Reversal of murine alcoholic steatohepatitis by pepducin-based functional blockade of interleukin-8 receptors

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ABSTRACT

Objective Alcohol steatohepatitis is a life-threatening condition with short-term mortality up to 40%. It features hepatic neutrophil infiltration and blood neutrophilia, and may evolve from ethanol-induced breakdown of the enteric barrier and consequent bacteraemia. Signalling through CXCR1/2 G-protein-coupled-receptors (GPCRs), the interleukin (IL)-8 receptors, is critical for the recruitment and activation of neutrophils. We have developed short lipopeptides (pepducins), which inhibit post-ligand GPCR activation precisely targeting individual GPCRs.

Design Experimental alcoholic liver disease was induced by administering alcohol and a Lieber–DeCarli high-fat diet. CXCR1/2 GPCRs were blocked via pepducins either from onset of the experiment or after disease was fully established. Hepatic inflammatory infiltration, hepatocyte lipid accumulation and overall survival were assessed as primary outcome parameters. Neutrophil activation was assessed by myeloperoxidase activity and liver cell damage by aspartate aminotransferase and alanine aminotransferase levels. Chemotaxis assays were performed to identify chemotactic signals derived from alcohol-exposed hepatocytes.

Results Here, we show that experimental alcoholic liver disease is driven by CXCR1/2-dependent activation of neutrophils. CXCR1/2-specific pepducins not only protected mice from liver inflammation, weight loss and mortality associated with experimental alcoholic liver disease, but therapeutic administration cured disease and prevented further mortality in fully established disease. Hepatic neutrophil infiltration and triglyceride accumulation was abrogated by CXCR1/2 blockade. Moreover, CXCL-1 plasma levels were decreased with the pepducin therapy as was the transcription of hepatic IL-1β mRNA.

Conclusions We propose that high circulating IL-8 in human alcoholic hepatitis may cause pathogenic overzealous neutrophil activation, and therapeutic blockade via pepducins merits clinical study.

OBJECTIVE

Morbidity and mortality due to excessive alcohol consumption is a major health problem worldwide affecting millions of people. The risk of alcohol-induced liver disease (ALD) increases proportionately with consumption, causing a spectrum of liver diseases ranging from steatosis to terminal liver disease and cirrhosis. At 6-month mortality rates up to 50%, the acutely deadliness manifestation of ALD is severe alcoholic hepatitis (AH). AH is estimated to affect 10–35% of heavy drinkers at some point in time. Severe AH exhibits a very specific presentation with hepatocyte steatosis, neutrophilic liver inflammation and necrosis, a characteristic peripheral blood neutrophilia, and manifests as liver failure including thrombocytopenia and coagulation disorders. Patients surviving severe AH have a high risk of developing fibrosis and cirrhosis, in turn increasing risk for hepatocellular carcinoma. No therapies exist that improve long-term survival of AH. On this dire background, it has indeed been suggested that patients with severe AH should be routinely randomised into experimental trials.

A common working model suggests that ethanol might induce inflammatory chemokines and cytokines via generation of reactive oxygen species and acetaldehydes, which may impact on endotoxin clearance in the liver. This, along with intestinal barrier dysfunction consequent to chronic alcohol
exposure, leads to systemic endotoxaemia, whose extent indeed correlates with progression of liver disease, establishing a vicious circle of pro-inflammatory signalling.

Expression of chemokines that are involved in neutrophil recruitment and activation is massively increased in human AH. Worse prognosis associates with blood neutrophilia, and elevated mRNA expression of interleukin (IL)-8 (CXCL8), CXCL5, Gro-γ (CXCL3) and CXCL6, all ligands of CXCR1/ CXCR2 chemokine receptors. Multivariate analysis revealed hepatic IL-8 protein levels as an independent predictor of 90-day mortality in AH. Neutrophils release an array of bactericidal molecules that might cause tissue destruction. Despite these associative data, a causal role of neutrophils and/or CXCR1/2 signalling in severe AH has not yet been established.

CXCR1 and CXCR2 are G protein-coupled receptors (GPCRs). GPCRs consist of seven transmembrane domains joined by three intracellular loops and a C-terminal domain (referred to as i1-i4) that are important for interaction with GTP and activation of G proteins involved in signal transduction. ‘Pepducins’ are short peptides coupled to a lipid moiety that can be designed as agonists or antagonists with precise specificity in vitro and in vivo for a given GPCR. Pepducins integrate into the inner plasma membrane via a flip-flop mechanism and affect GPCR-dependent G-protein activation. We have reported pepducin ‘x1/2pal-i1’, specific for CXCR1 and CXCR2, which reverses established experimental systemic inflammatory syndrome by inhibiting neutrophil activation. x1/2pal-i1 also abolishes tumour growth in a model of ovarian cancer and inhibits adenoma formation in Apemini mice.

In an experimental model that phenocopies human AH, here we report that CXCR1/2 blockade cures established steatohepatitis, identifying a critical role of CXCR1/2 signalling and neutrophils in propagating this disease.

