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# IN BRIEF

#### **HEPATITIS**

HCV virions incorporate host regulators to avoid lysis Patients with chronic HCV often have cross-reactive neutralizing agents but are unable to clear the virus, possibly because the HCV virions are able to avoid lysis. A study has shown that cultured HCV particles incorporate host CD59, a regulator of lysis by complement activation, into their cell membrane. These virions were able to resist antibody-dependent complement-mediated lysis. Addition of CD59 blockers to

**Original article** Amet, T. *et al.* CD59 incorporation protects hepatitis C virus against complement-mediated destruction. *Hepatology* doi:10.1002/hep.24686

#### HEPATOCELLULAR CARCINOMA

Lymphocyte infiltration predicts survival

the culture reversed this phenomenon.

Few methods can accurately predict survival of patients with hepatocellular carcinoma. Using quantitative PCR and samples from 57 patients from Singapore and a validation cohort of 98 patients, a signature of 14 immune genes has been identified that can predict survival of these patients, particularly during the early stages of the disease. The signature includes the chemokine genes *CXCL10*, *CCL5* and *CCL2*. These chemokines drive lymphocyte infiltration of the tumors, which results in increased death of cancer cells.

**Original article** Chew, V. et al. Chemokine-driven lymphocyte infiltration: an early intratumoural event determining long-term survival in resectable hepatocelluar carcinoma. *Gut* doi:10.1136/gutjnl-2011-300509

#### **GUT MICROBIOTA**

## Tolerance of commensal microbiota

The thymus prevents autoimmunity by eliminating or differentiating self-reactive T cells; however, it is unknown whether T cells can be altered to recognize antigens from commensal bacteria, which may prevent diseases such as IBD. Findings from a mouse study indicate that antigen-specific T<sub>REG</sub> cells are generated in response to an individual's microbiota. This occurs after T cells have been altered in the thymus and could be a mechanism by which a host can tolerate commensal bacteria.

**Original article** Lathrop, S. K. *et al.* Peripheral education of the immune system by colonic commensal microbiota. *Nature* doi:10.1038/nature10434

#### **HEPATITIS**

### New therapy for hepatitis C

A combination of the protease inhibitor BI 201335 (120 mg per day), the polymerase inhibitor BI 207127 (400 mg or 600 mg three times a day) and ribavirin (1,000–1,200 mg per day) for 4 weeks was safe and efficacious in 32 treatment-naive patients chronically infected with HCV genotype 1. In the group given 600 mg of BI 207127, the virological response was 100% by day 29, compared with 73% in those given 400 mg. The response to this combination was rapid and strong, and none of the patients experienced a severe adverse event.

 $\label{eq:continuous} \begin{tabular}{ll} \textbf{Original article} Zeuzem, S.\ et\ al.\ Efficacy\ of\ the\ protease\ inhibitor\ BI\ 201335, polymerase\ inhibitor\ BI\ 207127,\ and\ ribavirin\ in\ patients\ with\ chronic\ HCV\ infection.\ Gastroenterology\ doi:10.1053/j.gastro.2011.08.051 \end{tabular}$