biliary cirrhosis; ANIT, α-naphthylisothiocyanate; PBS, phosphate buffered saline; FITC, fluorescein isothiocyanate; PE, phycoerythrin; RT-PCR, reverse transcription-polymerase chain reaction; TNF, tumour necrosis factor; FACS, fluorescence activated cell sorter; ConA, concanavalin A; NK, natural killer; GAPDH, glyceraldehyde-3-phosphate dehydrogenase; PBMC, peripheral blood mononuclear cells.
handling stress and maintain low baseline serum cortico-
steroid levels. Animals were sacrificed 14 days following ANIT
administration and ANIT fed rats for 14 days according to Sarlis
and colleagues.\textsuperscript{17} Rat truncal blood was collected in
the morning (AM: between 8 and 9 am) and in the evening 
(PM: between 7 and 8 pm).

**Serum corticosterone and DHEA levels**
Serum corticosterone (the major adrenal glucocorticoid found
in rodents) and DHEA levels of control and ANIT treated rats
were determined using commercially available radioimmu-
noassays (ICN Biomedicals, Inc. Costa Mesa, California, USA,
and Diagnostic System Laboratories, Inc., Webster, Texas, USA,
respectively).

**Hepatic cytokine levels**
Livers of control and ANIT treated rats were perfused with ice
cold phosphate buffered saline (PBS; pH 7.4) and extracted for
cytokine protein expression. Experiments were also performed to
determine serum corticosterone kinetics: (i) serum corticosterone
levels were also measured four and seven days following ANIT
administration and (ii) corticosterone diurnal variations were
examined in control rats and rats treated with ANIT for 14
days according to Sarlis and colleagues.\textsuperscript{17} Rat truncal blood
was collected in the morning (AM: between 8 and 9 am) and
in the evening (PM: between 7 and 8 pm).

**Figure 1** Serum corticosterone and dehydroepiandrosterone
(DHEA) levels in control and \(\alpha\)-naphthylisothiocyanate (ANIT)
treated rats. Data are mean (SEM) of at least four animals per group.
(A) Corticosterone levels; (B) DHEA levels; and (C) serum
corticosterone/DHEA ratio. *\(p<0.05\) and **\(p<0.001\) versus
controls.

**Figure 2.** (A) Circulating corticosterone levels in control animals
and rats treated with \(\alpha\)-naphthylisothiocyanate (ANIT) for 4, 7, and
14 days. Data are mean (SEM) of at least four animals per group.
***\(p<0.001\) versus controls; †\(p<0.01\) versus day 4 and 7.
(B) Morning and evening plasma corticosterone levels of control and
ANIT fed rats for 14 days. Data are mean (SEM) of six animals per
control group, and at least 13 rats per ANIT group. **\(p<0.01\),
***\(p<0.001\) versus control AM levels.
mRNA/protein expression and cellular surface markers. T lymphocytes from the spleen and peripheral blood of control and ANIT treated rats were isolated with slightly modified protocols. Spleens were excised, minced, and homogenised in PBS. The spleen homogenate was then filtered through a 30 micron nylon mesh (Small Parts Inc.). Cell suspensions were layered on Lympholyte Rat (Cedarlane, Hornby, Ontario, Canada) and centrifuged at 1400 g for 30 minutes at room temperature. Peripheral blood was collected in sterile tubes containing anticoagulant (0.14 mol/l citric acid, 0.2 mol/l sodium citrate, and 0.22 mol/l dextrose) and centrifuged at 1400 g for 10 minutes. The pellets were resuspended in PBS to twice the original volume collected. Cell suspension was then layered on Lympholyte Mammals (Cedarlane) and centrifuged at 800 g for 30 minutes at room temperature. Splenic and peripheral blood T lymphocytes were collected from the interface layer, washed, and counted. Purified T lymphocytes were either used immediately or stored at −70°C.

