Research

Open Access

Distinct cellular responses differentiating alcohol- and hepatitis C virus-induced liver cirrhosis

Sharon L Lederer¹, Kathie-Anne Walters¹, Sean Proll¹, Bryan Paeper¹, Shahar Robinzon^{1,2}, Loreto Boix³, Nelson Fausto⁴, Jordi Bruix³ and Michael G Katze^{*1}

Address: ¹Department of Microbiology, University of Washington, Seattle, WA, USA, ²Ben-Gurion University, Beer-Sheva, Israel, ³Barcelona Clinic Liver Cancer Group, Liver Unit, Hospital Clinic IDIBAPS, Barcelona, Spain and ⁴Department of Pathology, University of Washington, Seattle, WA, USA

Email: Sharon L Lederer - leders@u.washington.edu; Kathie-Anne Walters - kathiw@u.washington.edu; Sean Proll - proll@u.washington.edu; Bryan Paeper - paeper@u.washington.edu; Shahar Robinzon - robinzon@bgu.ac.il; Loreto Boix - LBOIX@clinic.ub.es; Nelson Fausto - nfausto@u.washington.edu; Jordi Bruix - bruix@ub.edu; Michael G Katze* - honey@u.washington.edu * Corresponding author

Published: 22 November 2006

Virology Journal 2006, 3:98 doi:10.1186/1743-422X-3-98

This article is available from: http://www.virologyj.com/content/3/1/98

© 2006 Lederer et al; licensee BioMed Central Ltd.

This is an Open Access article distributed under the terms of the Creative Commons Attribution License (<u>http://creativecommons.org/licenses/by/2.0</u>), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Received: 18 October 2006 Accepted: 22 November 2006

Abstract

Background: Little is known at the molecular level concerning the differences and/or similarities between alcohol and hepatitis C virus induced liver disease. Global transcriptional profiling using oligonucleotide microarrays was therefore performed on liver biopsies from patients with cirrhosis caused by either chronic alcohol consumption or chronic hepatitis C virus (HCV).

Results: Global gene expression patterns varied significantly depending upon etiology of liver disease, with a greater number of differentially regulated genes seen in HCV-infected patients. Many of the gene expression changes specifically observed in HCV-infected cirrhotic livers were expectedly associated with activation of the innate antiviral immune response. We also compared severity (CTP class) of cirrhosis for each etiology and identified gene expression patterns that differentiated ethanol-induced cirrhosis by class. CTP class A ethanol-cirrhotic livers showed unique expression patterns for genes implicated in the inflammatory response, including those related to macrophage activation and migration, as well as lipid metabolism and oxidative stress genes.

Conclusion: Stages of liver cirrhosis could be differentiated based on gene expression patterns in ethanol-induced, but not HCV-induced, disease. In addition to genes specifically regulating the innate antiviral immune response, mechanisms responsible for differentiating chronic liver damage due to HCV or ethanol may be closely related to regulation of lipid metabolism and to effects of macrophage activation on deposition of extracellular matrix components.

Background

More than 170 million people worldwide are chronically infected with hepatitis C virus (HCV) [1] and about 20%

of those infected will develop cirrhosis [2]. The incidence of cirrhosis development due to chronic alcohol exposure is similar, and half the causes of death following end-stage liver disease are due to this etiology [3,4]. In alcoholic liver disease (ALD), the metabolism of ethanol plays a role in pathogenesis [5-8], and there is increasing evidence for specific roles played by the immune response [4][5,9-15]. Ethanol can suppress innate immunity in mice by attenuating the TLR3-ds RNA signaling pathway and subsequent interferon (IFN) response [11], as well as in actively drinking patients by decreasing expression of CD28/B7 co-stimulatory pathways [10]. In addition, β chemokine production and migration of mononuclear cells has been observed during later stages of ALD [13].

Early cirrhosis due to HCV infection can be difficult to recognize, as there are often only nonspecific symptoms associated with its progression. The standard method of determining the severity of disease and appropriate treatment has been the liver biopsy [16], but histological examination of the biopsy gives limited information on what is occurring at the molecular level. Therefore, identifying gene expression patterns in liver tissue during cirrhosis development could add valuable information to clinical tests. Functional genomics is increasingly being utilized to investigate the mechanisms underlying the development of liver disease. During chronic HCV infection in humans, broad activation of the endogenous type I IFN pathway has been observed by genomic analysis [17]. Microarray analysis of liver biopsies from HCVinfected patients has also revealed genes associated with activated lymphocytes, extracellular matrix, and macrophages, that could be possible markers for fibrosis progression[18,19]. In mice, expression of the HCV core protein results in down regulation in the expression of lipid metabolism-associated genes [20]. Other virus-host interactions, including SARS-CoV [21], influenza virus [22], and Ebola [23], have also been successfully studied by functional genomics. Gene expression profiling using microarray technology clearly makes it possible to obtain global transcriptional changes associated with a diseased state caused by many factors, which could help identify pathways implicated in progression of disease [24].