**Design**

Pepducins directed against CXCR1/2 (x1/2pal-i1) and a nonsense-scrambled pepducin (scram-i1) were synthesised with carboxy-terminal amides by standard iMOC solid-phase methods at the Tufts University Core Facility (Tufts University, Boston, Massachusetts, USA) or purchased from Peptide 2.0 (Chantilly, Virginia, USA). Lieber–DeCarli diet (LDC) was purchased from Dyets (Bethlehem, Pennsylvania, USA). The IL-8 ELISA was from R&D systems, the SYBR-green PCR mix was purchased from Eurogentec (Southampton, UK). Naphthol AS-D chlorocetate, the myeloperoxidase (MPO) assay kit and all other chemicals used in this study were purchased from Sigma Aldrich (St Louis, Missouri, USA). Oil-red-O was from Amaresco (Solon, Ohio, USA); Hep3B and HepG2 cells were purchased from ATCC (Manassas, Virginia, USA). Cell culture media were from Gibco Life Technologies (Grand Island, New York, USA).

**Animal model of alcoholic steatohepatitis**

Mouse protocols were approved by the relevant authorities and all procedures were performed in accordance with the institutional guidelines. Also, 5–6-week-old female C57Bl/6 mice were purchased from Charles River Laboratories. The animals were housed under specific pathogen-free conditions. Animals were allowed to acclimatise for 7–10 days before the start of the experiments. Mice were subjected to a high-fat LDC (44% fat-derived, 16% protein-derived and 40% carbohydrate or ethanol-derived calories). Ethanol was introduced after 5 days, starting 2% (v/v) ethanol, and was then increased every other day to a final concentration of 6.5% (v/v) ethanol on the 10th day of the experiment. Control animals received LDC without any alcohol. As outlined in figure 1, we started to inject the x1/2pal-i1 pepducin (2.5 mg/kg, every other day) either concomitantly with the introduction of ethanol (day 5) in a preventative mode or after mice had been on alcohol containing LDC for 4 weeks. The latter therapeutically treated mice were injected with x1/2pal-i1 (5 mg/kg) every day for the remaining 7 days of the experiment. The vehicle (10% dimethyl sulfoxide (DMSO)) was injected as a control. Mice received a single low-dose injection of endotoxin (2.5 mg/kg) intraperitoneally 24 h before sacrificing. Human alcoholic steatohepatitis (ASH) is characterised by high blood endotoxin levels, which cannot be observed in the murine model. Upon completion of the study, the animals’ weight was taken before terminal anaesthesia with ketamine/xylazine. Cardiac punctures were performed to collect blood. Livers were weighed and either embedded in paraffin, frozen or prepared for RNA extraction, triglyceride extraction or MPO assessment.

**Measurement of serum and liver cytokine levels**

Serum CXCL8 levels were measured by ELISA. Liver cytokine mRNA levels were measured by real-time quantitative SYBR-green PCR. Primers were designed as follows: tumour necrosis factor (TNF): (forward) and (reverse), CXCL1: (forward) and (reverse), IL-1β: (forward) and (reverse) (reverse).

MPO was extracted from liver tissue and measured by ELISA. Aspartate aminotransferase (AST) and alanine aminotransferase (ALT) were measured by a colorimetric enzyme activity assay (Sigma Aldrich).

**Histological assessments of murine liver tissue**

Liver sections were embedded in paraffin and cut into 5–15-μm-thick slices by subsequent staining with H&E. Liver morphology was then assessed microscopically. To compare the numbers of neutrophils in the livers of pepducin-treated versus vehicle-treated animals, paraffin was removed and slides were stained with naphthol AS-D chlorocetate (Sigma Aldrich). Neutrophil numbers were counted in four representative microscopic fields from 10 individual mice per group.

For the assessment of liver tissue lipid content, frozen sections were stained with Oil-red-O (Sigma Aldrich). Accumulated lipid appears as bright red droplets in the liver tissue and the amount of these bright red areas was then compared microscopically.

**Measurement of liver triglycerides from liver tissue**

The lipid liver fraction was extracted by the Folch method. Briefly, 100 mg of liver tissue were homogenised in the presence of 3:1 chloroform:methanol. After filtration and two washing steps, the lipid extracts were analysed by a photometric assay (Roche/Hitachi).

**Immunoblotting of caspase-1**

Snap frozen liver samples were homogenised and separated by 12% sodium dodecyl sulfate polyacrylamide gel electrophoresis. After blotting, membranes were probed by a polyclonal anti-caspase-1 p20 antibody (Santa Cruz, clone M19).

**In vitro neutrophil chemotaxis**

Human neutrophils were obtained from EDTA anticoagulated blood of healthy volunteers. After density gradient
centrifugation, remaining red blood cells were lysed. Cell preparations typically yielded >95% neutrophils with almost 100% viability.

Neutrophil migration was measured using a 48-well microchemotaxis chamber (Neuroprobe, Gaithersburg, Maryland, USA). Cells were allowed to migrate for 30 min into a 5 μm pore-sized nitrocellulose membrane and were stained subsequently. Migration depth was quantified microscopically by evaluating the difference between cells that did not migrate and the leading front of neutrophils.