**Flow cytometry**

To determine the cellular phenotypes and distribution of isolated liver derived lymphocytes in control and ANIT treated animals, we stained these cells for commonly expressed specific cell surface markers and analysed them by flow cytometry. Briefly, isolated hepatic lymphocytes (10⁶ cells) were incubated for 30 minutes at room temperature with 10 μl of fluorescein isothiocyanate (FITC) conjugated mouse antirat CD4 (clone W3/25; Serotec Inc., Raleigh, New Jersey, USA), mouse antirat CD8 (clone MRC OX-8; Serotec), mouse antirat monocytes/macrophages (clone ED1; Serotec), or mouse antirat B cell leucocyte common antigen (clone MRC OX-33; Cedarlane). Hepatic natural killer (NK) cells were also determined using phycoerythrin (PE) conjugated mouse antirat NKR-P1A cell surface marker (clone 10/78; Pharmingen, Mississauga, Ontario, Canada), which were labelled together with either FITC conjugated anti-CD4 or anti-CD8 monoclonal antibodies (see above). Mouse FITC (Serotec) and PE (Pharmingen) conjugated IgG1 negative isotype controls were also included in the analysis. Cells were washed twice in PBS following incubation, and resuspended in 0.2 ml of 1% formalin in PBS. Fixed cells were stored overnight at 4°C and analysed the next day by flow cytometry (Becton-Dickinson, Mountain View, California, USA). Cells were counted using an electronic gate set on the lymphocyte cluster on the forward and side scatter plot and analysed using CellQuest software (Becton-Dickinson).

**Semiquantitative RT-PCR**

RNA was extracted from purified T lymphocytes isolated from various sources (that is, liver, spleen, and peripheral blood) in Trizol (Molecular Research Center, Inc., Cincinnati, Ohio, USA) according to the manufacturer’s protocol. Briefly, isolated cells were sonicated in 1 ml of ice cold Trizol for 2–3 minutes (Fisher sonic dismembrator model 300; Farmingdale, New York, USA) and RNA extracted by adding 0.2 ml chloroform. Following centrifugation at 12 000 g for 15 minutes at 4°C, the aqueous phase was transferred to an Eppendorf tube and precipitated in 0.5 ml isopropanol. Further centrifugation at 12 000 g for 15 minutes at 4°C resulted in an RNA pellet that was washed with 1 ml of 75% ethanol and centrifuged at 7500 g for five minutes. The pellet was air dried and dissolved in diethyl pyrocarbonate treated water (Research Genetics, Burlington, Ontario, Canada) and stored at −70°C. The final RNA concentration was determined spectrophotometrically using a Gene Quant spectrophotometer (Pharmacia, Piscataway, New Jersey, USA). All reagents were from Sigma, unless otherwise indicated.

Steady state T lymphocyte GR mRNA levels were determined using the “primer dropping” method previously described. Reverse transcription-polymerase chain reaction (RT-PCR) was performed in an Amplitron I Thermal Cycler (Barnsted-Thermolyne, Dubuque, Iowa, USA). Briefly, complementary DNA (cDNA) was generated by an RT reaction by preincubating 2 μg of sample RNA with the appropriate RT reaction mixture at 21°C for 10 minutes, 42°C for 50 minutes, and 95°C for five minutes to terminate the reaction. Multiple

![Figure 3](https://example.com/figure3.png)
PCR reactions were performed using 2 µl of the newly synthesised cDNA as a template and amplified in a 50 µl PCR reaction mixture containing 1×PCR buffer, 80 µM of each deoxynucleotide triphosphates (dATP, dGTP, dCTP, and dTTP), and 20 pmol each of rat GR primers (307 bp; Biognostik, Germany). Each PCR cycle consisted of a heat denaturation step at 94°C for one minute, a primer annealing step at 60°C for 30 seconds, and a polymerisation step at 72°C for one minute. For proper amplification and to ensure that PCR amplification was in the linear range, the number of PCR cycles was predetermined for each sample (liver, 34 cycles; spleen and peripheral blood, 33 cycles). Two units of Taq DNA Polymerase (Pharmacia) were used.

**Figure 4** Glucocorticoid receptor (GR) expression in hepatic T lymphocytes isolated from control animals and rats treated with α-naphthylisothiocyanate (ANIT) for 14 days. Hepatic T lymphocyte GR mRNA levels were determined by reverse transcription-polymerase chain reaction (top panel; n=3-5) and protein expression by western blotting (lower panel; n=4). *p<0.05 and ***p<0.001 versus controls. GAPDH, glyceraldehyde-3-phosphate dehydrogenase.

**Figure 5** Splenic T lymphocyte glucocorticoid receptor (GR) expression in control and rats receiving α-naphthylisothiocyanate (ANIT) for 14 days. Splenic T lymphocyte GR mRNA levels are shown in the top panel (n=9) and protein expression in the lower panel (n=4). Similar splenic T lymphocyte GR mRNA and protein levels were observed between the control rats and those receiving ANIT for 14 days. GAPDH, glyceraldehyde-3-phosphate dehydrogenase.
added during the first denaturation step. Equal aliquots of internal standard glyceraldehyde-3-phosphate dehydrogenase (GAPDH) (222 bp; see Swain and colleagues20) were amplified through 24–26 amplification cycles. Aliquots of PCR reaction products were electrophoresed through 2% agarose gels containing 0.2 µg/ml of ethidium bromide. Gels were visualised under UV light, and analysed by computer densitometric scanning of the images using Quantity One software (Biorad, Hercules, California, USA). All bands were expressed in arbitrary densitometric units relative to the internal standard signal.

