Toll-like receptors 2 and 4 are differentially involved in Fas dependent apoptosis in Peyer’s patch and the liver at an early stage after bile duct ligation in mice

A Ogawa, T Tagawa, H Nishimura, T Yajima, T Abe, T Arai, M Taniguchi, K Takeda, S Akira, Y Nimura, Y Yoshikai

Background and aims: Surgical management of extraphelial cholestasis is frequently complicated by bacterial translocation and severe liver injury. The aim of this study was to clarify the involvement of Toll-like receptors (TLR) in the pathogenesis of bacterial translocation and liver injury in obstructive cholestasis.

Methods: TLR2 deficient (TLR2−/−), MyD88−/−, Jα281−/−, gld/gld, and lpr/lpr mice, all of which have a C57BL/6 background, and C3H/HeN and TLR4 mutated C3H/HeJ mice were subjected to bile duct ligation (BDL). Faecal IgA and serum alanine aminotransferase levels were determined after BDL. Apoptosis was examined by histological and flow cytometric analyses of cells from Peyer’s patches and the liver.

Results: The size and number of B cells in Peyer’s patch markedly decreased on day 3 after BDL. Increased apoptosis in Peyer’s patch B cells was evident on day 1 after BDL in control mice but not in lpr/lpr, MyD88−/−, or C3H/HeJ mice. On the other hand, TLR2 and Fas ligand expression on intrahepatic NK1.1+ T cells increased on day 1 after BDL in C57BL/6 mice. Liver injury and apoptosis were evident on day 1 after BDL in control and C3H/HeJ mice but were significantly reduced in TLR2−/−, Jα281−/−, gld/gld, and lpr/lpr mice.

Conclusions: TLR4 and TLR2 may play important roles in Fas dependent apoptosis in Peyer’s patch B cells and hepatocytes, respectively, at an early stage after BDL in mice.

MATERIALS AND METHODS

Mice

Seven to eight week old female C57BL/6, gld/gld, and lpr/lpr mice with a B6 background, C3H/HeN, and C3H/HeJ mice were purchased from Japan SLC, Inc. (Hamamatsu, Japan). Mutant mice deficient in TLR2, MyD88, or Jα281 with a C57BL/6 background were generated by gene targeting, as described previously.5 6 7 8 Age and sex matched groups of homozygous mice and their littermates (TLR2−/−) mice were used for the experiments. In some experiments, siblings from the same mother were used. All of the mice used for the experiments. In some experiments, siblings from the same mother were used.
experiments were maintained under specific pathogen free conditions in our animal facility.

**Surgical procedure**

After seven days of acclimatisation, surgery was performed under sterile conditions. Mice were anaesthetised by intraperitoneal pentobarbital injection (50 mg/kg). An abdominal midline incision was made, and the common bile duct was identified, ligated, and divided as previously described. Control animals underwent a sham procedure in which the common bile duct was identified and exposed but not ligated.

**Preparation of cells**

Fresh liver was immediately perfused with sterile Hank’s balanced salt solution (HBSS) through the portal vein to wash out all remaining peripheral blood and then meshed with stainless steel mesh. After coarse pieces had been removed by centrifugation at 50 g for one minute, cell suspensions were again centrifuged, resuspended in 8 ml of 45% Percoll (Amersham Biosciences, Uppsala, Sweden), and layered on 5 ml of 66.6% Percoll. The gradients were centrifuged at 600 g for 20 minutes at 20°C. Lymphocytes at the interface were harvested and washed twice with HBSS. Peyer’s patches (4–6 patches per mouse) were excised aseptically from the exposed small intestine.

