Identification of novel molecules and pathogenic pathways in primary biliary cirrhosis: cDNA array analysis of intrahepatic differential gene expression

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Abstract

Background—Primary biliary cirrhosis (PBC) is an autoimmune disease in which the pathogenesis of progressive liver injury is poorly understood.

Aim—To provide novel insights into the pathogenesis of PBC related liver injury using cDNA array analysis, which simultaneously examines expression of many genes.

Methods—Utilising cDNA arrays of 874 genes, PBC was compared with primary sclerosing cholangitis (PSC) associated cirrhosis and non-diseased liver. Differential expression of 10 genes was confirmed by real time quantitative reverse transcriptase-polymerase chain reaction (RT-PCR).

Results—Array analysis identified many differentially expressed genes that are important in inflammation, fibrosis, proliferation, signalling, apoptosis, and oxidative stress. PBC was associated with increased expression of both Th1 and Th2 type molecules of the immune response. Fibrosis related gene expression featured upregulation of connective tissue growth factor and transforming growth factor beta3. Many more apoptosis associated molecules exhibited increased expression, consistent with apoptosis being a more active and regulated process, in PSC associated cirrhosis than in PBC. Increased expression of many genes of the Wnt and notch pathways implicated these highly conserved and linked pathways in PBC pathogenesis. The observed increases in expression of c-jun, c-myc, and c-fos related antigen 1 are consistent with increased Wnt pathway activity in PBC. Differential expression of four components of the Wnt pathway, Wnt-5a, Wnt-13, FRTZ, and beta-catenin, was confirmed by quantitative RT-PCR.

Conclusion—Many genes implicated in intrahepatic inflammation, fibrosis, and regeneration were upregulated in PBC cirrhosis. In particular, increased expression of a number of Drosophila homologues was seen in PBC.

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The pathogenesis of primary biliary cirrhosis (PBC) involves immune mediated injury of bile ducts and is characterised by multiple auto-antibodies to mitochondrial antigens (antimitochondrial antibodies (AMA)).1–3 Investigation into the pathogenesis of PBC has focused principally on the mitochondrial antigens,1 AMA characterisation,2 and the nature of the immune infiltrate around bile ducts.1 Targeting of the biliary epithelium may be explained by abnormal expression of mitochondrial antigens on the luminal surface of biliary epithelial cells.1 The aberrant mitochondrial antigen expression is thought to result in a breakdown of tolerance leading to immune mediated liver injury.1 An alternative hypothesis that PBC results from an immune response initiated due to an infectious agent is under investigation.1

In PBC, damage to the biliary epithelium and progressive liver injury leading to cirrhosis appears to be mainly due to a cell mediated immune response. The mononuclear cell infiltrate in PBC is characterised by activated CD4+ and CD8+ T lymphocytes with a predominant Th1 response.4,5 However, we have observed intrahepatic upregulation of interleukin (IL)-2 but not interferon (IFN)-γ in PBC, which suggests that PBC does not universally conform to the Th1 paradigm.6

Abbreviations used in this paper: ABI, Applied Biosystems Incorporated; AMA, antimitochondrial antibodies; BDNF, brain derived neurotrophic factor; CDK, cyclin dependant kinase; cDNA, complementary DNA; CTGF, connective tissue growth factor; CXCR, CXC chemokine receptor; dATP, 2’ deoxy -adenosine 5’ triphosphate; dCTP, 2’ deoxy -cytidine 5’ triphosphate; dGTP, 2’ deoxy -guanosine 5’ triphosphate; dNTP, deoxy-nucleotides; dTTP, 2’ deoxy-thymidine 5’ triphosphate; EMMPRIN, extracellular matrix metalloproteinase inducer; EGF, epidermal growth factor; FGF, fibroblast growth factor; FRITZ, secreted frizzled related protein; Fz, frizzled; GDNF, glial cell line derived neurotrophic factor; HGF, hepatocyte growth factor; HSC, hepatic stellate cell; HSP, heat shock protein; IAP, inhibitor of apoptosis protein; IFN, interferon; IGF/IR, insulin-like growth factor binding protein; IL, interleukin; MCP, monocyte chemoattractant protein; MMP, matrix metalloproteinase; MRP-8, migration inhibitory factor related protein 8; NGF, nerve growth factor; NT, neurotrophin; PBC, primary biliary cirrhosis; PCR, polymerase chain reaction; PDGF, platelet derived growth factor; PSC, primary sclerosing cholangitis; PTG, patch homologue; RT, reverse transcriptase; SCL, stem cell protein; TAF, TATA box binding protein associated factor; TAL-1, stem cell protein; TGF, transforming growth factor.
T cells of most patients recognise purified oxo-acid dehydrogenase mitochondrial multi-
enzyme complex antigens but the role of autoantibodies to these antigens is controver-
sial. There is some evidence of “molecular mimicry” where a cross reactive antigen is target-
ged by an immune response initially induced by a micro-organism. The presence of a cross reactive antigen is possible given that the titre of AMA and the T cell response to mitochondrial antigens do not always correlate with disease severity or even the presence of disease. Binding of AMA to purified biliary epithelial cells differs markedly for various antibodies suggesting that there are multiple epithelial antigens with epitope cross reactivity.