In the current report we utilized functional genomic analysis to compare gene expression patterns in cirrhotic livers, including various Child-Turcotte-Pugh (CTP)classifications, associated with either chronic alcohol consumption or HCV infection. The pathology of alcoholand HCV-induced cirrhosis is very similar and characteristic fibrosis patterns leading to cirrhosis can overlap among these two etiologies [25]. Therefore, it is particularly useful to now have the tools for genomic analysis, in order to determine how cirrhosis development may differ between these two distinct etiologies. Upon analysis, we indeed found differences in the expression of interferon-related genes and lymphocyte-specific genes between alcoholic and viral etiologies. Interestingly, a clear distinction existed between CTP classes of ALD, but not HCV-associated, cirrhosis. Different expression patterns depending on severity of alcohol-induced cirrhosis were found in genes associated with the inflammatory response, lipid metabolism, and oxidative stress. This information could be integral in increasing understanding of underlying mechanisms at work during end-stage liver disease.

Results

Global gene expression patterns distinguish ethanol- and HCV-induced cirrhosis

To analyze gene expression changes in cirrhotic livers associated with HCV (n = 8) or ethanol (n = 7), microarray experiments were performed comparing each type of cirrhotic liver to a reference pool of normal liver tissue. Cirrhosis severity was evaluated according to the CTP clinical scoring system (A-C, least to most severe, respectively) and patient scores are summarized in Table 1. The global gene expression profiles of all experiments are shown in Figure 1. In total, 2,965 genes were differentially expressed (\geq 2-fold, P \leq 0.05) in at least 3 of 15 samples. The clustering algorithm demonstrates that global gene expression profiles clearly distinguish ethanol- and HCVassociated cirrhotic samples (indicated with black and blue text, respectively). Substantially more gene expression changes were observed in HCV-related samples compared to ethanol-related, ranging from 574-1622 in HCV samples as compared to 79-683 in ethanol samples. Interestingly, the gene expression patterns observed were relatively uniform among HCV-cirrhotic samples and did not appear to differ between CTP classes. In contrast, the global gene expression profiles demonstrated distinct patterns among CTP classes in ethanol-related cirrhosis.

Gene expression changes specific to HCV-infected cirrhotic livers are associated with induction of the innate antiviral immune response

To further classify functional pathways specifically associated with etiology, we performed a one-way errorweighted ANOVA comparison of HCV- and ethanol-associated cirrhotic liver biopsies. A set of 946 genes was identified as being significantly ($P \le 0.05$) different with respect to expression level between the two groups. We then used Fatigo software to analyze functional groups that differed between cirrhosis etiologies. Genes associated with metabolism, the immune response, and cell death were differentially regulated in HCV- and alcoholrelated cirrhosis. Many of the genes specifically induced in HCV-associated cirrhotic livers were not up-regulated in ethanol-cirrhotic livers. This appears to be at least partly related to activation of the innate antiviral immune response, as many of these up-regulated genes are known to be regulated by IFN (Fig. 2A). Specifically, a subset of genes highly induced in HCV-associated cirrhotic samples were IFN α/β -inducible (IRF7, OAS2, GIP2, GIP3) or take

Patient #	Sex	Age	Etiology	Child-Pugh	Length of alcohol consumption	Quantity of alcohol consumed	ALT/AST
1239	М	68	Alcohol	А	several years	> 100 g/day	28/37
1240	М	70	Alcohol	А	several years	> 100 g/day	36/44
1245	М	67	Alcohol	А	unknown	unknown	19/19
1263	М	72	Alcohol	А	> 20 years	80 g/day	82/89
1164	М	62	Alcohol	В	> 30 years	> 80 g/day	33/27
1227	М	59	Alcohol	С	39 years active at OLT	> 100 g/day	28/54
1228	М	44	Alcohol	С	18 years	> 100 g/day	18/27
1015	М	47	HCV	А	N/A	none	96/45
1019	М	66	HCV	А	N/A	none	77/99
1022	М	61	HCV	А	N/A	none	113/143
1023	М	67	HCV	А	N/A	none	90/115
1037	F	54	HCV	В	N/A	none	57/78
1038	М	55	HCV	В	N/A	none	69/108
1035	М	59	HCV	С	N/A	none	87/100
1039	Μ	66	HCV	С	N/A	none	90/35

Table 1: Patient profile

part in its activation. Additional IFN-type I genes – OAS1, IFITM1, IFI35, IFIT1, MX1, IFI27 – were highly up-regulated in HCV-associated samples but not included in this ANOVA. A small set of genes (ISG20, IFI16, TRIM22, STAT5A) were also highly up-regulated in nearly all samples, including ethanol-cirrhotic livers, suggesting these may play a role in an inflammatory response related to cirrhosis development regardless of etiology.

Significant infiltration of lymphocytes into the liver is observed during chronic HCV infection [26]. In this study, up-regulation of gene markers for lymphocytes was indeed much more pronounced in HCV-infected liver samples than in ethanol-induced livers (Fig. 2B, Additional file 1). Specifically, RANTES, which is chemotactic for T cells *in vitro* was significantly up-regulated in all HCV-infected livers (1.8–4.5 fold, $P \ge 0.05$). In contrast, circulating levels of natural killer cells have been shown to be suppressed under the influence of alcohol [27,28]. Consistent with this, we observed decreased expression of natural killer cell-related genes, including CD44, IL15R, CD2 and CCL5, in alcohol-associated cirrhotic liver samples.