**Bacterial growth in organs**

Peritoneal exudates were obtained from the peritoneal cavity by lavage with 3 ml of HBSS. For enumeration of viable bacteria in the liver, the liver was perfused with 8 ml of sterile HBSS to wash out bacteria in the blood vessels immediately after mice had been bled. The liver and spleen were removed and separated into sterile Teflon coated homogenisers (Asahi Techno Glass Co., Tokyo, Japan), each containing 2 ml of cold phosphate buffered saline (PBS). After each organ had been homogenised thoroughly, bacterial counts in homogenates were established by plating serial 10-fold dilutions in sterile distilled water on blood agar and MacConkey agar (Nissui, Tokyo, Japan). Colonies were counted 24 hours later, after incubation at 37°C.

**Flow cytometric analysis**

Cells were preincubated with a culture supernatant from rat-IgG1, producing hybridoma to prevent non-specific binding. Cells were preincubated with a culture supernatant from a hybridoma. Flow cytometric analysis was performed using an in situ apoptosis detection kit (Apoptag, Intergen, Purchase, New York, USA). All steps were performed using an in situ apoptosis detection kit (Apoptag, Intergen, Purchase, New York, USA). All steps were performed according to the manufacturer’s instructions. Briefly, paraffin embedded sections were deparaffinised and rehydrated and then permeabilised by incubation with 0.55 unit/ml proteinase K (Wako, Tokyo, Japan) for 15 minutes at 37°C. Tissue sections were then washed in distilled water. Inactivation of endogenous peroxidases was accomplished by immersing the tissue sections in 3% hydrogen peroxide diluted in methanol for 10 minutes at room temperature in the dark, after which 400 ml of 1× binding buffer were added, as recommended by the manufacturer. For propidium iodide (PI) staining, cells were washed twice with PBS and fixed with ice cold 70% ethanol/PBS. Cells were then kept on ice for at least one hour. Subsequently, the medium was removed by centrifugation, and pellets were resuspended in 100 μl PBS. Cells were then incubated in the dark for 30 minutes at 4°C in the presence of PI (50 μg/ml; Sigma, St Louis, Missouri, USA) and DNase-free RNase A (250 μg/ml; Roche, Indianapolis, Indiana, USA). Thereafter, cell cycle status and apoptosis were determined using a FACSCalibur flow cytometer (Becton Dickinson & Co., San Jose, California, USA).

**IgA in faecal samples**

Faecal samples (0.02 g) was each incubated with 1 ml of PBS at room temperature for 60 minutes, vortexed, left for 15 minutes, revortexed until all materials were suspended, and centrifuged at 2000 g for 10 minutes. The supernatants were removed and tested for total IgA by enzyme linked immunosorbent assay.

**Histopathological examination**

The liver and Peyer’s patches were removed and fixed in 10% buffered formalin. After paraffin embedding, tissue sections were cut (3 μm) and stained with haematoxylin-eosin for morphological evaluation. An experienced hepatopathologist evaluated the sections and graded them as follows: grade 0, normal histomorphology; grade 1, minor inflammatory infiltrates with occasional liver cell necrosis; grade 2, moderate liver damage with inflammatory infiltrates and focal necroses; and grade 3, extensive liver damage accompanied by diffusely distributed cell necroses. At least three separate sections were assessed per liver in each group of five mice in a blinded fashion. In situ terminal deoxynucleotidyl transferase dUTP nick end labelling (TUNEL) assay was performed using an in situ apoptosis detection kit (ApopTag, Intergen, Purchase, New York, USA). All steps were performed according to the manufacturer’s instructions. Briefly, paraffin embedded sections were deparaffinised and rehydrated and then permeabilised by incubation with 0.55 unit/ml proteinase K (Wako, Tokyo, Japan) for 15 minutes at 37°C. Tissue sections were then washed in distilled water. Inactivation of endogenous peroxidases was accomplished by immersing the tissue sections in 3% hydrogen peroxide diluted in methanol for 10 minutes at room temperature in the dark, after which 400 ml of 1× binding buffer were added, as recommended by the manufacturer. For propidium iodide (PI) staining, cells were washed twice with PBS and fixed with ice cold 70% ethanol/PBS. Cells were then kept on ice for at least one hour. Subsequently, the medium was removed by centrifugation, and pellets were resuspended in 100 μl PBS. Cells were then incubated in the dark for 30 minutes at 4°C in the presence of PI (50 μg/ml; Sigma, St Louis, Missouri, USA) and DNase-free RNase A (250 μg/ml; Roche, Indianapolis, Indiana, USA). Thereafter, cell cycle status and apoptosis were determined using a FACSCalibur flow cytometer (Becton Dickinson & Co., San Jose, California, USA).