The immune response in PBC results in progressive bile duct damage and tissue fibrosis. Many fibrotic mediators in PBC may arise from damage to biliary epithelium with release of fibrogenic growth factors such as epidermal growth factor (EGF), basic fibro-
blast growth factor (FGF), transforming growth factor (TGF) beta, and platelet derived growth factor (PDGF). These fibrotic mediators lead to activation of the hepatic stel-
late cell (HSC) with release of basic FGF, hepatocyte growth factor (HGF), and TGF-
beta. Autocrine stimulation of the activated HSC with PDGF, basic FGF, and TGF-beta, and stimulation of the biliary epithelium with EGF and HGF help maintain the initiated fi-
brogenic stimulus. Inflammatory media-
sors such as monocyte chemotactic factor 1 (MCP-1) are thought to stimulate the fibrotic response in PBC.

Analysis of differential gene expression is an important approach to understanding liver injury pathogenesis. Classical approaches have examined mRNA expression of individual can-
didate genes such as growth factors (for exam-
ple, connective tissue growth factor (CTGF), cytokines (for example, IL-8), and chemokines (for example, IP-10)). Unfortunately, methods of detecting differential gene expression such as northern blot analysis and semi-quantitative reverse transcriptase-
polymerase chain reaction (RT-PCR) are labo-
rious and have explored few genes. Pathogenic processes involved in liver injury, such as inflammation, proliferation, apoptosis, and fibrosis, are inter-related and increasing num-
bers of genes are being identified as important in each pathogenic process. Clearly, appro-
aches that can simultaneously examine dif-
ferential expression of many genes are likely to lead to significant advances in understanding pathogenic pathways involved in liver injury. Therefore, in this study we used complemen-
tary DNA (cDNA) array analysis to examine intrahaptic differential gene expression in PBC. This paper describes multiple novel observations of differentially expressed genes and pathways implicated in the above men-
tioned pathogenic processes. Interestingly, the Wnt pathway was strongly implicated in PBC pathology for the first time.

Methods

Tissue and RNA isolation

Total RNA was isolated from end stage cirrhotic PBC (n=6) and end stage cirrhotic primary sclerosing cholangitis (PSC) (n=4) tissue ob-
tained from liver explants. Non-diseased tissue was obtained from transplant donor liver biopsies (n=4) and from normal liver (distant from the tumour margin) during hepatic metas-
tasis resection (n=4). Tissue was obtained following institutional ethics committee approval and Australian Medical Research Coun-
cil guidelines. Chronic inflammatory activity and end stage cirrhosis were evident in all of the cirrhosis specimens. RNA extraction using gua-
nidine isothiocyanate dissolution and isopropan-
ol precipitation has been described previ-
ously. Poly A+ mRNA for array probe synthesis was isolated from pools of four individual RNA samples using oligo-dT<sub>18</sub> cellulose (Roche Mo-
lecular Systems Inc., Branchburg, New Jersey, USA) by standard methods. The pooling of samples is a means of normalising for individual differences in array analysis.

cDNA array analysis

Two nylon membrane based cDNA arrays were used: (a) ATLAS Human Gene Array 1.0 (588 genes and nine housekeeping genes) and (b) ATLAS Cytokine/Receptor Array (268 genes and nine housekeeping genes) (Clontech Laboratories, Inc., Palo Alto, California, USA) (fig 1). A list of all of the genes on these two arrays with summaries of gene functions is at www.clontech.com/atlas.

Probes from cirrhotic PBC, cirrhotic PSC, donor liver, and normal liver tissues were hydridised to both an ATLAS Cytokine Array and an ATLAS Human Gene Array. Probe synthesis was performed as recommended by the manufacturer with some modifications. During first strand cDNA synthesis, 1–2 µg of poly A+ mRNA were labelled in a 20 µl reaction with 7.5 µl of α<sup>33</sup>P-dCTP (10 µCi/µl, 3000 Ci/mmol; NEN Life Science Products, Inc., Boston, Massachusetts, USA) with 1 µl of Superscript II RT (Gibco-BRL Gaithersburg, Maryland, USA), 4 µl of 5× reaction buffer, 2 µl of 100 mM dithiothreitol, 0.5 µl of RNase inhibitor (Promega Corp., Madison, Wisconsin, USA) with 1 µl of gene specific primer mix (Clontech), water, and 2 µl of deoxy-
nucleotides (dNTPs) (5 mM 2’ deoxy-
adenosine 5’ triphosphate (dATP), 2’ deoxy-
thymidine 5’ triphosphate (dTTP), 2’ deoxy-
guanosine 5’ triphosphate (dGTP), and 50 µM 2’ deoxy-cytidine 5’ triphosphate (dTCP); Amersham Pharmacia Biotech UK Ltd, Buck-
inghamshire, UK) at 48°C for 40 minutes. Unincorporated nucleotides were removed using resin based Chroma spin-200 columns as recommended by the manufacturer (Clontech). Probes at a final concentration of 2–4×10<sup>7</sup> CPM/ml were hybridised to each array membrane for approximately 15 hours in Express Hyb hybridisation solution at 68°C (Clontech). The array membranes were washed in decreasing concentrations of SSC. Signals from the array membrane were quantifi-
ced using a GS-525 Molecular Imager and

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Molecular Analyst Software (Bio-Rad Laboratories Inc., Hercules, California, USA). Additionally, all arrays were exposed to film (MS Film with an MS Screen; Eastman Kodak Company, New Haven, Connecticut, USA). The signal from a blank portion of the cDNA array was taken as background. Individual gene expression on each array was adjusted for background.