Gene expression patterns differentiate ethanol-associated cirrhosis by CTP class

Whereas HCV-associated cirrhotic samples looked very similar regardless of CTP class (Fig. 1), global gene expression profiles indicated that CTP class A ethanol-associated samples exhibited unique expression patterns from CTP class B and C samples. In order to further identify which gene expression patterns were specific depending on severity of ethanol-associated cirrhosis, we performed a one-way error-weighted ANOVA ($P \le 0.05$) of less severe (CTP class A) cirrhotic samples versus more severe (CTP classes B and C) samples (Fig. 3A, Additional file 2). Expression profiles distinguished the ethanol-related CTP

class A samples from ethanol-related CTP class B and C samples. In general, class A samples demonstrated up-regulation of more genes than in samples with more severe cirrhosis (Class B and C). Distinct gene expression profiles were not observed between CTP classes B and C, but this may be due to small sample size. Ingenuity analysis of groups I and II (Fig. 3A) demonstrated that the number of genes associated with various biological functions differed significantly depending on severity of alcohol cirrhosis. Of 135 annotated genes up-regulated in group II, 60 were associated with cell death as compared to only 16 in group I. High numbers of genes related to cell movement, mainly of phagocytes, and tissue development were also up-regulated in more severe ethanol cirrhosis. Many genes in group I (less severe cirrhosis) were associated with lipid metabolism, specifically the metabolism of cholesterol. Genes involved in the function of the immune and lymphatic system, such as activation of mononuclear leukocytes and proliferation of macrophages, were also up-regulated in early, less severe cirrhosis.

A number of genes that were nearly exclusively up-regulated in the CTP-A class of ethanol cirrhosis included inflammatory-related genes (Fig. 3B). Some of these – VIPR1, CCL15, PLG, CCR3, CSF1 – are involved in macrophage activation and migration. Macrophage-related genes not in this ANOVA set but also highly up-regulated in CTP class A samples included FN1, C5, LPA, and SERPINA3.

Interestingly, severe ethanol cirrhosis differed from less severe ethanol cirrhosis in expression of genes associated with fatty and bile acid metabolism (Fig. 3C). Many of these genes were up-regulated during less severe cirrhosis but down-regulated by end-stage ethanol cirrhosis. Sterol 27-hydroxylase (CYP27A1) oxidizes cholesterol in the



Figure I

Global gene expression analysis of liver tissue from HCV and ethanol-associated cirrhosis. 2-D hierarchical clustering was performed using Resolver System software with agglomerative algorithm, average link heuristic criteria, and Euclidian correlation metric. Each column represents gene expression data from an individual experiment comparing cirrhotic and normal liver and the cluster represents 2965 genes that showed a >2-fold change (P value < 0.05) in at least 3 of 15 experiments. A full view is shown with green and red colors showing decreased or increased levels, respectively, of mRNA expression relative to normal (noncirrhotic, uninfected) tissue. HCV-infected samples are highlighted in blue.

alternative pathway of bile acid biosynthesis; a defect in this enzyme leads to excessive accumulation of sterols [29]. We observed significant up-regulation of this gene in livers of all patients with early ethanol cirrhosis and, as with other genes in this group, down-regulation during late (end-stage) cirrhosis (Figure 3C). Fatty acids are substrates for cytochrome CYP2E1 (P450 2E1) [5], which is a key enzyme in the microsomal ethanol oxidation system and plays a role in alcohol detoxification and nutritional support [30]. It is also a major source of oxidative stress [6,5], and both protein and mRNA levels increase dramat-



Figure 2

Differences in global gene expression profiles between HCV and ethanol-associated cirrhotic liver tissue. A. Expression of genes associated with IFN response. HCV- and EtOH-associated cirrhosis samples were clustered based on the expression of IFN-related genes that were at least 2-fold regulated with a P value \leq 0.05 in at least 3 of 15 experiments, comparing cirrhotic tissue to a normal liver reference. See legend of Fig. 1 for details on clustering, the applied color scheme for the fold changes in the mRNA levels, and experiment representation. B. Expression of genes associated with lymphocyte activation and proliferation. Cirrhotic samples were again clustered based on the expression of lymphocyte-related genes that were at least 2-fold regulated with a P value \leq 0.05 in at least 3 of 15 experiments comparing cirrhotic tissue to a normal liver reference. See legend of Fig. I for details on clustering, the applied color scheme for the fold changes in the mRNA levels, and experiment representation. See Supplementary Table 1 for a list of gene names represented in Figure 2B.

ically in actively drinking patients [31]. Therefore it was interesting that significant down-regulation of this gene was also observed in high grade (CTP class B and C) cirrhotic livers (Figure 3C). A similar pattern was observed for genes encoding alcohol dehydrogenases (ADH4, ADH1B, ADH1C).