<p>| Table 1 | Incidence of bacterial translocation in bile duct ligated (BDL) and sham operated mice |</p>
<table>
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<th>Days after operation</th>
<th>Liver</th>
<th>Mesenteric lymph nodes</th>
<th>Spleen</th>
<th>Peritoneal exudate</th>
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</table>

Values represent the incidence of bacterial translocation in the liver, mesenteric lymph nodes, spleen, peritoneal exudates, and bile of BDL and sham operated mice (n = 6) on the indicated days after operation.

*p<0.02
temperature. After the slides had been washed with distilled water, sections were incubated in equilibration buffer for five minutes. Sections were then incubated with a labelling solution containing terminal deoxynucleotidyl transferase in a humidified chamber for one hour at 37°C. Reactions were terminated by rinsing the sections in a stop/wash buffer. The incorporated digoxigen-dUTP was detected by incubation with antidigoxigenin peroxidase at room temperature for 30 minutes, and positive reactions were revealed using 3,3-diaminobenzidine. Haematoxylin was used for counterstaining nuclei, and slides were coverslipped.

Real time polymerase chain reaction (PCR)
mRNA was obtained from EPICS sorted NK, NKT, or T cells in the liver using the Quick Prep Micro mRNA Purification kit (Amersham Biosciences, Buckinghamshire, UK). Each RNA sample was reverse transcribed into cDNA using random primers and Moloney murine leukaemia virus transcriptase. Real time PCR was performed using a Taq Man PCR kit (Applied Biosystems, Foster City, California, USA) and primers for Fas ligand (FasL). Glyceraldehyde-3-phosphate dehydrogenase (GAPDH) primers were used as a control for RNA isolation and integrity. Real time PCR was performed using the ABI PRISM 7000 sequence detector thermal cycler (Applied Biosystem). Results were expressed as a ratio of product copies/ml to copies/ml of the housekeeping gene GAPDH from the same RNA (respective cDNA) sample and PCR run.

Assay for serum alanine aminotransaminase (ALT) and caspase 3 activity in liver tissues
Serum ALT activity was determined using a transaminase test kit (Wako, Osaka, Japan). Serum total bilirubin levels were measured using a commercially available kit following the manufacturer’s instructions (Sigma Diagnostics Kit No550 for bilirubin; Sigma Chemical Co). The activity of caspase 3 was determined by the Fluorometric CaspACE Assay System (Promega, Madison, Wisconsin, USA) according to the manufacturer’s instructions. Briefly, fresh isolated liver was meshed with a stainless steel mesh and resuspended in hypotonic lysis buffer. Cells were lysed by subjecting them to four cycles of freezing and thawing. After centrifuging, the activity of caspase 3 in the supernatant was measured with fluorogenic caspase 3 substrate (Ac-DEVD-AMC) at an excitation wavelength of 360 nm and an emission wavelength of 460 nm.

Statistical analysis
Data were analysed by the Student’s t test and Fisher’s exact test, and a Bonferroni correction was applied for multiple comparison. The value of p<0.05 was considered statistically significant.

RESULTS
Bacterial translocation after BDL
It has been reported that cholestasis promoted bacterial translocation from the gut. To determine whether bacterial translocation occurs in choledochal mice, we examined the translocation of Enterobacteriaceae in the liver,
mesenteric lymph nodes, spleen, peritoneal exudate, and bile in mice that had undergone BDL. Bacterial translocation was found three days after BDL in the liver and mesenteric lymph nodes whereas no bacteria were detected in sham operated mice (p<0.05) (table 1). Most of the bacteria were Escherichia coli and Enterococcus faecalis and a few were Klebsiella pneumonia and Staphylococcus aureus, all of which are present in bacterial flora. These results suggest that bacterial translocation from the intestine often occurs in BDL mice.