**SELECTION OF HOUSEKEEPER GENE**

On the regression plots, such as that depicted in fig 1D, ubiquitin did not exhibit significant differential expression and was selected as the housekeeper gene. Data on the other eight housekeeper genes were not used because phospholipase A2 and hypoxanthine-guanine phosphoribosyltransferase signals were less than twice the background, and brain specific tubulin alpha 1 subunit, cytoplasmic beta-actin, MHC class I antigen C-4 alpha subunit, and glyceraldehyde 3-phosphate dehydrogenase were differentially expressed in cirrhosis.

**SENSITIVITY AND REPRODUCIBILITY OF cDNA ARRAYS**

Over 95% of the cDNA spots on the array had detectable signals following hybridisation. Signal intensity on the arrays varied by four orders of magnitude. Low abundance signals were defined as signals less than twice background and represented at least 61% of the signals on the arrays. Given the use of $^{32}$P labelled probes to maximise sensitivity, 90 signals were excluded from subsequent analysis due to blooming or membrane contamination. The raw housekeeping gene signals were greater from the cirrhotic liver probes than from the normal or donor liver probes, causing our experiments to preferentially detect up- rather than downregulation. Data were used only when at least one of the signals was 50% or more above background.

Pairwise comparative gene expression was determined from signal intensities with the four different probes (PBC cirrhosis, PSC cirrhosis, donor tissue, and normal tissue). Data are presented as a ratio generated by one of two methods:

1. Signal intensities converted to a ratio adjusted for background and housekeeper gene (ubiquitin) expression:

   \[
   \frac{(\text{gene A probe A} - \text{background from probe A})/(\text{housekeeper gene probe A} - \text{background from probe A})}{(\text{gene A probe B} - \text{background from probe B})/(\text{housekeeper gene probe B} - \text{background from probe B})}.
   \]

2. Regression analysis, a method that did not require adjustment for background or housekeeper gene signal (fig 1D). A non-linear power
regression analysis on a logarithmic scale was used because the signal intensities varied by four orders of magnitude. Each observed signal was divided by each calculated signal, calculated from the regression line formula derived from all signals in each pairwise tissue comparison.

The reported ratio comparing expression of each gene used the method that generated the greater value. Individual genes that exhibited greater than 1.5-fold upregulation were categorized with the pathogenic processes of inflammation, fibrosis, proliferation, stress response/oxidative stress, apoptosis, and intracellular signalling pathways. Genes were categorized by pathogenic process according to the array manufacturer’s guidelines and known gene functions. Many genes could be included in multiple categories, so genes were grouped into pathogenic processes using this descending hierarchy: (i) intracellular signalling and nuclear, (ii) apoptosis, (iii) inflammation, (iv) proliferation, (v) fibrosis and (vi) stress response/oxidative stress associated genes, and (vii) uncategorised genes.

In addition, comparative expression data on individual genes were plotted graphically; genes were clustered according to the nature and extent of their differential expression over the three comparisons. Clustering used the Cluster and TreeView software (M Eisen, Stanford University). This hierarchical clustering method uses pairwise average linkage analysis and no a priori knowledge of gene function.

QUANTITATIVE RT-PCR
To confirm differential expression in individual patient specimens, mRNA of a selection of genes was quantified by real time RT-PCR using a modification of published methods. The genes assayed were: Wnt-5a, Wnt-13, CTGF, extracellular matrix metalloproteinase inducer (EMMPRIN), stromal cell-derived factor 1 receptor (CXCR4), follistatin, secreted frizzled related protein 3 (FRITZ), beta-catenin, Jagged-1, and patch homologue (PTC).

Briefly, 5 µl of total RNA (5 µg) was primed with 3 µl (500 ng) of oligo-dT18 primer (Roche Diagnostics, Indianapolis, Indiana, USA) and then reverse transcribed at 42°C for 1 hour. The reaction mix (20 µl) also contained 4 µl of 5× reaction buffer, 2 µl of 100 mM dithiothreitol, 0.5 µl of RNase inhibitor (Promega), 1 µl of 10 mM dNTPs (Amersham), 1 µl of Superscript II (Gibco-BRL), and 3.5 µl of H2O. After cDNA synthesis the total volume was made up to 80 µl with H2O. Two cDNA samples were prepared from each RNA sample and then stored at −70°C until needed. Each real time PCR reaction contained 1 µl of CDNA, 2.5 µl of 10× TaqMan reaction buffer A (PE Biosystems, Foster City, California, USA), 4 µl of 25 mM MgCl2, 0.5 µl of 10 mM dNTPs, 1 µl of 5 mM forward and reverse primer, 0.18 µl (5.5 U/µl) of AmpliTaq Gold polymerase (PE Biosystems), 0.25 µl of 50× Sybr Green1 (Molecular Probes, Eugene, Oregon, USA), and H2O for a 25 µl total volume. Cycling parameters were 95°C for 10 minutes followed by 40 cycles of 95°C for 30 seconds, 40 second ramp, 60°C for 30 seconds, 20 second ramp, 72°C for one minute, 17 second ramp, and 30 seconds for fluorescence measurement. This measurement resolved the gene of interest from non-specific products of amplification such as primer-dimers and was achieved using a measurement temperature that ranged from 82 to 86°C (determined by melting point analysis of each product). The reactions were performed in duplicate using an Applied Biosystems Incorporated (ABI) Model 7700 Sequence Detector (PE Biosystems) and analysed using ABI Prism Sequence Detector Software v1.6.3 (PE Biosystems). A gene specific DNA standard was included in each assay. Confirmation of a single PCR product was performed by gel analysis and the identity of the amplified PCR product was confirmed by sequencing (ABI 373 Sequencer using dye terminator chemistry; PE Biosystems). Ubiquitin, the housekeeper gene, was quantified in all cDNAs.

