Vaccination. One infant (who received 3 vaccinations) is HBsAg negative with a viral load of 1.7 × 10^3 IU/ml. 2 of 15 (13.3%) infants had an adequate response to HBIG at birth. 7 mothers had a HBV DNA level > 10^7 IU/ml. 2 were HBeAg positive mothers. All 15 infants born to HBeAg positive mothers were given HBIG at birth.

**Introduction**

Antenatal screening for Hepatitis B (HBV) has been offered to all pregnant women in the UK since 2000. Immunoprophylaxis of infants is essential to reduce the risk of vertical transmission. It is recommended that HBV vaccination (4 serial doses) be given to all infants born to HBV positive mothers. In addition, Hepatitis B immunoglobulin (HBIG) is given to all infants of Hepatitis B e antigen (HBeAg) positive mothers. Infants should have post-vaccination testing between 9–18 months. Hepatitis B surface antigen (HBsAg) negative infants with anti-HBs levels > 10 IU/ml need no further management. If anti-HBs is < 10 IU/ml infants should receive a second vaccination series. Our aim was to evaluate the management of infants born to HBV positive mothers.

**Methods**

All HBV positive pregnant women seen in our hospital between January 2008 and November 2011 were identified from an obstetric database. We examined if the infants received the recommended vaccinations, HBIG and post-vaccination testing.

**Results**

From a total of 99 pregnancies data was available for 76 infants. All 15 infants born to HBeAg positive mothers were given HBIG at birth. 7 mothers had a HBV DNA > 10^7 IU/ml. 2 were treated with antiviral therapy during pregnancy. 58 (76.3%) infants received a full vaccination course. Only 35 (54.7%) of the 64 infants who should have had their post-vaccination status checked to date have had this completed. 34 (97.1%) had an adequate response to vaccination. One infant (who received 3 vaccinations) is HBsAg positive with a viral load of 5.4 × 10^6 IU/ml. The mother was HBeAg negative with a viral load of 1.7 × 10^5 IU/ml.

**Conclusion**

HBIG was administered appropriately to the infants at highest risk of vertical transmission of HBV. However, completion of the 4 dose HBV vaccination in infants was suboptimal postpartum and post- vaccination testing was inadequate. Efforts to improve this are now in place and include: prospective data collection to improve quality of data; development of a central reminder system to advise family doctors 2 weeks before each vaccination dose is due; and introduction of dry blood spot testing of the infants to improve the acceptability of testing.

**Disclosure of Interest**

None Declared

**PTU-116**

**TRANSPANTATION FOR ALD: LESSONS FROM THE EXPLANT**

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**Introduction**

Alcoholic liver disease (ALD) remains one of the commonest indications for liver transplantation in Europe. The histological features of ALD vary, depending on extent and stage of injury. No features are reliably pathognomonic of ALD. We describe the histological spectrum of explants of well characterised cohort of patients undergoing transplantation for ALD.

**Methods**

Consecutive explants (n = 84) of patients transplanted for ALD in our institution between 2002 and 2011 were selected for retrospective histological assessment. Explants were scored blinded by two pathologists using a predetermined pro-forma. Histological assessment including the presence and degree of cirrhosis, steatosis, inflammation, inclusions, siderosis and neoplastic changes were scored semi-quantitatively.

**Results**

Median age was 54 and the majority (70%) were male. All patients had a long history of alcohol excess but reported abstinance for at least 6 months by transplantation. The aetiology was ALD (n = 80) and mixed ALD/HCV (n = 4). The majority (n = 83) had a mixed or macronodular cirrhosis with evidence of re-modelling in a significant number; one had pre-septal cirrhosis. Alpha-1 antitrypsin inclusion bodies were seen in 9 (10.7%); only 4 of these had serum α1AT levels below normal. Parenchymal siderosis was present in 39 (46.4%); in 19 (22.6%) this was grade 3–4. Amongst these, only single mutations of the HFE gene were identified. Induced cell change was seen in 67 (79.8%) and 47 (56%) had the “abstinent cell” phenotype. While 46 (54.8%) had Mallory-denek bodies (MDB), 22 (26.1% of total) patients had both “abstinent cells” and MDB. Ballooning (n = 45, 53.6%) and steatosis (n = 31, 36.9%) were also seen. HCC was present in 14 (16.7%), with dysplastic nodules in 15 (17.9%), small-cell change in 20 (23.8%) and large-cell change 50 (59.5%). Phlebosclerosis and parenchymal extension were universal findings.

**Conclusion**

We describe a wide spectrum of histological features in a large cohort transplanted for end-stage ALD. We demonstrate that despite abstinence, over half have residual MDB and ballooning. Conversely, over half had the recently described “abstinent cell” phenotype. Therefore, the presence of MDB should not be used as evidence of continued alcohol consumption; the presence of induced or abstinent cells correlates more strongly with reported abstinence.

**Disclosure of Interest**

None Declared

**PTU-117**

**IMAGING HEPATIC NEUTROPHIL MIGRATION WITH INDIUM-111-(111In)-RADIOLABELLED LEUCOCYTES: A NOVEL NON-INVASIVE DIAGNOSTIC TEST FOR SEVERE ALCOHOLIC HEPATITIS**

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**Introduction**

Clinical diagnosis of severe alcoholic hepatitis (SAH) can be unreliable and liver biopsy often problematic. Neutrophil infiltration is a key histological feature of SAH and predicts corticosteroid (CS) response (1). We assessed the use of radiolabelled leukocyte scintigraphy to diagnose SAH through in vivo imaging of hepatic neutrophil migration. Liver signals at 111In-leucocyte scintigraphy portray margination at 30min post injection (PI) and destruction at 24h, and are normally matched. We hypothesised that both these functions would be impaired in SAH and liver activity would increase between 30min and 24h due to neutrophil migration.