Increased faecal IgA and apoptosis in Peyer’s patch B lymphocytes after BDL

IgA production is associated with bacterial translocation in BDL mice. We first examined IgA levels in the intestine after BDL. As shown in fig 1, levels of IgA in faeces transiently increased on day 1 after BDL (p<0.01) and then decreased to control levels.

We next examined the size and cellularity of Peyer’s patches after BDL. The sizes of Peyer’s patches on day 1 after BDL were compared with those of sham operated groups. The size of Peyer’s patches in BDL mice was significantly smaller than that in sham operated groups (p<0.01) (table 2).

Figure 3  Apoptosis in Peyer’s patch lymphocytes after bile duct ligation (BDL). (A) Apoptotic cells in situ in Peyer’s patches after sham operation (a, b) or BDL (c, d). Peyer’s patches in C57BL/6 mice were obtained on day 1 after BDL. Detection of DNA fragmentation was performed using the in situ terminal deoxynucleotidyl transferase dUTP nick end labelling (TUNEL) method, as described in the materials and methods. TUNEL positive cells are dark brown. Data shown are representative of three independent experiments. (Original magnification (left panel) ×100; (right panel) ×400.) (B) Flow cytometric analysis of Peyer’s patch cells stained with annexin V. Peyer’s patch cells obtained from BDL mice on day 1 after operation were stained with anti-B220 monoclonal antibody (mAb) and anti-annexin V mAb. Analysis gates were set on B220+ corresponding to B cells. The percentage of annexin V positive cells are indicated. Data shown are representative of three independent experiments.

Figure 4  Flow cytometric analysis of Peyer’s patch cells in lpr/lpr, TLR2−/−, and C3H/HeJ mice after bile duct ligation (BDL). (A) Expression of Fas, CD40, and Fas ligand (FasL) on Peyer’s patch cells after BDL. Peyer’s patches in C57BL/6 mice were obtained on day 1 after BDL and stained with anti-B220 and anti-CD3 monoclonal antibody (mAb) and anti-Fas, FasL, or CD40 mAb. The shaded area shows the BDL groups and unshaded area the sham operated groups. Data shown are representative of three independent experiments. (B) Flow cytometric analysis of lymphocytes of Peyer’s patches in lpr/lpr, C3H/HeN, C3H/HeJ, and TLR2−/− mice after BDL. Peyer’s patch cells were obtained on day 3 after operation and stained with anti-B220 and anti-CD3 mAb. Analysis gate was set on lymphocytes using forward and side scatter. The percentage of each fraction in total lymphocytes is indicated.
BDL were similar to those in sham operated mice (data not shown) but were markedly reduced on day 3 after BDL (fig 2A). Total cell numbers of Peyer’s patches were significantly decreased by day 3 after BDL and thereafter (p<0.05) (fig 2B). Flow cytometric analysis revealed that the number of B220<sup>+</sup> B cells decreased on day 3 after BDL (fig 2C). We next attempted to assess the extent of apoptosis in Peyer’s patches by the TUNEL technique. As shown in fig 3A, a large number of TUNEL positive cells were observed on day 1 after BDL. Cells showed characteristics of apoptosis, such as shrinking, chromatin clumping, and nuclear fragmentation, and they were mostly located in the germinal centres but not in the marginal zone, which is abundant in T cells. There were a few apoptotic cells in other organs such as the spleen on day 1 after BDL (data not shown). To determine the number of Peyer’s patch cells undergoing apoptosis, we stained cells with anti-B220, anti-CD3<sub>e</sub>, and anti-annexin V mAbs. Analysis gates were set on B220<sup>+</sup> cells corresponding to B cells. A major increase in annexin V positive cells was observed on day 1 after BDL in Peyer’s patch B cells (fig 3B) but not in T cells (data not shown). Thus increased apoptosis in Peyer’s patches of BDL mice is restricted to cells of the B lineage.

