Abstract OC-029 Figure 1

0.0001) [Figure i]. HE grade improved significantly following therapy (p < 0.0001) [Figure ii].

Child Pugh score fell significantly following therapy (p < 0.0001) [Figure iii], as did MELD and UKELD scores: 15 ± 7 vs. 13 ± 5 (p = 0.03) and 55 ± 6 vs. 51 ± 5 (p < 0.02), respectively. This is noteworthy as the MELD score does not include HE as a parameter and is based on bilirubin, INR and creatinine.

Conclusion Our UK multi-centre experience is that rifaximin is well-tolerated and an efficacious treatment for the secondary prevention of HE. Rifaximin significantly reduced both hospital re-admission rates after 3 months treatment, impacting significantly on the NHS resource burden of HE, and reduced overall liver disease severity raising the possibility that its therapeutic effect may extend beyond reducing gut ammonia production.


OC-030 EFFECTIVE STRATIFICATION OF HEPATOCELLULAR CARCINOMA RISK IN PRIMARY BILIARY CIRRHOSIS: RESULTS OF A MULTI-CENTRE INTERNATIONAL STUDY

1A Trivedi,1 W Lammas,1 H van Buuren,1 H Jansen,1 P Invernielli,1 PM Battezzati,1 A Floreani,1 A Pares,1 C Ponsioen,1 C Coppechot,1 B Poupop,1 M Mayo,1 T Talwalkar,1 I A Burroughs,1 I N V eros,1 A Mason,1 T Bruns,1 K K U,1 K Kwédy,1 K Yamashita,1 A Cheung,1 A Lie,1 N Cazagon,1 F Franceschet,1 C Caballera,1 K Boons,1 T de Vries,1 M Imam,1 G Piet,1 I P Kanwar,1 K Under,1 B Hansen,1 G Hirschfeld on behalf of Global PBC Study Group, 1NIHR Biomedical Research Unit and Centre for Liver Research, University of Birmingham, Birmingham, UK; 2Department of Gastroenterology and Hepatology, Erasmus Medical Centre, Rotterdam, The Netherlands; 3Liver Clinic, Toronto Western and General Hospital, University Health Network, Toronto, Canada; 4Liver Unit and Center for Autoimmune Liver Diseases, Humanitas Clinical and Research Center, Rozzano, Italy; 5Department of Health Sciences, Università Degli Studi Di Milano, Milano, Italy; 6Department of Surgical, Oncological and Gastroenterological, University of Padua, Padua, Italy; 7Liver Unit, Hospital Clinic, CIBERReh, IDIBAPS, University of Barcelona, Barcelona, Spain; 8Department of Gastroenterology and Hepatology, Academic Medical Center, Amsterdam, The Netherlands; 9Centre de Référence Des Maladies Inflammatoires Des Voies Biliaires, Hôpital Saint-Antoine, Paris, France; 10Digestive and Liver Diseases, UT Southwestern Medical Center, Dallas, USA; 11Department of Gastroenterology and Hepatology, Mayo Clinic, Rochester, USA; 12The Sheila Sherlock Liver Centre, The Royal Free Hospital, London, UK; 13Department of Hepatology, University Hospitals Leuven, Leuven, Belgium; 14Division of Gastroenterology and Hepatology, University of Alberta, Edmonton, Canada; 15Department of Internal Medicine IV, Jena University Hospital, Friedrich Schiller University Jena, Jena, Germany; 16Liver Center of Excellence, Digestive Disease Institute, Virginia Mason Medical Center, Seattle, USA; 17Liver Clinic, Toronto Western and General Hospital, Toronto, Canada; 18Liver Unit and Center for Autoimmune Liver Diseases, Humanitas Clinical and Research Center, Rozzano, Italy; 19Department of Scienze Chirurgiche Oncologiche E Gastroenterologiche, Università Degli Studi Di Milano, Milano, Italy; 20Department of Surgical, Oncological and Gastroenterological, University of Padua, Padua, Italy; 21Primary Healthcare Centre Premià de Mar, Catalonia

Introduction Hepatocellular carcinoma (HCC) is an important but infrequent outcome in primary biliary cirrhosis (PBC). Improved risk evaluation is an important goal for stratified surveillance.

Methods Risk-factor analysis of the ‘Global PBC Study Group’ comprising 15 centres across North America and Europe spanning >40-years follow-up was performed using Cox proportional hazards model, logistic regression and Kaplan-Meier estimates (SPSSv21).