**Western blotting**

Isolated T lymphocyte GR protein levels were quantitated using western blotting.21 Briefly, isolated T lymphocytes were incubated in lysis buffer (10 mM Tris, pH 7.4, 150 mM NaCl, 0.5% Triton-X) containing protease inhibitor cocktail (1 mM 4-(2-aminoethyl)benzenesulphonyl fluoride, 0.04 mM bestatin, 0.014 mM E-64 (trans-epoxysuccinyl-L-leucylamido(4-guanidino)butane), 0.022 mM leupeptin, 0.015 mM pepstatin A, 0.8 µM aprotinin, and 1 mM sodium orthovanadate; Sigma). The protein content of each sample was determined using a Biorad protein assay kit with bovine serum albumin standard (Biorad). Protein samples were equilibrated with lysis buffer, mixed with Laemmli sample buffer (Biorad), and boiled for five minutes. Proteins were loaded into wells and separated in 8% polyacrylamide gel containing sodium dodecyl sulphate, which then electrophoretically transferred to nitrocellulose membrane (Biorad). Membrane blots were blocked with 5% skim milk at room temperature for one hour to prevent non-specific binding of antibodies before incubating them with the primary antibody (mouse monoclonal antibody against rat GR: 1/3000; Jackson Immuno Research Laboratories Inc., West Grove, Pennsylvania, USA). Positive immunoreactivity on blots was detected following exposure to a chemiluminescent producing substrate (Pierce, Rockford, Illinois, USA). GR protein band density was quantitated using computer densitometric scanning of the images with Quantity One software (Biorad).

**GR sensitivity assay**

To determine whether reduced hepatic T cell GR number in chronic cholangitis is associated with reduced GR sensitivity to glucocorticoid, surface IL-2 receptor (IL-2R) expression (IL-2R expression is a widely used marker of lymphocyte activation22) on hepatic CD4+ lymphocytes was determined using flow cytometry, as described previously.23 Briefly, liver derived lymphocytes from control and ANIT fed rats were incubated in RPMI 1640 media supplemented with 5% fetal calf serum, 100 U/ml penicillin, 100 µg/ml streptomycin (all from Gibco BRL), and 2 mM L-glutamine (Sigma). Liver derived lymphocytes (4×10⁶ cells/ml) were stimulated in 24 well plates with concanavalin A (ConA 2.5 µg/ml; Sigma) for 24 hours at 37°C and 5% CO₂. Then, dexamethasone (10⁻⁶ M; Sigma) or vehicle was added to the cell culture and the plates were further incubated for an additional 24 hours. This dose of dexamethasone has previously been shown to inhibit IL-2R expression during antigen induced mouse splenic T cell activation.24 Cells were harvested, washed, and double stained with FITC conjugated mouse antirat CD4 (clone W3/25; Serotec) followed by PE conjugated mouse antirat CD25 (IL-2R) monoclonal antibody (clone MRC OX-9; Serotec). FITC and PE conjugated mouse IgG1 negative controls (Serotec) were also included in the analysis.

**Statistics**

All data are expressed as mean (SEM). The Student’s t test was used to compare differences between the two groups. Multiple
comparisons were performed by one way analysis of variance followed by the Student-Newman-Keul post hoc test. A p value of <0.05 was considered significant.

RESULTS
Serum corticosterone and DHEA levels
We observed a significant increase in serum corticosterone levels in ANIT treated rats compared with control animals (fig 1A). Serum corticosterone levels were markedly elevated in ANIT fed rats as early as day 4 post ANIT treatment and the elevations were maintained over the remaining 14 days of ANIT treatment (fig 2A). In contrast, serum DHEA levels of rats treated with ANIT were significantly reduced when compared with control levels (fig 1B). The serum corticosterone to DHEA ratio was significantly higher in ANIT treated rats than in controls (fig 1C). Circulating corticosterone levels in control animals were low in the morning (between 8–9 am) but significantly increased in the evening (7–8 pm) (fig 2B). In contrast, diurnal variation of corticosterone levels were absent in ANIT fed rats; serum corticosterone levels were elevated in the morning and remained high in the evening.