Involvement of Fas and TLR4 in apoptosis of Peyer’s patch B cells after BDL

The Fas/FasL system is known to be involved in B cell apoptosis. We therefore examined expression levels of Fas on Peyer’s patch B cells after BDL. As shown in fig 4A, expression levels of Fas were significantly higher in B cells of Peyer’s patches on day 1 after BDL than those of sham operated control mice. We found no difference in expression levels of CD40 on Peyer’s patch B cells and FasL on Peyer’s patch T cells on day 1 after BDL compared with those of sham operated mice.

To determine whether Fas signalling is involved in Peyer’s patch B cell apoptosis in BDL mice, we examined apoptosis of B cells in Peyer’s patches from lpr/lpr mice after BDL. The number of Peyer’s patch B cells in lpr/lpr mice was not reduced after BDL, and apoptosis was not evident in lpr/lpr mice after BDL (fig 4B, fig 5). Thus Fas signalling is important for apoptosis of B cells in Peyer’s patches after BDL. We next examined whether TLR signalling is involved in B cell apoptosis after BDL using TLR4 mutated C3H/HeJ mice or mice deficient in TLR2 or MyD88, a key adaptor protein for most of the TLRs. As shown in fig 4B, the number of Peyer’s patch B cells was reduced in C3H/HeJ and TLR2<sup>−/−</sup> mice but not in C3H/HeJ mice. Apoptosis in Peyer’s patch was not evident in C3H/HeJ or MyD88<sup>−/−</sup> mice after BDL (fig 5). These results suggest that MyD88 dependent TLR4 signalling plays an important role in Peyer’s patch B cell apoptosis after BDL.

Increased expression of FasL and TLR2 on intrahepatic NKT cells after BDL

We previously reported that FasL expression was selectively induced on NK T cells via TLR2 signalling and it was partly responsible for liver injury caused by Gram negative bacterial infection. To determine whether expression of FasL and TLR2 was induced on NK T cells in vivo by BDL, we analysed expression of FasL and TLR2 on intrahepatic lymphocytes of C57BL/6 mice on day 1 after BDL by flow cytometry. No change in the number of NK and NK T cells was observed between before and after BDL in C57BL/6 mice (data not shown). As shown in fig 6A, expression levels of FasL and TLR2 on NK T cells were apparently increased after BDL. Although NK cells expressed slightly increased levels of FasL and TLR2 after BDL, T cells did not express these even after BDL. We further examined FasL mRNA expression by real time PCR and found that FasL mRNA was increased in NK T cells of BDL mice (fig 6B).

Involvement of FasL, NK T cells, and TLR2 in liver injury after BDL

BDL causes early hepatocellular injury. Peak ALT levels were observed on day 1 after BDL in C57BL/6 mice (fig 7A). To investigate the involvement of Fas/FasL and TLR2 on liver NKT cells in liver injury following BDL, we compared serum...
ALT levels in Vx14 NKT cell deficient (Jx281−/−), lpr/lpr, gld/gld, TLR2−/−, MyD88−/−, C57BL/6, C3H/HeJ, and C3H/HeN mice after BDL. Serum ALT levels in Jx281−/−, gld/gld, and TLR2−/− mice were significantly lower than those in control C57BL/6 mice on day 1 after BDL (fig 7B). On the other hand, there was no difference in ALT levels between TLR4 mutated C3H/HeJ and control C3H/HeN mice after BDL (fig 7B). Differences in serum ALT serum among Jx281−/−, gld/gld, and TLR2−/− mice and control C57BL/6 mice could not be attributed to differences in the magnitude of cholestasis because bilirubin levels were almost the same in these groups (13.5 (5.6) mg/dl for Jx281−/− mice, 12.5 (3.6) mg/dl for gld/gld mice, 14.5 (3.9) mg/dl for TLR2−/− mice, and 13.2 (4.6) mg/dl for C57BL/6 mice).