Results Of 3546 patients with PBC (med. follow-up 8.6 yrs; IQR 4.4–14.1), 131 developed HCC. Excluding those who developed HCC within 12 months of PBC diagnosis (n = 23), median time to HCC was 12.7 yrs (6.9–16.8) and subsequent survival 1.1 yrs. (0.2–7.7). At diagnosis, factors associated with HCC development were male gender (adj. HR: 4.1;1.4–12.2, p = 0.014) and thrombocytopenia (adj. HR: 4.5;1.4–14.8, p = 0.012). Use of ursoodeoxycholic acid per-se was not associated with future risk of HCC, but stratification of risk by biochemical response at 12 months was effective by Rotterdam (adj. HR: 8.9;2.1–37.3, P = 0.003), Paris-I (adj. HR: 7.6;2.0–29.0, p = 0.003) or Toronto criteria (HR: 5.6;1.6–18.8, P = 0.006). Five (4.6 vs. 0.2%) and 10-year (13%/vs.1.9%) HCC incidence was significantly increased for biochemical non-responders (p = 2.2 × 10⁻³), and by multivariate analysis non-response remained the only significant risk factor.

Conclusion Our uniquely powered cohort allows robust demonstration that 12-month biochemical non-response is associated with an increased risk of developing HCC in PBC. Routine surveillance in those achieving biochemical response is unlikely cost-effective.

Disclosure of Interest None Declared.

OC-031 RELAXIN MODULATES CIRRHOSIS-INDUCED RENAL VASCULAR ENDOTHELIAL DYSFUNCTION

1VK Snowdon*,2 VWF Hadoke,1 NV Mungall,1 A Thomsen,1 T Kendall,1 D Webb,1 P Heddle,1 JA Fallowfield,1 MRC Centre for Inflammation Research, University of Edinburgh, Edinburgh, UK; 2BHF University of Edinburgh Centre for Cardiovascular Science, Edinburgh, UK; 3Biomedical Research Resources, University of Edinburgh, Edinburgh, UK

Introduction Hepatorenal syndrome (HRS) is a feared complication of cirrhosis characterised by intense renal vasoconstriction. The pathophysiology remains unclear, pharmacotherapy is limited and mortality is high. We investigated vascular responsiveness and the pathogenesis of renal vasoconstriction in models of advanced rat cirrhosis. Additionally, we determined the mechanism of action of the vasoactive peptide relaxin (RLN), previously shown to increase renal blood flow (RBF) in experimental cirrhosis (Snowdon V et al., BSG 2013).

Methods We induced cirrhosis, reduced RBF and renal dysfunction in male SD rats by carbon tetrachloride (16 wk) or bile duct ligation (4 wk). Arteries from renal (renal, segmental, interlobar) and splanchnic circulation were isolated for functional assessment using wire myography. qPCR array for vasoactive signalling genes,
western blot for eNOS signalling proteins and NO activity assay were undertaken in cirrhotic and control kidneys. Markers of oxidative stress and inflammatory cytokines were measured in serum by ELISA. We studied the effects of s.c. infusion of recombinant human RN (seralaxin; 72 h, 4 μg/h) on these parameters. Doppler ultrasound measured changes in cardiac output (CO) and renal arterial resistive index (RRI) in response to i.v. RLN (4 μg). Kidney endothelial morphology was assessed by electron microscopy, H+E and PAS stained kidney by light microscopy.

**Results** In renal arteries from control and cirrhotic rats endothelial vasodilation was eNOS-dependent. In cirrhotic rats endothelium-dependent relaxation (acetylcholine; 10−9–3 × 10−5 M) was dramatically reduced (p < 0.0001) in all renal arteries; with only a modest reduction seen in the mesenteric arteries. Endothelium-independent relaxation (sodium nitroprusside; 10−9–3 × 10−5 M) and vasoconstriction (phenylephrine; 10−9–3 × 10−5 M) were unaltered. In cirrhotic kidneys, total eNOS expression was up-regulated, as were arginase2 and caveolin1 (negative regulators of eNOS), and NOS activity was reduced (p < 0.05). Acute RLN had no effect on CO but decreased RRI (p < 0.05). Extended RLN restored endothelium-dependent relaxation, increased kidney NOS activity (p < 0.05), increased phosphorylated Akt and eNOS, and reduced serum TNFα levels.

**Conclusion** Renal vascular endothelial dysfunction characterises experimental cirrhosis, through a reduction in renal eNOS activity. This impairment may contribute to the renal vasoconstriction seen in cirrhosis and is a promising target for therapeutic modulation. RLN treatment restored renal endothelial vasodilatation. The potential for recombinant forms of RLN as a haemodynamic modulator in human cirrhosis and HRS merits investigation in translational studies.