The primer sequences were: Wnt-5a forward 5' GCAATGTCTTCCAAGTTTC 3', reverse 5' AAGTGCGCAGTTTCTTC 3'; Wnt-13 forward 5' GAGTGTGACGACAAATTCC 3', reverse 5' CACCCCGATCAAAGTCTC 3'; CTGF forward 5' TCCACCACAAATTCAAAAC 3', reverse 5' CAGATTAGCAAGGCATATTACGG3'; CXCR4 forward 5' TCAGTGGAGCAGTCAG 3', reverse 5' TCCCAATTGAAGGCAG 3'; EMMPRIN forward 5' ACAAGATCGCTGACTCTGAGAC 3', reverse 5' TTCTCAAGTGTAGCTCTGACC 3'; follistatin forward 5' CCTCAACCCATCTTTCAAC 3', reverse 5' CCCCTTCTGATTTTCTCC 3'; beta-catenin forward 5' CATTACACTCTTCACAACCC 3', reverse 5' CAGATAGCACTTTCCAGC 3'; FRITZ forward 5' CAGTAGGTGGAGGTGAAGGAG 3', reverse 5' GAGTCCCAAAGATGAGGAAG 3'; PTC forward 5' CCCCAACAAAAATTCACACC 3', reverse 5' CATCATCCACACCAACACC 3'; Jagged-1 forward 5' GGACTATGAGGGCAAGAAC 3', reverse 5' CGGTTCCTTACATTATGGT 3'; and ubiquitin forward 5' GGTTAGCTTCTGGTGGAAAAC 3', reverse 5' AATGCGCTTCCTTGCTCCTG 3'.

STATISTICAL ANALYSIS
All results are expressed as mean (SEM). Statistical comparisons were performed using a non-parametric Mann-Whitney U test. Statistical analysis was performed using Statview 4.5.1 (Abacus Concepts California, USA) and regression plots were generated using Kaleideagraph 3.0 (Synergy Software, Pennsylvania, USA).

Results
GROUNDBASE OF PBC
Clustering of genes based on their relative expression (fig 2) indicated gene groups that were similarly upregulated or downregulated and was a rapid screen for gene groups of interest. This clustering analysis revealed that greater than 70% of genes had a similar pattern of expression in PBC and PSC associated cirrhosis compared with non-diseased tissue.

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Although the patterns of gene expression were similar, there were differences in the degree of differential gene expression of both PBC and PSC-associated cirrhosis compared with non-diseased tissue. All differential expression data are depicted by colour intensity in both fig 2 and the supplemental material clustering figure (please see website for supplemental data, at www.gutjnl.com). Additionally, the supplemental material clustering figure includes the vertical (gene) dendrogram labelled with individual gene names.

**DIFFERENTIAL GENE EXPRESSION IN PBC**

Of interest among the most upregulated genes, Wnt-13, follistatin, and Jagged-1 expression levels were greater in PBC than in either non-diseased or PSC-associated cirrhotic liver (supplemental material—see website). Brain-derived neurotrophic factor (BDNF) receptor (Trk3), protein serine/threonine kinase 1, stem cell protein (SCL or TAL-1), and Wnt-2 were greatly increased in both PBC and PSC-associated cirrhosis compared with non-diseased tissue (supplemental material—see website). Ephrin A3 was the most strikingly downregulated gene and migration inhibitory factor related protein 8 (MRP-8/calgranulin A) and the receptor for cytotoxic tumour necrosis factor related apoptosis inducing ligand were markedly downregulated in both PBC and PSC-associated cirrhosis (supplemental material—see website).

Gene upregulation at 2.0-fold or greater is presented in figs 3 and 4. The supplemental material tables list genes that exhibited 1.5-fold or greater differential expression while the entire cDNA array differential expression data is in the supplemental material clustering figure (see website).

**DIFFERENTIAL EXPRESSION IN PBC COMPARED WITH NON-DISEASED LIVER**

*Inflammation (fig 3A).*

CXCR4 and flt3 receptor were the most dramatically increased inflammation associated genes, each with greater than sixfold increases in expression in PBC compared with non-diseased tissue. Neither a Th1 nor a Th2 predominant cytokine response was observed. The Th2 associated cytokines IL-4, IL-5, and IL-13 and the Th1 associated molecules lymphotoxin-beta, IFN-γ receptor beta subunit, and tumour necrosis factor receptors were increased in PBC compared with non-diseased liver. Increased expression in PBC of the chemokines CXCR4 and MCP-1 was observed. CXCR4, flt3 receptor, myeloid cell nuclear differentiation antigen, IL-3, IL-5, IL-15, IL-17, and granulocyte-macrophage colony stimulating factor were upregulated greater than threefold in PBC compared with non-diseased liver.