Histopathological examination with haematoxylin-eosin staining of liver specimens on day 1 after BDL demonstrated bile duct proliferation, portal oedema, and mild portal infiltrates, indicating similar ductular responses of the liver to large bile duct obstruction in all groups. However, necroinflammatory foci, cell death of hepatocytes characterised by cell shrinkage and chromat in condensation, and lymphocyte infiltration were obviously less in Jx281−/−, lpr, gld, TLR2−/−, and C3H/HeJ mice than in control wild-type mice after BDL (fig 8A, B). Confluent foci of hepatocyte feathery degeneration due to bile acid cytotoxicity ( bile infarcts) were significantly reduced in Jx281−/− (6.7 (5.7) foci per low power field), lpr (9.3 (3.1) foci), gld/gld (12.5 (3.5) foci), and TLR2−/− mice (8.3 (4.2) foci) compared with control mice (19.3 (4.2) foci) 24 hours after BDL (p<0.05). To confirm the difference in frequency of apoptotic cells among these mutant mice, we identified apoptotic cells using the TUNEL technique and the numbers of TUNEL positive cells per field were quantified (fig 8C). Greater numbers of cells undergoing apoptosis were observed in the livers of control and C3H/HeJ mice whereas the numbers of apoptotic cells in Jx281−/−, lpr/lpr, gld/gld, TLR2−/−, and
MyD88<sup>−/−</sup> mice remained at smaller levels (p < 0.05) (fig 8C). To further compare the degree of apoptosis in the liver of these mutant mice subjected to BDL, we examined the activities of caspase 3, a downstream caspase in apoptosis, in liver tissues on postoperative day 1. Consistent with serum ALT levels and TUNEL positive cells, caspase 3 activity in Jα281<sup>−/−</sup>, lpr/lpr, gld/gld, MyD88<sup>−/−</sup>, TLR2<sup>−/−</sup>, C57BL/6, C3H/HeN, and C3H/HeJ mice was significantly lower than that in control mice on day 1 after BDL (p < 0.05) (fig 8D). Taken together, these results indicate that FasL, NKT cells, and TLR2 contribute to liver injury in the early stage after BDL.

**DISCUSSION**

We have shown that increased apoptosis in Peyer’s patch B cells is evident on day 1 after BDL in normal mice but not in lpr/lpr mice, MyD88<sup>−/−</sup>, or C3H/HeJ mice. Our findings suggest that signalling from both Fas and TLR4 are involved in Peyer’s patch B cell apoptosis after BDL. Although Fas is expressed only weakly on naive B cells, it is strongly expressed in B cells after antigen stimulation and is involved in negative selection in the germinal centre. Fas expression is also upregulated by various stimuli such as stimulation by CD40. We found that expression levels of Fas on Peyer’s patch B cells were upregulated at the early stage after BDL, and the increase in apoptosis in Peyer’s patch B cells appears to be an example of activation induced apoptosis of lymphocytes. We speculate that upregulation of Fas expression on B cells is in part a response to altered exposure to enteric antigens released due to increased gut permeability and microflora explosion observed during cholestasis. The production of IgA is a result of activation of Peyer’s patch B cells.

We found that faecal IgA levels transiently increased very early after BDL, suggesting early activation of Peyer’s patch B cells after BDL. Ayala et al recently reported that Fas was involved in Peyer’s patch B cell apoptosis during...
polymicrobial sepsis caused by caecal ligation and puncture (CLP) and that the increase in apoptosis of Peyer’s patch cells was coincident with an augmented endogenous IgA secretory response after CLP. Consistent with this finding, increased IgA production at the very early stage after BDL appears to be associated with an increase in activation induced apoptosis of Peyer’s patch B cells via the Fas dependent pathway. On the other hand, they showed that TLR4 mediated process was not involved in increased B cell apoptosis during sepsis caused by CLP, which is inconsistent with our present results in obstructive cholestasis. It remains to be determined how TLR4 signalling is involved in Fas dependent apoptosis in B cells in obstructive cholestasis.