**Disclosure of Interest** None Declared.

**OC-032** SURVEILLANCE LEADS TO IMPROVED OUTCOMES FOR PATIENTS WITH HEPATOCELLULAR CARCINOMA (HCC)

1TJS Cross*, 2W Ding, 1PD Richardson, 2F Yousel, 2D Palmer, 2E Jeeves, 3F Farrell, 1J Evans.
1Department of Gastroenterology and Hepatology, The Royal Liverpool Hospital, Liverpool, UK; 2Academic Department of Oncology, The University of Liverpool, Liverpool, UK; 3Department of Radiology, The Royal Liverpool Hospital, Liverpool, UK

10.1136/gutjnl-2014-307263.32

**Introduction** 6-monthly ultrasound surveillance is recommended in cirrhotic patients at risk of HCC. The benefit of surveillance has never been demonstrated in a western population.

**Methods** A retrospective, single centre cohort analysis in patients diagnosed with HCC from 2008–2013. From 2008 an automated recall system for 6-monthly ultrasound was instigated by the radiology department, in preference ad hoc ultrasound requests. Patients with abnormal lesions proceeded to CT, MRI or liver biopsy according to defined international criteria. The primary end-points evaluated were stage of cancer detection (early i.e. BCLC 0 or A), versus late presentation (BCLC B-D) and patient survival from time of diagnosis to 12 months and 60 months.

**Results** 160 patients were identified. Surveillance status was known in 132 patients. Median patient age was 68 years (57–75), median number of lesions was one, diameter of largest lesion 30 mm (19–50), and AFP 19.5 (5–250). Patients under surveillance were more likely to have disease at a curative stage 67 vs. 39% (p = 0.006, OR 0.39 (0.41–0.84), and had better survival at 1 year 80 vs. 62% (p = 0.04, OR 0.77 (0.62–0.97), and at 5 years 60 vs. 41% (p = 0.046, OR 0.69 (0.48–0.98). On univariate analysis the following variables on survival were evaluated: Age (p = 0.11), Number HCC nodules (p = 0.31), Total diameter of lesions (p = 0.001), Diameter of largest lesion (p < 0.001), AFP (p < 0.001). The presence on imaging of extra-hepatic metastases (p = 0.006), lymph nodes (p = 0.004), and portal vein thrombosis (p < 0.001), were associated with poorer survival.

**Conclusion** Surveillance for hepatocellular carcinoma leads to earlier diagnosis and improved survival.

**Disclosure of Interest** None Declared.

**OC-033** THE TWEAK AND FN14 PATHWAY AS POTENTIAL MEDIATOR OF LIVER FIBROSIS

1A Wilhelm*, 1M Munir, 1E Humphreys, 1D Adams, 1J Burkly, 1S Aford, 1C Weston.
1Centre for Liver Research, University of Birmingham, Birmingham, UK; 2Biogen Idec, Cambridge, USA

10.1136/gutjnl-2014-307263.33

**Introduction** The TNF superfamily ligand TWEAK and its cognate receptor Fn14 have been implicated in the pathogenesis of liver disease and have been predominantly associated with liver progenitor cell proliferation and ductular reaction. We hypothesised that TWEAK and Fn14 may also be involved in the establishment and progression of fibrosis via a direct effect on hepatic stellate cell (HSC) function.

**Methods** TWEAK and Fn14 expression was studied by qPCR, western blot and immunostaining in tissue and stromal cells from explanted human liver specimens and normal donor livers surplus to surgical requirements, or as a byproduct of surgical resection. The responses of HSCs to TWEAK were investigated by western blot, live cell imaging and proliferation assays. TWEAK was measured in HSC supernatant by ELISA.

**Results** Confocal microscopy revealed localisation of Fn14 to cells expressing stromal markers in normal human livers, with significant upregulation in diseased livers. Fn14 expression was confirmed in both primary human HSCs and myofibroblasts in vitro. Stimulation with recombinant TWEAK led to an upregulation of NF-kB signalling and induced proliferation in cultured HSCs. TWEAK immunostaining localised the protein to the fibrotic areas of ALD and NASH liver sections suggestive of an autocrine regulation of Fn14 signalling. We confirmed that HSCs express TWEAK and release it into their environment by qPCR and ELISA, and demonstrated that function-blocking TWEAK antibodies reduced the proliferative capacity of HSC.

**Conclusion** Our study suggests that TWEAK/Fn14 promotes liver fibrosis via enhanced proliferation of HSC, possibly through an autocrine mechanism driven by HSC production of TWEAK.

**Disclosure of Interest** None Declared.

**OC-034** OUTCOME OF PATIENTS CONSIDERED UNSUITABLE FOR LIVER TRANSPLANTATION – A MISSED OPPORTUNITY FOR PALLIATIVE CARE

A Phoolchund*, S Murray, B Hogan, J O’Beirne. Royal Free Hospital and UCL Institute of Liver and Digestive Health, Royal Free Hospital, London, UK

10.1136/gutjnl-2014-307263.34