**Methods**

16 patients with SAH (Discriminant Function 54.8 ± 16.3, 14 biopsied), 14 with inactive alcoholic cirrhosis (Child-Pugh score 5.4 ± 0.9) and 11 controls were recruited. Abdominal gamma camera images were obtained 50min and 24h after IV injection of autologous 111In-labelled leucocytes and change in liver activity expressed as a 24h:30min liver ratio (PI) and destruction at 24h, and are normally matched. We hypothesised that both these functions would be impaired in SAH and liver activity would increase between 30min and 24h due to neutrophil migration.

**Results**

Liver activity significantly increased in SAH but was static or fell in cirrhotic and normal control groups (24h:30min liver ratio 2.18 (IQR 1.62) versus 0.97 (0.28) and 0.78 (0.15) respectively, p < 0.001). Figure 1 shows example gamma camera images. Liver activity ratios in SAH correlated with histological neutrophil infiltration (p = 0.571, p = 0.041) and microautoradiography demonstrated intact intrahepatic radiolabelled leucocytes as the likely source of 24h liver activity. Prominent 30min liver activity was a consistent finding in SAH, providing in vivo evidence of neutrophil
priming. There was no correlation between imaging/histology and CS response, likely due to the small heterogeneous study group.

Abstract PTU-117 Figure 1

Conclusion In-labelled leucocyte scintigraphy is a novel technique for assessment of hepatic neutrophil migration in SAH. It has potential to be a non-invasive diagnostic tool and may help to prospectively identify those likely to respond to CS.

Disclosure of Interest None Declared

REFERENCE


PTU-118 LONG-TERM OUTCOME IN SEVERE ALCOHOLIC HEPATITIS

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Introduction Although short-term outcome in severe alcoholic hepatitis (SAH) is well described, its long-term course is largely unknown. Our aim was to assess long-term outcome in SAH.

Methods Cohort study of patients with SAH (Discriminant Function ≥32) admitted to our institute, identified retrospectively 2007–2009 then prospectively until August 2011. Clinical and laboratory parameters were recorded at accession and subsequent follow-up. Kaplan-Meier (KM) and Cox proportional hazards analyses were performed. Data are presented as mean ±SD or median (IQR).

Results 109 consecutive patients with SAH were included (63.3% men, age 49.6 ± 9.4 yrs) with median follow-up of 40.7 mths (95% CI 37.2–44.3). At accession median DF was 58 (54), MELD 23 (6) and Glasgow Alcoholic Hepatitis Score 8 (2). 55.0% drank spirits, 86.2% had established cirrhosis and 65.1% received corticosteroids to improve abstinence following SAH are urgently needed, especially in view of the increasing numbers of deaths due to alcohol-related liver disease in the UK.

Disclosure of Interest None Declared

PTU-119 HISTOPATHOLOGY AND LONG TERM CLINICAL PROGNOSIS IN HCV. DOES THE BIOPSY STILL HAVE ANYTHING TO ADD?

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Introduction Hepatitis C (HCV) was first characterised in 1989 and most UK sufferers are infected in early adulthood. While a cirrhosis rate of 20% over 20 years is often quoted, life-long outcomes and factors predicting these are not fully defined. With increasing availability of noninvasive markers of fibrosis, liver biopsy is now rarely performed at diagnosis.

This study aims to explore the value of baseline liver biopsy in determining long-term clinical prognosis.

Methods Patients were identified from a historical HCV study cohort (n = 202) at one centre who had a diagnostic liver biopsy at baseline (between 1984 and 2004) and available follow-up data. Clinical and histologic data were recorded. Kaplan-Meier plots of time to cirrhosis and multivariate Cox regression were performed.

Results 146 patients had adequate follow-up data. The mean duration of follow-up from presumed date of infection was 22 (SD 8.51) years with a mean follow-up from biopsy of 10 years (SD 5.3). The majority of patients were male 90 (62%) and had been infected through IV drug use 81 (55%). 56 patients (40%) had Genotype 1 infection. From baseline biopsies, 44 (30%) had moderate and 12 (8%) severe steatosis. 93 (64%) had Ishak fibrosis scores of 0 or 1 at baseline, 54 (35%) 2–3 and 10 (7%) 4–5. 9 patients (6%) had cirrhosis at base.

A total of 31 (21%) developed clinical cirrhosis during follow up. From these, 11 compensated and 5 developed hepatocellular carcinoma. Factors associated with shorter time to cirrhosis were later age at diagnosis (HR: 1.08, 95% CI 1.03–1.13) and increasing Ishak fibrosis score (HR: 2.15, 95% CI 1.59–2.85). The 10 year cirrhosis free survival from baseline biopsy was 93% for Ishak fibrosis scores of 0 or 1, compared with 78% and 27% for scores 263 and 485 respectively (see Fig 1.). 95 (65%) patients were treated, of whom 64 (67%) achieved a sustained virological response (SVR) with standard treatment. History of alcohol excess, genotype 1 and severe steatosis were significant negative predictors of SVR.