Toxic bile salts such as glycochenodeoxycholate (GCDC) have been reported to increase the sensitivity of Fas mediated apoptosis. Therefore, bile acids in synergy with TLR4 ligands may induce efficient apoptosis in Peyer’s patch B cells after BDL through elevation of Fas expression and/or increase in sensitivity to Fas mediated apoptosis.

Another notable finding in the present study is that TLR2, FasL, and NKT cells are responsible for the pathogenesis of liver injury at the early stage after BDL in mice. We have recently found that freshly isolated NKT T cells from livers of naive mice expressed relatively high levels of TLR2 mRNA and that freshly isolated NKT cells respond to synthetic lipoprotein, a ligand for TLR2, by inducing FasL on their surfaces, suggesting that TLR2 on NK T cells can be expressed on their surfaces and function to induce FasL expression on NK T cells. In the present study, we found that expression levels of TLR2 on NKT cells significantly increased after BDL. We have recently reported that TLR2 expressed by NK T cells in the liver contribute to liver injury induced by Gram negative bacteria such as E coli and Salmonella choleraesuis in a FasL/Fas dependent manner. Increased gut permeability and microflora explosion may be evident prior to bacterial translocation after BDL. It is possible that enteric antigens could be released from the gut to the liver in the early stage after BDL. Enteric bacterial stimulation in BDL mice may stimulate liver NKT cells via TLR2 to express FasL, aggravating liver injury at the early stage after BDL.

Miyoshi et al have recently reported that hepatocyte apoptosis after BDL in the mouse involves Fas. It has also been reported that bile acid induces hepatocyte apoptosis in the short term in vitro by a Fas dependent mechanism. Although T cells are a documented source of Fasl. and indispensable for Fas dependent apoptosis of various types of cells, Faubion et al suggested that apoptosis induced by toxic bile salts such as GCDC involves ligand independent oligomerisation of Fas, recruitment of FADD, activation of caspase 8, and subsequent activation of effector proteases, including downstream caspases by FasL independent mechanisms. However, Jz281 mice deficient in NKT cells and gld/gld mice with mutated FasL showed significantly reduced levels of liver injury after BDL. We speculate a totally different mechanism of killing hepatocytes from their assumption that FasL on NKT cells stimulates Fas on hepatocytes to induce apoptosis after BDL.

In conclusion, TLR4 and TLR2 are important for Peyer’s patch B cell apoptosis and NK T cell mediated liver injury, respectively, in a Fas dependent pathway. Our results should be useful for the development of a prophylactic approach to control pathogenesis associated with obstructive cholestasis.

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EDITOR’S QUIZ: GI SNAPSHOT

Answer

From question on page 104

The endoscopy image (fig 1) demonstrates florid duodenal varices. Figure 2 shows an atrophic calcified pancreas with a history of acute pancreatitis complicated by pseudocyst formation and subsequent chronic pancreatitis. Figure 3 illustrates splenic vein thrombosis at the splenic hilum along with extensive duodenal varices. Surprisingly, gastric varices were not demonstrated. The patient continued to be transfusion dependent and haemodynamically unstable and was transferred to the regional liver unit for a splenectomy. He remains well at follow up.

Splenic vein thrombosis complicates acute pancreatitis in up to 19% of cases. Other causes include pancreatic adenocarcinoma, pancreatic cysts, or pseudocysts. Rarely it may also be caused by hypercoagulable states, including nephrotic syndrome and sickle cell anaemia. Pathophysiologically, splenic blood is shunted into the portal circulation through the short gastric vessels, resulting in the formation of gastric and duodenal varices. Oesophageal varices are uncommon in this situation because the collateral circulation does not involve the oesophagus. Variceal bleeding occurs in only 4% of patients with splenic vein thrombosis and treatment consists of either a splenectomy or splenic artery embolisation.

Patient outcome depends on the aetiology of the splenic vein thrombosis.

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