Hepatic cytokine profile
Hepatic IFN-γ levels were significantly elevated in ANIT treated rats compared with controls, but in contrast, no significant difference in hepatic IL-4 levels was noted between control and ANIT treated rats (fig 3A, 3B, respectively). Hepatic IFN-γ/IL-4 ratio was significantly higher in ANIT treated rats than in controls (fig 1C). Circulating corticosterone levels in control animals were low in the morning (between 8–9 am) but significantly increased in the evening (7–8 pm) (fig 2B). In contrast, diurnal variation of corticosterone levels were absent in ANIT fed rats; serum corticosterone levels were elevated in the morning and remained high in the evening.

Liver derived lymphocytes
There was no marked difference in the cellular phenotypes and distribution of purified hepatic T lymphocytes obtained from control and ANIT treated rats. However, the total number of hepatic T lymphocytes extracted from ANIT treated rats was approximately five times higher than from control livers (data not shown). The isolated hepatic lymphocytes mainly consisted of CD4+ liver derived lymphocytes of control rats and rats treated with α-naphthylisothiocyanate (ANIT) for 14 days. Freshly isolated hepatic lymphocytes were stimulated with concanavalin A (Con A) in the presence or absence of 10^-6 M dexamethasone (Dex). Similar results were obtained in two additional experiments.

Figure 7 A representative dot plot of surface expression of interleukin 2 receptors (IL-2R) on CD4+ liver derived lymphocytes of control rats and rats treated with α-naphthylisothiocyanate (ANIT) for 14 days. Freshly isolated hepatic lymphocytes were stimulated with concanavalin A (Con A) in the presence or absence of 10^-6 M dexamethasone (Dex).

T lymphocyte GR expression
Hepatic T lymphocyte GR mRNA expression was significantly reduced in ANIT treated rats compared with control animals (fig 4, top panel). Furthermore, hepatic T lymphocyte GR downregulation in ANIT treated rats was also observed at the protein level (fig 4, bottom panel). To determine whether T lymphocyte GR downregulation in ANIT treated animals is a generalised phenomenon in response to elevated glucocorticoids, we studied GR mRNA and protein expression of extrahepatic T lymphocytes obtained from the spleen and peripheral blood. In contrast with hepatic T lymphocytes, both splenic and peripheral blood T lymphocytes of control and ANIT treated animals showed similar GR mRNA and protein levels (figs 5, 6).

Hepatic T cell IL-2R expression
In vitro activation of hepatic T cells with ConA resulted in upregulation of IL-2R expression in both control and ANIT fed animals. However, dexamethasone reduced IL-2R expression on activated hepatic T cells isolated from control rats, but failed to attenuate IL-2R upregulation in ANIT treated rats (fig 7).
**DISCUSSION**

Glucocorticoids have a wide range of biological effects with potent anti-inflammatory functions, in part through their ability to inhibit the expression of inflammatory cytokines such as IL-2 and IFN-γ. T lymphocytes are typically proinflammatory in nature and play a central role in cell mediated immunity. An augmented, and overzealous Th1 response may be detrimental to tissues, and is known to play an important role in the pathogenesis of several autoimmune diseases. Despite a significant rise in the serum glucocorticoid levels in rats with experimental cholangitis, we observed a predominant hepatic Th1 cytokine profile, which we have previously demonstrated to be due to enhanced IFN-α production by hepatic CD4+ T lymphocytes. Our data suggest that compared with controls, and we suggest that this is a consequence of the immunomodulatory actions of endogenous glucocorticoids. This insensitivity of hepatic T lymphocytes to glucocorticoid mediated suppressive effects would be expected to directly aggravate hepatic inflammation. Interestingly, clinical studies have shown that corticosteroid treatment in patients with chronic cholangitic diseases such as PSC and PBC, either alone or in conjunction with other drugs such as ursodeoxycholic acid, does not appear to be effective.

Resistance of T lymphocytes to glucocorticoid effects has been documented in a number of chronic inflammatory diseases including asthma and IBD. Steroid resistance in these conditions has been attributed, at least in part, to GR defects. In patients with IBD, GR numbers in peripheral blood mononuclear cells (PBMCs) are significantly lower in steroid non-responsive patients compared with steroid responders and healthy subjects. Moreover, IBD patients have been shown to have elevated serum cortisol levels with lower levels of serum DHEA compared with control subjects, a similar finding to our observation in rats with experimental cholangitis. Our results demonstrate that hepatic T lymphocytes isolated from ANIT treated rats have decreased GR expression compared with controls and we suggest that this is a direct consequence of hepatic T lymphocyte glucocorticoid insensitivity of these cells. GR downregulation does not appear to be a consequence of elevated circulating glucocorticoids as similar GR expression was documented in T lymphocytes isolated from the spleen and peripheral blood of control and ANIT treated rats.