*Fibrosis (fig 3B).*

The most dramatic increases in the fibrosis category were CTGF and TGF-beta3, which were increased greater than 10-fold in PBC. Notably, platelet activating factor receptor, FGF receptor 1, FGF receptor 3, cysteine rich FGF receptor, acidic FGF, and PDGF receptor were increased in PBC. Interestingly, upregulated mediators of fibrosis included several TGF family members, TGF-beta3, TGF-beta2, and TGF-beta1 (1.5-fold, see supplemental material at the website). Importantly, recognised fibrotic mediators such as EMMPRIN were increased in PBC (1.7-fold, see supplemental material at the website). Clearly, integrin alpha 5 and 6 (VLA-5 and 6; CD49e and CD49f), which we categorised under inflammation (fig 3A), are also implicated in fibrogenesis.
Regeneration, growth, and proliferation (supplemental material—see website)

Expression of many highly conserved genes of the Wnt and notch pathways and four neurotrophin associated genes were increased in PBC. Indeed, of the 16 genes in this category that exhibited greater than threefold increases in expression, 13 were either Drosophila homologues (fig 4) or neural genes. The Drosophila homologues included Wnt-2, Wnt-13, Wnt-5a, and Wnt-10b/Wnt-12 of the Wnt pathway, and Jagged-1, Jagged-2, notch-1, notch-2, and notch-3 of the notch pathway (fig 4). The upregulated neurotrophin associated genes were BDNF receptor/Trk3, BDNF, glial cell line derived neurotrophic factor (GDNF), and ret proto-oncogene (part of the GDNF receptor complex) (supplemental material—see website). Many genes of the insulin growth axis such as insulin-like growth factor binding protein (IGFBP)-6, IGFBP-2, and insulin receptor, were increased in PBC. Furthermore, cyclin-D1, cyclin dependant kinase inhibitor (CDKN)-1C, CDKN-1A, and CDKN-3 were increased in PBC. Delta-like protein, ataxia telangiectasia, Erb B4, CDKN-1C, and IGFBP-6 were increased threefold or greater in PBC compared with non-diseased tissue.

Intracellular signalling and nuclear (supplemental material—see website)

Synapse associated protein 97 and stem cell protein (SCL or Tal-1) were increased greater than ninefold in PBC. Many genes involved in intracellular signalling pathways had increased expression in PBC. They included genes that are recognised downstream targets of Wnt signalling, including c-Jun, c-Myc, c-Fos related antigen 1, and beta-catenin. Many transcription factors including helix-loop-helix protein HLH 1R2 and zinc finger in the multiple endocrine neoplasia 1 locus and mediators of several pathways, including the Erb B pathway, of which transducer of Erb B2 is a component, were increased in PBC.

Stress response/oxidative stress (supplemental material—see website)

The stress response was characterised by increased expression of the heat shock proteins (HSP)70.1, HSP90A, and HSP27 and genes involved in oxidative stress including glutathione-S-transferase M1, dioxin inducible cytochrome P450 1B1, and cytosolic superoxide dismutase 1.

Uncategorised genes (supplemental material—see website)

Many genes were not categorised because they have either multiple or poorly characterised functions. Follistatin, prorelaxin-H2, and ephrin associated genes including ephrin receptor 2 and 3 and ephrin associated genes including ephrin type A receptors were increased in PBC. Indeed, of the 16 genes in this category that exhibited greater than threefold increases in expression, 13 were either Drosophila homologues (fig 4) or neural genes. The stress response was characterised by increased expression of the heat shock proteins (HSP)70.1, HSP90A, and HSP27 and genes involved in oxidative stress including glutathione-S-transferase M1, dioxin inducible cytochrome P450 1B1, and cytosolic superoxide dismutase 1.

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cDNA array analysis of PBC

Apoptosis genes (fig 3C)

Only the proapoptotic early response protein NAK1/TR3 orphan receptor was increased greater than twofold in PBC.

Differential expression in PBC compared with PSC

Gene upregulation in PBC compared with PSC associated cirrhosis helped identify genes preferentially involved in the pathogenesis of PBC compared with another type of "biliary" cirrhosis. The inflammation associated genes IL-3, IL-4, IL-5, IL-15, IL-17, and CXCR4 had greater expression in PBC compared with PSC associated cirrhosis (fig 3A). CTGF and TGF-beta3 expression was greater in PBC than in PSC associated cirrhosis (fig 3B).

Many mammalian homologues of highly conserved genes involved in growth and regeneration originally identified in Drosophila were increased in PBC compared with PSC associated cirrhosis. These included FRITZ, Wnt-13, Wnt-5a, notch-1, notch-2, notch-4, Jagged-1, Jagged-2, and smoothened (fig 4).

Additionally, the GDNF receptor ret was increased in PBC compared with PSC. Differential expression was seen in some stress related genes, notably HSP27.

Data on cytoplasmic signalling and nuclear associated genes revealed striking differences between PBC and PSC associated cirrhosis. An increase of greater than 80-fold of transcription initiation factor 250 kDa subunit (TAFII250) was seen in PBC compared with PSC associated cirrhosis. Interestingly, expression of c-src, Rad52, and c-myb was greater in PBC compared with PSC. In contrast, some signalling associated genes such as neural-cadherin and several oncogenes, such as SmaN and p56-ck, were markedly upregulated in PSC compared with PBC (supplemental material—see website).

Notable uncategorised genes that exhibited greater expression in PBC compared with PSC were follistatin, FRITZ, prorelaxin H2, renin binding protein, angiotensin II type 1A receptor, and thrombomodulin.