Figure 3

Differences in global gene expression profiles between different CTP classes of ethanol-associated cirrhotic liver tissue. A. Expression profiles of genes differentially expressed in ethanol-related cirrhosis CTP class A versus CTP BC (identified from one-way ANOVA). Each column represents gene expression data from an individual experiment comparing cirrhotic and normal liver and the cluster represents genes that showed a >2-fold change (P value < 0.05) in at least 3 of 7 experiments. See legend of Fig. 1 for details on clustering, the applied color scheme for the fold changes in the mRNA levels, and experiment representation. Grouping (I and II) is discussed in the text and demonstrates differences between CTP classes in gene expression of genes associated with various biological processes. Refer to Supplementary Table 2 for gene names in each group. **B.** Expression of genes associated with the inflammatory response. Cluster depicts expression of genes that were at least 2-fold regulated with a *P* value \leq 0.05 in at least 3 of 7 experiments comparing cirrhotic tissue to a normal liver reference. See legend of Fig. 1 for details on clustering, the applied color scheme for the fold changes in the mRNA levels, and experiment representation **C**. Expression of genes involved in fatty acid and bile acid metabolism. Expression of genes that were at least 2-fold regulated with a *P* value \leq 0.05 in at least 3 of 15 experiments comparing cirrhotic tissue to a normal liver reference. See legend of Fig. 1 for details on clustering, the applied color scheme for the fold changes in the mRNA levels, and experiment representation **C**. Expression of genes involved in fatty acid and bile acid metabolism. Expression of genes that were at least 2-fold regulated with a *P* value \leq 0.05 in at least 3 of 15 experiments comparing cirrhotic tissue to a normal liver reference. See legend of Fig. 1 for details on clustering, the applied color scheme for the fold changes in the mRNA levels, and experiment representation.

Differential gene expression in ethanol-induced class A cirrhosis

CTP class A alcohol-associated samples appeared to have unique expression profiles compared to all other samples regardless of etiology (Fig. 1). We therefore performed a one-way error-weighted ANOVA ($P \ge 0.05$) of less severe (CTP class A alcohol-associated) versus more severe ethanol-related and HCV-associated cirrhotic samples to identify genes with expression levels unique to CTP class A alcohol. Of 607 total genes in this set, a cluster of 273 were highly up-regulated in all samples but ethanol CTP class A (Fig. 4, group I), and a cluster of 109 were highly up-regulated specifically in class A (Fig. 4, group II); see Additional file 3 for more information on gene names. Functional analysis (Ingenuity) revealed that genes associated with cell death, movement, and the immune response were much more abundant in group I than in group II. Most immune response genes up-regulated in group I were generally involved in proliferation and migration of leukocytes, while those in group II were again related to activation and infiltration of macrophages. The majority of genes up-regulated in group II of early ethanol cirrhosis are involved in the modification of lipids. The response to oxidative stress was also significantly different in early ethanol cirrhosis as related to all other samples. Two genes in particular were highly up-regulated in CTP class A livers - PRDX2 (up to 4.7 fold) and APOE (up to 17.5 fold) - and either less up-regulated or even down-regulated in the majority of the remaining samples.

Discussion

Global gene expression profiling of liver biopsy material revealed significant differences between HCV and ethanol-induced cirrhosis. In general, gene expression changes in cirrhotic livers are more numerous in HCV-infected livers compared to alcohol-induced diseased livers. Gene expression patterns can also distinguish classes of ethanol-associated cirrhosis. Additionally, unique differences in gene expression existed between early ethanol cirrhosis and all other liver samples.

The majority of these differences between HCV and ethanol etiologies involved the expression of genes associated with the immune response, including the IFN response and infiltration of immune cells. This suggests that the immune response may play a more prominent role in viral versus ethanol-induced cirrhosis. Our data showing significant up-regulation of IFN-type I genes specifically in HCV-associated samples are also consistent with previous studies that demonstrated activation of innate antiviral signaling pathways during HCV infections in chimps [32] and humans [17,18,26,33-37].



Figure 4

Differences in global gene expression profiles between ethanol-induced (Class A) cirrhotic tissue versus HCV (Class A, B, C) and ethanol (Class B and C)-induced cirrhotic tissue. Expression profiles of genes differentially expressed in early ethanol-related cirrhosis versus all other samples; as identified by ANOVA. Each column represents gene expression data from an individual experiment comparing cirrhotic and normal liver and the cluster represents genes that showed a >2-fold change (P value < 0.05) in at least 3 of 15 experiments. See legend of Fig. 1 for details on clustering and the applied color scheme. Refer to Supplementary Table 3 for names of genes represented in this figure.

In this report, gene markers for lymphocytes were also very pronounced in HCV-infected livers. Increased expression of RANTES – chemotactic for T cells – has been observed in livers of chronically-infected HCV patients [38] and a correlation has been found between increased expression of RANTES in the liver and the extent of liver damage [39]. That we observed significant up-regulation of genes related to lymphocyte infiltration exclusively in HCV-infected cirrhotic livers may have some significance for how liver injury differs in HCV- and ethanol-cirrhosis. It has already been shown that liver injury during chronic HCV infection is associated with increased expression of TH1 cytokines [40,41]. However, T-cell activation is also seen in patients during chronic alcohol consumption [42]. The present study points to a predominant role of the immune response in liver pathogenesis well into the latest stages of HCV-associated disease, and T cells seem to be important in this response. Though the activation of T cells appears to contribute to liver injury in chronic ALD, our observations suggest that the immune response could be more important in the stages before cirrhosis develops, but no longer predominant during cirrhosis progression.

Significant differences in gene expression patterns between classes of ethanol-associated liver cirrhosis point to the role of macrophages in progression of ethanolinduced liver disease. Macrophages and other cells of the innate immune system have already been implicated in liver injury during alcoholic liver disease [43-47,12-14]. Induction of cytokine production by Kupffer cells activates hepatic stellate cells, leading to excess accumulation of extracellular matrix and fibrosis development. Production of cytokines and reactive oxygen species from monocytes has been shown to increase in response to chronic alcohol intake but decrease with acute alcohol intake [43]. We found an increase in the expression of macrophagerelated genes in early less severe (CTP A) ethanol cirrhosis, but not in late more severe (CTP B and C) ethanol cirrhosis. Though sample number does not allow for strictly defining less or more severe cirrhotics, the trend observed may have some significance related to regulation of metabolism.