GR expression may vary in different tissues or cell populations. Therefore, we ensured that similar cell populations were being compared. Fluorescence activated cell sorter (FACS) analysis of isolated liver derived lymphocytes demonstrated that GR downregulation in hepatic T cells from ANIT treated rats was not due to differences in cell populations between control and ANIT treated rats, as both control and ANIT groups showed comparable cell types and distribution. Liver derived lymphocytes from both control and ANIT fed rats consist mainly of T and NK cells with low or negligible numbers of monocytes/macrophages and B cells. A similar observation was noted in liver derived lymphocytes from patients with PBC and PSC. Interestingly, our double staining study showed that the great majority of NK positive cells in liver derived lymphocytes isolated from control and ANIT treated animals were also CD8 positive, similar to the observations of others. Although NK cells typically do not express CD3, they have been shown to express CD4 or CD8 surface markers. In our study, liver derived lymphocytes in control and ANIT treated rats consisted of a much lower percentage of NK/CD4+ cells than NK/CD8+ cells. Although we did not measure the relative percentage of NK positive cells in the peripheral blood of ANIT fed rats, we did observe a lower percentage of circulating NK positive cells in control rats. Accumulation of NK cells in the liver may play a direct or indirect role in liver immune dysfunction; however, their role in the development of chronic cholangitis is unknown.

Reduced cellular sensitivity to GC mediated suppression may be associated with lowered GR number and/or affinity. However, reduced GR numbers may not always translate into reduced GR function. Schlaghecke et al demonstrated that GR function (that is, GC inhibition of lymphocyte proliferation and cytokine release) is not compromised despite reduced GR number in PBMC of patients with rheumatoid arthritis. To determine whether hepatic T cell GR downregulation in rats with chronic cholangitis was associated with reduced GR function, hepatic T cell ConA stimulated IL-2R expression was analysed in the presence or absence of dexamethasone using FACS analysis. GCs have been shown to inhibit cytokine receptor (for example, IL-2R) expression on activated T cells. In our study, dexamethasone attenuated ConA induced IL-2R expression on hepatic T cells of control rats but failed to inhibit ConA stimulated IL-2R expression on hepatic T cells from ANIT treated rats. Therefore, our results suggest that in rats with experimental chronic cholangitis, hepatic T cell GR downregulation appears to impair GR function (that is, inhibition of IL-2R expression).

The mechanism for GR downregulation in hepatic T lymphocytes of ANIT fed rats is not known. Various factors such as ligand induced downregulation, inflammatory cytokines, and cellular activation may influence GR expression. Glucocorticoids are known to regulate their own receptor expression at various levels (reviewed by Oakley and Cidlowski). Given that GR downregulation in ANIT treated rats is tissue specific, and not a generalised effect in response to elevated circulating glucocorticoid levels, it seems unlikely that hepatic T lymphocyte GR downregulation in ANIT treated rats is simply due to glucocorticoid mediated auto-regulation of receptor expression.

In recent years, a number of cytokines have been shown to modulate GR number and/or affinity, thereby contributing to glucocorticoid resistance. Kam et al demonstrated that incubation of human PBMCs with both IL-2 and IL-4 together resulted in a decrease in GR number without a change in GR number. However, IL-2 and IL-4 are unlikely to play a significant role in decreased hepatic T cell GR expression in ANIT fed rats for two reasons. Firstly, these cytokines do not appear to cause a reduction in GR number. Secondly, hepatic IL-4 levels in ANIT treated rats did not differ from those observed in control rats. However, other cytokines may play a role in modulating GR expression in hepatic T cells of rats with chronic cholangitis.

In addition, stimulation of immune cells with phorbol esters and the cytokine tumour necrosis factor-α (TNF-α) have been shown to modulate cellular GR number. However, we found that incubation of splenic T cells with either TNF-α or phorbol-12-myristate 13-acetate for up to 48 hours did not decrease cellular GR numbers (Tjandra K, unpublished data). In summary, hepatic T lymphocyte resistance to elevated circulating glucocorticoids occurs in rats with experimental cholangitis. The mechanism for steroid resistance in this model may be attributed, at least in part, to downregulation of GR in hepatic T lymphocytes. Hepatic T cell GR downregulation in ANIT induced experimental cholangitis is tissue specific and appears to be independent of circulating glucocorticoid levels. Currently, the mechanism underlying GR downregulation in hepatic T lymphocytes during experimental cholangitis is unknown but is of obvious potential clinical relevance in treating these diseases. Our findings may explain, at least in part, why patients with chronic cholangitis do not respond clinically to glucocorticoid therapy.

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