An interesting difference between PBC and PSC associated cirrhosis was that in contrast with PBC, many apoptosis associated genes were upregulated in PSC (fig 3D). These included inhibitor of apoptosis protein (IAP)-1, IAP-2, IAP-3, four anti-apoptotic Bcl-2 related proteins, Bcl-2, Bcl-2 A1, Bcl-x, and Bcl-2 adenovirus E1B 19 kDa interacting protein, and the pro-apoptotic molecules Bak, Bik, and Bax of the Bcl-2 family.

Statistical analysis of array data

Correlation coefficients following regression analysis for the ATLAS Human Gene Array and ATLAS Cytokine Receptor Array data were, respectively: PBC compared with normal liver \( r^2 = 0.77 \) and 0.85; PBC compared with donor liver \( r^2 = 0.84 \) and 0.93; and PBC compared with PSC \( r^2 = 0.86 \) and 0.85, with p<0.001 for all three comparisons.

There were 190 genes common to both cDNA arrays. With the four probes on the two arrays there was a strong correlation of the 760 duplicate signals \( (r^2 = 0.76, p<0.001) \). Similarly, the 760 duplicate signals were compared by subtracting the regression ratio determined in comparisons of PBC or PSC associated cirrhosis with non-diseased tissue. The 760 duplicate signals showed a mean difference of only 0.084 (0.017).

The two methods of determining differential gene expression were compared by subtracting, for each gene, the ratio determined by regression analysis from the ratio determined by adjustment for ubiquitin expression (see methods). The three comparisons, of PBC with normal, donor, and PSC, used a total of 2359 determinations of differential gene expression and found that the mean difference between the two methods was 0.29 (0.058) ratio units. In addition, we subtracted, for each gene, the ratios generated by the donor and normal liver samples when each were compared with PBC. The 1572 duplicate comparisons showed a mean difference between donor and normal liver of only 0.10 (0.058) ratio units, indicating the close similarity of the two types of non-diseased liver.

Quantitative RT-PCR

Increased expression of 10 selected genes, mainly genes associated with fibrosis and the Wnt pathway, was confirmed on individual patient specimens, including all specimens analysed on the cDNA arrays (fig 5). Gene expression was quantified by real time RT-PCR twice on two separate cDNA syntheses. CTGF was the most abundantly expressed gene with \( 4.6 \times 10^6 \) (0.4 \times 10^6) \( \times \) relative gene copies per \( \mu g \) of total RNA in PBC. Wnt-13 was the least abundant transcript quantified, with only 64 copies per \( \mu g \) of total RNA in non-diseased liver. The cDNA arrays comparing PBC with non-diseased liver showed increases of 13.6x for CTGF, 7.2x for CXCR4, 1.7x for EMMPRIN, 10.7x for follistatin, 5.7x for Wnt-13, 3.6x for Wnt-5a, 1.8x for beta catenin, 4.4x for FRITZ, 7.9x for Jagged-1, and 6.7x for PTC. Thus the array and RT-PCR data were concordant.

Discussion

Intrahepatic cDNA array analysis is a new approach to the study of gene expression in cirrhosis. Utilising this approach we have identified many novel interesting differences at the mRNA level in the molecular pathways of PBC compared with non-diseased liver and PSC associated cirrhotic liver. Identifying differential expression of genes associated with inflammation, fibrosis, apoptosis, proliferation, regeneration, neural cells, and the Wnt and notch pathways enabled identification of molecular pathways that may be involved in PBC pathogenesis.

The novel observations of differential expression included many genes of the Wnt and notch pathways, both pathways being highly conserved and initially identified in Drosophila development and differentiation. The observations are consistent with increased Wnt signalling rather than increased turnover of Wnt.
pathway components. In addition to the downstream increase in beta-catenin expression, most of the genes classically activated by the Wnt pathway were increased in PBC, including c-myc, c-jun, c-fos related antigen 1, and cyclin-D1. The exact function of the Wnt pathway in cirrhosis is unknown. Multiple Wnt pathway genes have been implicated in tumorigenesis but the incidence of hepatocellular carcinoma in PBC is less than in other types of cirrhosis. The physiological role of the Wnt pathway in adult organs is poorly understood. The Wnt response transcription factor T cell factor 4 is greatly expressed in normal adult liver but its physiological function there is unknown.\textsuperscript{30} Differentiation of haematopoietic progenitor cells involves Wnt pathway genes,\textsuperscript{31} mesenchymal to epithelial conversion in the metanephros requires Wnt expression,\textsuperscript{32} and a regulator role for the Wnt pathway and T cell factor 4 in differentiation within the vertebrate gut epithelium is likely.\textsuperscript{33} Replacement of the resident fibroblast population in the synovium of rheumatoid arthritis patients with immature bone marrow and mesenchymal cells is accompanied by increased Wnt-5a/Fz5 expression.\textsuperscript{33} The increased expression of Wnt-5a, FRITZ, and beta-catenin in PBC, PSC, and HCV cirrhosis (fig 5) suggests that the Wnt pathway is active in all forms of cirrhosis. However, the frequency of Wnt pathway gene detection in PBC suggests that it may play a more specific role in PBC pathogenesis (fig 4). Clearly a more detailed examination of the Wnt pathway in cirrhosis is needed.