Genes specifically linked to metabolism also demonstrated differential regulation depending on cirrhosis stage in alcoholic liver disease. The metabolism of alcohol leads to fat accumulation and oxidative stress which, through the activation of hepatic stellate cells, eventually leads to liver fibrosis [48]. As the CYP system is the most important enzyme system for drug metabolism, it is interesting that several genes encoding these proteins were upregulated only in livers exhibiting early ethanol cirrhosis. CYP27A1, an oxidizer of cholesterol, and CYP2E1, a key enzyme in the microsomal ethanol oxidation system, were both up-regulated in most early (CTP class A) cirrhosis samples but significantly down-regulated by more severe, end-stage cirrhosis. As CYP27A1 is an acute phase gene repressed by lipopolysaccharide and cytokines, it is possible that the down-regulation of this gene is linked to levels of cytokine activity during late-stage ethanol cirrhosis [49].

We know that Kupffer cells harbor CYP2E1 [50], become active in ALD, and produce inflammatory cytokines [4]. It could therefore be consistent that we observe both an increase in expression of macrophage-related genes as well as genes involved in cholesterol oxidation at the same (early) stages of cirrhosis. Macrophages may be an important source for lipid metabolism enzyme activity at the beginning stages of cirrhosis, but become less necessary as there is decreased demand for active metabolism of alcohol. Again, this is only speculation as we realize that sample size is limiting our ability to definitively group CTP class B with CTP class C as late/severe cirrhotics. Alternatively, this down-regulation of genes during later stages of ethanol cirrhosis may reflect widespread cell death and shut-down of liver function.

It is interesting that there were no significant changes in gene expression patterns observed between classes of HCV-induced cirrhotic livers. This may be evidence for a uniform host response throughout cirrhosis development in this etiology. However, it is more likely that the similarities seen among HCV-infected livers are due to the antiviral host response to HCV infection.

Conclusion

It is clear that gene expression profiles in ethanol-induced and HCV-induced liver cirrhosis are distinct. We have identified functional groups that differ both between etiologies and within ethanol-induced cirrhosis, depending on the extent of liver damage. The data suggest that oxidative stress, lipid metabolism, and macrophage activity may be more associated with severe liver injury leading to cirrhosis in ALD than in viral-induced disease. In contrast, the adaptive immune response, characterized by T cellmediated killing of HCV-infected cells, appears to be more associated with the development of liver cirrhosis in viral-, but not alcohol-mediated disease. The study supports the idea that though these different etiologies lead to similar pathologies, the cellular mechanisms responsible for disease progression may be unique. Although we realize the limitations of a small cross-sectional study, the gene expression patterns identified here are from a large set of genes in order to give a broad global perspective. These initial findings will contribute to further studies to distinguish which pathways can differentiate development of liver damage due to etiology, which could lead to better-targeted therapies for end-stage liver disease.

Methods

Human Liver Tissue Samples

Cirrhotic liver samples from 7 alcoholic and 8 HCVinfected patients were obtained from liver disease patients at the University of Barcelona, Spain who underwent liver transplant or hepatic resection. None of the HCV-infected patients in this study were undergoing anti-viral treatment at the time the liver sample was obtained. Alcohol consumption information was based on data obtained in patient interviews. The severity of cirrhosis was clinically graded according to Child-Turcotte-Pugh (CTP) scores (summarized in Table 1). Normal, uninfected liver tissue from 8 patients who underwent tumor resection due to metastases of non-liver origin was collected at the same clinic (non-tumor material was used). These uninfected samples were pooled to create a standard normal liver reference (pNL) that was used for all microarray experiments. All patients gave informed prior consent to protocols approved by the University of Barcelona, Spain.

Total RNA Isolation and mRNA Amplification

Frozen liver tissues were disrupted in Trizol Reagent (Invitrogen, Carlsbad, CA) using a Polytron homogenizer (PowerGene 700, Fisher Scientific), and total RNA was isolated according to the Trizol protocol. All total RNA samples were double amplified using the RiboAmp RNA Amplification kit (Arcturus, Mountain View, CA). The quality of amplified RNA was evaluated by capillary electrophoresis using an Agilent 2100 Bioanalyzer (Agilent Technologies, Palo Alto, CA).

Expression Microarray Format and Data Analysis

Microarray format, protocols for probe labeling, hybridization, slide treatment, and scanning were performed as previously described [35] and are also available at our website [51]. Human cDNA set 1 expression arrays carrying 13,026 unique cDNA clones were purchased from Agilent. A single experiment comparing two samples, one from a liver with infection and one from the normal liver reference, was performed with four replicate arrays using the dye label reverse technique as described before, thus providing mean ratios between the expression levels of each gene in the analyzed sample pair, standard deviations, and P values [35,52-54]. All data were entered into a custom-designed database, Expression Array Manager, and then uploaded into Rosetta Resolver System 4.0 (Rosetta Biosoftware, Kirkland, WA) and Spotfire Decision Suite 7.1.1 (Spotfire, Somerville, MA). Data normalization and the Resolver System Error model specifically developed for this array format are described on the web site [51]. This web site is also used to publish all primary data in accordance with the proposed standards [55]. The annotation sites Fatigo [56] and Ingenuity [57] were used to classify genes into functional groups.

Additional material

Additional File 1

"Genes differentially regulated in Figure Figure 2B" The table lists gene names for heat map shown in Figure 2B.

Click here for file

[http://www.biomedcentral.com/content/supplementary/1743-422X-3-98-S1.xls]

Additional File 2

"Goup I and II gene names for Figure 3A" The table list gene names for biological function groups in Figure 3A. Click here for file [http://www.biomedcentral.com/content/supplementary/1743-422X-3-98-S2.xls]

Additional File 3

"Group I and II gene names for Figure 4" The table lists gene names for groups differentiating cirrhosis class in Figure 4. Click here for file [http://www.biomedcentral.com/content/supplementary/1743-422X-3-98-S3.xls]

Acknowledgements

We thank Dr. Maria Smith for technical and analytical assistance. Financial support by Public Health Service grants R01CA074131, U19Al48214, P30DA01562501, R37DA004334, R01DA12568, and R01DA16078 from the National Institutes of Health.

References

- Koike K: Hepatitis C as a metabolic disease: Implication for the pathogenesis of NASH. Hepatol Res 2005, 33(2):145-150.
- 2. Alter HJ, Seeff LB: Recovery, persistence, and sequelae in hepatitis C virus infection: a perspective on long-term outcome. Semin Liver Dis 2000, 20:17-35.
- Hines IN, Wheeler MD: Recent advances in alcoholic liver disease III. Role of the innate immune response in alcoholic hepatitis. Am J Physiol Gastrointest Liver Physiol 2004, 287:G310-G314.
- Saito H, Ishii H: Recent understanding of immunological aspects in alcoholic hepatitis. *Hepatol Res* 2004, 30:193-198.
- 5. Lieber CS: Alcoholic fatty liver: its pathogenesis and mechanism of progression to inflammation and fibrosis. Alcohol 2004, **34**:9-19.
- Nanji AA: Role of different dietary fatty acids in the pathogenesis of experimental alcoholic liver disease. Alcohol 2004, 34:21-25.
- Polavarapu R, Spitz DR, Sim JE, Follansbee MH, Oberley LW, Rahemtulla A, Nanji AA: Increased lipid peroxidation and impaired antioxidant enzyme function is associated with pathological liver injury in experimental alcoholic liver disease in rats fed diets high in corn oil and fish oil. *Hepatology* 1998, 27:1317-1323.
- 8. Sies H: Strategies of antioxidant defense. Eur J Biochem 1993, 215:213-219.
- 9. Thiele GM, Freeman TL, Klassen LW: Immunologic mechanisms of alcoholic liver injury. *Semin Liver Dis* 2004, **24:**273-287.
- Laso FJ, Iglesias-Osma C, Ciudad J, Lopez A, Pastor I, Torres E, Orfao A: Alcoholic liver cirrhosis is associated with a decreased expression of the CD28 costimulatory molecule, a lower ability of T cells to bind exogenous IL-2, and increased soluble CD8 levels. Cytometry 2000, 42:290-295.
- 11. Pruett SB, Schwab C, Zheng Q, Fan R: Suppression of innate immunity by acute ethanol administration: a global perspective and a new mechanism beginning with inhibition of signaling through TLR3. J Immunol 173:2715-2724. 8-15-2004
- Afford SC, Fisher NC, Neil DA, Fear J, Brun P, Hubscher SG, Adams DH: Distinct patterns of chemokine expression are associated with leukocyte recruitment in alcoholic hepatitis and alcoholic cirrhosis. J Pathol 1998, 186:82-89.
- Bautista AP: Impact of alcohol on the ability of Kupffer cells to produce chemokines and its role in alcoholic liver disease. J Gastroenterol Hepatol 2000, 15:349-356.
- Maltby J, Wright S, Bird G, Sheron N: Chemokine levels in human liver homogenates: associations between GRO alpha and histopathological evidence of alcoholic hepatitis. *Hepatology* 1996, 24:1156-1160.