Figure 5 Differential gene expression examined by reverse transcriptase-polymerase chain reaction (RT-PCR). Quantitative real-time RT-PCR data (mean (SEM)) on mRNA from primary biliary cirrhosis (PBC) (n=6), primary sclerosing cholangitis (PSC) associated cirrhosis (n=4), hepatitis C virus (HCV) cirrhosis (n=6), and non-diseased donor liver (n=4). The depicted differential expression data are in two groups: (A) follistatin, CXC chemokine receptor 4 (CXCR4), connective tissue growth factor (CTGF), and extracellular matrix metalloproteinase inducer (EMMPRIN) and (B) Drosophila homologues including Wnt-13, secreted frizzled related protein 3 (FRITZ), beta-catenin, and Jagged 1.

\begin{figure}
\centering
\includegraphics[width=\textwidth]{figure5}
\caption{Differential gene expression examined by reverse transcriptase-polymerase chain reaction (RT-PCR). Quantitative real-time RT-PCR data (mean (SEM)) on mRNA from primary biliary cirrhosis (PBC) (n=6), primary sclerosing cholangitis (PSC) associated cirrhosis (n=4), hepatitis C virus (HCV) cirrhosis (n=6), and non-diseased donor liver (n=4). The depicted differential expression data are in two groups: (A) follistatin, CXC chemokine receptor 4 (CXCR4), connective tissue growth factor (CTGF), and extracellular matrix metalloproteinase inducer (EMMPRIN) and (B) Drosophila homologues including Wnt-13, secreted frizzled related protein 3 (FRITZ), beta-catenin, and Jagged 1.}
\end{figure}
The notch (receptor)/Jagged (ligand) pathway has been partially characterised in human disease.\textsuperscript{12} Mutations in the Jagged-1 gene are responsible for Alagille syndrome.\textsuperscript{13} Jagged-1 expression in the adult liver is localised to zone three hepatocytes and biliary epithelium.\textsuperscript{36} Notch-1, -2, and -3 as well as Jagged-1 and -2 are implicated in the determination of both gamma-delta and alpha-beta T cells as well as the CD4\textsuperscript{+} versus CD8\textsuperscript{+} thymocyte lineage choice.\textsuperscript{37} The cell fate determination function that the Wnt and notch pathways share is partially explained by the interaction of Wnt proteins with notch receptors and the binding of the regulatory protein dishevelled to both frizzled and notch receptors.\textsuperscript{39} 40 The observed expression relationship to PBC pathogenesis remains to be determined.

A documented function of Wnt and notch pathways is regulation of the growth and regeneration related molecules, follistatin and activin. It is therefore of interest that follistatin and activin were increased in both PBC and PSC associated cirrhosis. Wnt and FGF expression inhibit activin activity while the Wnt and notch pathways may regulate follistatin.\textsuperscript{41} 42 Intrahepatic follistatin stimulates proliferation and is antiapoptotic. Follistatin can accelerate liver regeneration following partial hepatectomy.\textsuperscript{43} 44 Conversely, activin is an autocrine inhibitor of hepatocyte DNA synthesis,\textsuperscript{45} retards hepatocyte proliferation, and functions in the maintenance of constant liver mass.\textsuperscript{46} Follistatin is an activin binding protein and generally the effects of follistatin are antagonised by activin.\textsuperscript{47} 48 Activin infused intravenously leads to apoptosis around the central vein of the hepatic lobule, an effect ablated by adding follistatin.\textsuperscript{49} The mechanism of follistatin mediated control of liver growth is not understood. EGF, TGF-beta, phentylephrine, and glucagon, all of which are known to influence hepatocyte regeneration, upregulate follistatin mRNA expression in hepatocytes.\textsuperscript{50} The possible relationship of the Wnt, notch/jagged, and follistatin/activin pathways may be a common link in influencing hepatocyte proliferation, either specifically in PBC or more generally in any form of cirrhosis. CDNA array technology is an unparalleled means of simultaneously examining the mRNA expression of potentially functionally linked genes such as those of the Wnt, notch/jagged, and follistatin/activin pathways in hepatocyte proliferation.

A principle feature of PBC is intrahepatic inflammation. PBC was characterised by our cDNA array analysis as involving a mixed Th1 and Th2 immune response. Previous studies have generally suggested that PBC predominantly involves a Th1 response.\textsuperscript{51} In situ hybridisation studies in PBC patients\textsuperscript{1} and an animal model of PBC\textsuperscript{2} have shown increased IFN-\gamma expression consistent with a Th1 response. However, mixed Th1 and Th2 cytokine expression has been observed in a different animal model of PBC\textsuperscript{3} and we previously demonstrated significant increases in intrahepatic IL-2, IL-6, IL-8, and TGF-beta expression in PBC.\textsuperscript{52} Furthermore, PBC patient derived peripheral T cell clones are heterogeneous for Th1/Th2 profile.\textsuperscript{53} Examining few cytokines may misrepresent the Th1/Th2 profile. Our data highlight the principle strength of cDNA array analysis in simultaneously examining expression of many genes.