- Mosher B, Dean R, Harkema J, Remick D, Palma J, Crockett E: Inhibition of Kupffer cells reduced CXC chemokine production and liver injury. *J Surg Res* 2001, 99:201-210.
 Saadeh S, Cammell G, Carey WD, Younossi Z, Barnes D, Easley K:
- Saadeh S, Cammell G, Carey WD, Younossi Z, Barnes D, Easley K: The role of liver biopsy in chronic hepatitis C. Hepatology 2001, 33:196-200.
- Bieche I, Asselah T, Laurendeau I, Vidaud D, Degot C, Paradis V, Bedossa P, Valla DC, Marcellin P, Vidaud M: Molecular profiling of early stage liver fibrosis in patients with chronic hepatitis C virus infection. Virology 2005, 332:130-144.
- Smith MW, Walters K-Ä, Korth MJ, Fitzgibbon M, Proll SC, Thompson JC, Yeh MM, Shuhart MC, Furlong JC, Cox PP, Thomas DL, Phillips JD, Kushner JP, Fausto N, Carithers RL, Katze MG: Gene expression patterns that correlate with hepatitis C and early progression to fibrosis in liver transplant recipients. *Gastroenterol* 2006, 130:179-187.
- Walters KA, Joyce MA, Thompson JC, Smith MW, Yeh MM, Proll S, ZhuLin-Fu Gao TJ, Knetman NM, Tyrrell DL, Katze MG: Host-Specific Response to HCV Infection in the Chimeric SCID-beige/ Alb-uPA Mouse Model: Role of the Innate Antiviral Immune Response. PLOS Pathogens 2:591-601. 6-23-2006
- Yamaguchi A, Tazuma S, Nishioka T, Ohishi W, Hyogo H, Nomura S, Chayama K: Hepatitis C virus core protein modulates fatty acid metabolism and thereby causes lipid accumulation in the liver. *Dig Dis Sci* 2005, **50**:1361-1371.
- Baas T, Taubenberger J, Chong PY, Chui P, Katze MG: SARS-CoV virus-host interactions and comparative etiologies of acute respiratory distress syndrome as determined by transcriptional and cytokine profiling of formalin-fixed paraffinembedded tissues. J Interferon Cytokine Res 2006, 26:309-317.
- Korth MJ, Kash JC, Baskin CR, Katze MG: Insights into influenza virus-host interactions through global gene expression profiling: cell culture systems to animal models. 2006;341-360.
- Kash JC, Mühlberger É, Carter V, Grosch O, Perwitasari O, Proll SC, Thomas MJ, Weber F, Klenk H.-D, Katze MG: Global suppression of the host antiviral response by Ebola- and Marburgviruses: increased antagonism of the type I interferon response is associated with enhanced virulence. J Virol 2006, 80:3009-3020.
- 24. Korth MJ, Kash JC, Furlong JC, Katze MG: Virus infection and the interferon response: a global view through functional genomics. *Methods Mol Med* 2005, 116:37-55.
- 25. Uchimura Y, Sata M, Kage M, Abe H, Tanikawa K: A histopathological study of alcoholics with chronic HCV infection: comparison with chronic hepatitis C and alcoholic liver disease. *Liver* 1995, **15**:300-306.
- Nuti S, Rosa D, Valiante NM, Saletti G, Caratozzolo M, Dellabona P, Barnaba V, Abrignani S: Dynamics of intra-hepatic lymphocytes in chronic hepatitis C: enrichment for Valpha24+ T cells and rapid elimination of effector cells by apoptosis. Eur J Immunol 1998, 28:3448-3455.
- 27. Laso FJ, Madruga JI, Lopez A, Ciudad J, varez-Mon M, San Miguel J, Orfao A: Distribution of peripheral blood lymphoid subsets in alcoholic liver cirrhosis: influence of ethanol intake. Alcohol Clin Exp Res 1996, 20:1564-1568.
- Chedid A, Mendenhall CL, Moritz TE, French SW, Chen TS, Morgan TR, Roselle GA, Nemchausky BA, Tamburro CH, Schiff ER: Cellmediated hepatic injury in alcoholic liver disease. Veterans Affairs Cooperative Study Group 275. Gastroenterology 1993, 105:254-266.
- 29. Chiang JY: Regulation of bile acid synthesis: pathways, nuclear receptors, and mechanisms. J Hepatol 2004, 40:539-551.
- Koop DR, Tierney DJ: Multiple mechanisms in the regulation of ethanol-inducible cytochrome P450IIE1. Bioessays 1990, 12:429-435.
- Takahashi T, Lasker JM, Rosman AS, Lieber CS: Induction of cytochrome P-4502E1 in the human liver by ethanol is caused by a corresponding increase in encoding messenger RNA. *Hepatology* 1993, 17:236-245.
- Bigger CB, Brasky KM, Lanford RE: DNA microarray analysis of chimpanzee liver during acute resolving hepatitis C virus infection. J Virol 2001, 75:7059-7066.
- Koziel MJ, Dudley D, Wong JT, Dienstag J, Houghton M, Ralston R, Walker BD: Intrahepatic cytotoxic T lymphocytes specific for hepatitis C virus in persons with chronic hepatitis. J Immunol 1992, 149:3339-3344.