Fibrosis associated gene expression was well defined by cDNA array analysis. Increases in growth factors such as FGF-1 and FGF-3 in PBC are consistent with known pathways of both paracrine and autocrine stimulation of HSCs.\textsuperscript{54} Biliary epithelial cells and hepatocytes, in response to the immune mediated damage or cytokine signals such as MCP-1, are likely additional sources of these profibrotic molecules.\textsuperscript{55} The increased expression of TGF-beta 2 and 3 concords with evidence that all TGF-beta isoforms mediate hepatic fibrosis.\textsuperscript{56} 57 TGF-beta, CTGF, and EMMPRIN were upregulated in both PBC and PSC (fig 3B). CTGF is a HSC product released following stimulation by TGF-beta, so their linked differential expression was expected.\textsuperscript{58} In contrast, EMMPRIN is not known to be a fibrotic mediator of human cirrhosis. EMMPRIN is expressed by tumour and epithelial cells and interacts with fibroblasts resulting in increased expression of MMP-1, MMP-2, (collagenase) and MMP-3 (stromelysin 1).\textsuperscript{59} 63

The above discussion has primarily focussed on differential gene expression in PBC compared with non-diseased liver. Such differential expression may be generally involved in cirrhosis or inflammation rather than be specific for PBC. Therefore, PBC was compared with another “biliary” cirrhosis, PSC. The overall gene profile data indicate that regulation of liver gene expression in these two diseases is somewhat similar with 70% of examined genes similarly expressed. However, there were differences consistent with differing pathogenic pathways in PBC. Most notable was the dramatic difference in expression of apoptosis associated genes between the two types of cirrhosis. This difference is consistent with apoptosis being more active and actively regulated in PSC associated cirrhosis than in PBC. There have been no direct comparisons of apoptosis between PBC and PSC. Apoptosis and cellular proliferation are clearly linked processes. Presently, studies have not implicated apoptosis in the proliferative response of PBC. However, abnormal apoptosis leading to persistence of dysplastic biliary epithelial cells is thought to be involved in the pathogenesis of cholangiocarcinoma complicating PSC.\textsuperscript{64}

Intracellular signalling arising from growth and differentiation responses is clearly important in the pathogenesis of human liver disease. The observed signalling associated differential gene expression was consistent with a PBC specific transcription associated gene profile. Greater TAFII250 expression in PBC than both non-diseased and PSC liver is intriguing. TAFII250 is a TATA box binding protein associated factor (TAF) that participates in the transcription factor TFIID assembly for transcription of cyclin genes, especially cyclin A
and cyclin D1.55 Cell lines without normal TAFII250 function arrest in G1 and transcription of cyclin genes is arrested.56 Why such a dramatic difference in TAFII250 expression is seen in PBC is unclear and warrants further study.

Detection of intrahepatic expression of neural genes was expected given the emerging connection of such genes with HSC differentiation. However, the large number of such genes identified was interesting. Novel differential expression of the neural genes Trk3, GDNF, ret, and synapse associated protein 97 was observed. Trk3, the BDNF receptor, which was increased in both PBC and PSC, is neurotrophin (NT) associated. The NT associated proteins include the ligands nerve growth factor (NGF), NT-3, NT-4, NT-6, and GDNF, and the receptors TrkB, TrkC, and the NGF receptor p75NRTR.67 GDNF has not been studied in liver but is implicated in development; mutations in its receptor ret are responsible for multiple endocrine neoplasia 2 and Hirschsprung disease.68–70 NT's are not neural cell specific; T cells, B cells, and macrophages produce NT's during inflammation.71–73 Increased expression of neural markers might also reflect nerve formation accompanying the increased angiogenesis in cirrhosis. However, the cellular origin of differentially expressed neural genes in liver may originate from HSCs, which have a neural phenotype that includes expression of p75NTR and may originate from the neural crest.54 74–79

A number of caveats relate to the use of cDNA arrays and experiments on whole liver tissue. The cDNA array data are expressed numerically and the statistical analyses convey its precision and reproducibility. However, the strength of this technique is the identification of differentially expressed genes rather than quantification of individual gene expression. To quantify and confirm increased individual gene expression, techniques such as northern blot or quantitative RT-PCR are required.80 81 Hepatocytes are the major contributor to preparations of whole liver mRNA. However, other cell types such as biliary epithelial cells, hepatic stellate cells, endothelial cells, Kupffer cells, T cells, and macrophages probably make significant contributions. In PBC with bile duct loss the proportion of the total mRNA pool derived from biliary epithelium is unknown. Genes expressed specifically by one cell type may be significantly differentially expressed despite small cDNA array ratios. Future studies will analyse mRNA from isolated subpopulations of cells. Such studies require careful interpretation because isolation methods may activate cells, especially when cell culture is involved, and removes interactions that are present in the solid organ, such as those with other cell types and the extracellular matrix. Additionally, differential expression of a mRNA need not lead to the differential expression of its corresponding biologically active protein.

In conclusion, the data presented here are a significant advance in documenting many new observations and multiple simultaneous observations of differential gene expression in PBC in both novel and previously identified pathogenic pathways. A number of these genes have been studied in vitro or in vivo in experiments unrelated to liver pathophysiology. However, given the central roles of the genes we have discussed in processes with hepatic equivalents such as inflammation, fibrosis, apoptosis, signalling, proliferation, and regeneration, they are very likely important in the pathogenesis of PBC. Many genes, such as EMMPRIN, CXCR4, follistatin, and CTGF, were associated with cirrhosis irrespective of aetiology. The use of PSC as a cirrhotic biliary inflammatory comparison helped identify genes potentially important in the differential pathogenesis of PBC and PSC. Finally, the observed differential gene expression of the Wnt and notch pathways particularly implicates these highly conserved Drosophila pathways in PBC pathogenesis. This is the first time the Wnt pathway has been implicated in any form of cirrhosis. Future studies will require localisation of gene expression and a direct analysis of function in pathways related to liver pathobiology.

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cDNA array analysis of PBC


Identification of novel molecules and pathogenic pathways in primary biliary cirrhosis: cDNA array analysis of intrahepatic differential gene expression

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