- Cerny A, Chisari FV: Pathogenesis of chronic hepatitis C: immunological features of hepatic injury and viral persistence. *Hepatology* 1999, 30:595-601.
- Smith MW, Yue ŽN, Korth MJ, Do HA, Boix L, Fausto N, Bruix J, Carithers RL Jr, Katze MG: Hepatitis C virus and liver disease: global transcriptional profiling and identification of potential markers. Hepatology 2003, 35:1458-1467.
- Mueller T, Mas-Marques A, Sarrazin C, Wiese M, Halangk J, Witt H, Ahlenstiel G, Spengler U, Goebel U, Wiedenmann B, Schreier E, Berg T: Influence of interleukin 12B (IL12B) polymorphisms on spontaneous and treatment-induced recovery from hepatitis C virus infection. J Hepatol 2004, 41:652-658.
- Nascimbeni M, Shin EC, Chiriboga L, Kleiner DE, Rehermann B: Peripheral CD4(+)CD8(+) T cells are differentiated effector memory cells with antiviral functions. *Blood* 2004, 104:478-486.
- Leroy V, Vigan I, Mosnier JF, Dufeu-Duchesne T, Pernollet M, Zarski JP, Marche PN, Jouvin-Marche E: Phenotypic and functional characterization of intrahepatic T lymphocytes during chronic hepatitis C. Hepatology 2003, 38:829-841.
- 39. Kusano F, Tanaka Y, Marumo F, Sato C: Expression of C-C chemokines is associated with portal and periportal inflammation in the liver of patients with chronic hepatitis C. Lab Invest 2000, 80:415-422.
- McCaughan GW, Gorrell MD, Bishop GA, Abbott CA, Shackel NA, McGuinness PH, Levy MT, Sharland AF, Bowen DG, Yu D, Slaitini L, Church WB, Napoli J: Molecular pathogenesis of liver disease: an approach to hepatic inflammation, cirrhosis and liver transplant tolerance. *Immunol Rev* 2000, 174:172-191.
- Napoli J, Bishop GA, McGuinness PH, Painter DM, McCaughan GW: Progressive liver injury in chronic hepatitis C infection correlates with increased intrahepatic expression of Th1-associated cytokines. Hepatology 1996, 24:759-765.
- Cook RT: Alcohol abuse, alcoholism, and damage to the immune system – a review. Alcohol Clin Exp Res 1998, 22:1927-1942.
- Zhang Z, Bagby GJ, Stoltz D, Oliver P, Schwarzenberger PO, Kolls JK: Prolonged ethanol treatment enhances lipopolysaccharide/ phorbol myristate acetate-induced tumor necrosis factoralpha production in human monocytic cells. Alcohol Clin Exp Res 2001, 25:444-449.
- 44. Pritchard MT, Nagy LE: Ethanol-induced liver injury: potential roles for egr-1. Alcohol Clin Exp Res 2005, 29:146S-150S.
- Takeyama Y, Kamimura S, Kuroiwa A, Sohda T, Irie M, Shijo H, Okumura M: Role of Kupffer cell-derived reactive oxygen intermediates in alcoholic liver disease in rats in vivo. Alcohol Clin Exp Res 1996, 20:335A-339A.
- Neuman MG, Brenner DA, Rehermann B, Taieb J, Chollet-Martin S, Cohard M, Garaud JJ, Poynard T, Katz GG, Cameron RG, Shear NH, Gao B, Takamatsu M, Yamauchi M, Ohata M, Saito S, Maeyama S, Uchikoshi T, Toda G, Kumagi T, Akbar SM, Abe M, Michitaka K, Horiike N, Onji M: Mechanisms of alcoholic liver disease: cytokines. Alcohol Clin Exp Res 2001, 25:251S-253S.
- 47. Thurman RG: II. Alcoholic liver injury involves activation of Kupffer cells by endotoxin. *Am J Physiol* 1998, **275**:G605-G611.
- Bataller R, Sancho-Bru P, Gines P, Brenner DA: Liver fibrogenesis: a new role for the renin-angiotensin system. Antioxid Redox Signal 2005, 7:1346-1355.
- 49. Memon RA, Moser AH, Shigenaga JK, Grunfeld C, Feingold KR: In vivo and in vitro regulation of sterol 27-hydroxylase in the liver during the acute phase response. potential role of hepatocyte nuclear factor-1. J Biol Chem 2001, 276:30118-30126.
- Koivisto T, Mishin VM, Mak KM, Cohen PA, Lieber CS: Induction of cytochrome P-4502E1 by ethanol in rat Kupffer cells. Alcohol Clin Exp Res 1996, 20:207-212.
- 51. Expression Array Manager [http://expression.microslu.washing ton.edu/expression/index.html]
- Geiss GK, An MC, Bumgarner RE, Hammersmark E, Cunningham D, Katze MG: Global impact of influenza virus on cellular pathways is mediated by both replication-dependent and -independent events. J Virol 2001, 75:4321-4331.
 Smith MW, Yue ZN, Geiss GK, Sadovnikova NY, Carter VS, Boix L,
- Smith MW, Yue ZN, Geiss GK, Sadovnikova NY, Carter VS, Boix L, Lazaro CA, Rosenberg GB, Bumgarner RE, Fausto N, Bruix J, Katze MG: Identification of novel tumor markers in hepatitis C virus-associated hepatocellular carcinoma. *Cancer Res* 2003, 75:859-864.

- Geiss GK, Carter VS, He Y, Kwieciszewski BK, Holzman T, Korth MJ, Lazaro CA, Fausto N, Bumgarner RE, Katze MG: Gene expression profiling of the cellular transcriptional network regulated by alpha/beta interferon and its partial attenuation by the hepatitis C virus nonstructural 5A protein. J Virol 2003, 77:6367-6375.
- 55. Brazma A, Hingamp P, Quackenbush J, Sherlock G, Spellman P, Stoeckert C, Aach J, Ansorge W, Ball CA, Causton HC, Gaasterland T, Glenisson P, Holstege FCP, Kim IF, Markowitz V, Matese JC, Parkinson H, Robinson A, Sarkans U, Schulze-Kremer S, Stewart J, Taylor R, Vilo J, Vingron M: Minimum information about a microarray experiment (MIAME) toward standards for microarray data. Nature Genet 2001, 29:365-371.
- 56. Fatigo [http://fatigo.bioinfo.cipf.es]
- 57. Ingenuity Systems [http://ingenuity.com]