Cell culture
Hep3b cells were cultured in Dulbecco’s modified Eagle’s medium (DMEM) cell culture media containing 10% fetal bovine serum (FBS). HepG2 cells were cultured in Roswell Park Memorial Institute medium containing 10% FBS. For some experiments, Hep3b cells were stimulated with ethanol or TNFα. IL-8 levels were measured in the supernatants of Hep3b cells and subjected to chemotaxis assays. To indicate generalisability, some of the experiments were repeated in HepG2 cells. To assess potential cytotoxicity of pepducins on hepatocytes, we performed MTT tests. Hep3B cells were incubated with DMEM/0.2% FBS for 48 h in 96-well plates. Thereafter, cells were incubated with different concentrations of the x1/2pal-i1 pepducin, control medium or doxorubicin IC50 (300 nM) for another 48 h. MTT (10 μg/well) was added and incubated for another 5 h. After the addition of 100 μL DMSO, the extinction was measured photometrically (wavelength 550 nm).

RESULTS
High-fat diet supplemented with alcohol induces experimental ASH
We administered C57Bl/6 mice with an LDC high-fat diet, supplemented with ethanol, followed by intraperitoneal lipopolysaccharide (LPS) injection 24 h prior to sacrifice as detailed in figure 1A. H&E stainings of liver sections from mice receiving an ethanol-enriched LDC diet (LDC30%), but not LDC

Figure 1 x1/2pal-i1 treatment inhibited the development and progression of alcoholic liver disease. (A) Experimental approach. Mice received the Lieber–DeCarli (LDC) diet. Pepducin therapy was commenced either with the introduction of ethanol (day 5; ‘preventative setting’, 2.5 mg/kg x1/2pal-i1 subcutaneous every other day) or after mice had established disease (day 29, ‘therapeutic setting’, 5.0 mg/kg x1/2pal-i1 subcutaneous once a day). On day 35, mice received LPS (2.5 mg/kg intraperitoneal) and were assessed 24 h later. (B) x1/2pal-i1 prevents development of liver steatosis. Representative liver sections stained with H&E (n=15). (C) Therapeutic x1/2pal-i1 reverts liver steatosis. Representative liver sections stained with H&E (n=15). (D) Histological disease activity. H&E-stained sections for hepatocyte ballooning, leucocyte infiltration and Mallory–Denk bodies, Oil-red-O for steatosis. Statistical analysis: Mann–Whitney U after Kruskal–Wallis; n=15 per group; *p<0.05 (LDC vs LDC-EtOH); †p<0.05 (LDC-EtOH vs LDC-EtOH-x1/2pal-i1). (E) x1/2pal-i1 treatment prevents from alcohol-induced mortality. Mice received 2.5 mg/kg of x1/2pal-i1, scram-i1 (2.5 mg/kg) or vehicle control every other day. Statistical analysis: Mantel–Cox test p<0.0001. n=15. (F) x1/2pal-i1 therapy reduces alcohol-induced mortality and prevents from weight loss. Mice received 5 mg/kg of x1/2pal-i1, scram-i1 or vehicle control every day from day 29 until the conclusion of the experiment. The animals’ weight was taken every other day and weight curves were compared. Statistical analysis: Mantel–Cox test p<0.0001. n=15 (n=7 for scram-i1).
diet alone (LDC\textsuperscript{Ctrl}), resulted in severe steatohepatitis, characterised by neutrophilic infiltration and hepatocyte ballooning, lipid accumulation and Mallory–Denk bodies (figure 1B–D). While macrovesicular steatosis was observed in most of the liver sections of LDC\textsuperscript{Ctrl} mice (figure 1B, C), addition of alcohol resulted in a macrovesicular pattern with hepatocyte ballooning in LDC\textsuperscript{EtOH} mice (figure 1B, C). These features (figure 1D) closely resemble the characteristic histological picture observed in human ASH.\textsuperscript{35–37} LDC\textsuperscript{EtOH} mice exhibited significant mortality and developed cachexia (figure 1E, F) over the course of the experiment, even prior to LPS administration (figure 1F). At the end of the experiment, the liver-to-body weight ratio in LDC\textsuperscript{EtOH} mice was more than twofold higher compared with LDC\textsuperscript{Ctrl} mice (figure 2A, B), which was associated with profoundly increased AST and ALT serum levels, reflecting increased hepatocyte injury (figure 2A, B). Altogether, this experimental model faithfully recapitulates many critical features of human ASH.

**Figure 2.** x1/2pal-i1 treatment normalises liver–body weight ratio and prevents from hepatic necrosis. (A) Prophylactic and therapeutic x1/2pal-i1 administration result in a normal liver–body weight ratio. Statistical analysis: Mann–Whitney U after Kruskal–Wallis. n=15 per group (n=7 for scram-i1 experiment). (B and C) Prophylactic and therapeutic x1/2pal-i1 treatment prevents from liver cell necrosis. Aspartate aminotransferase (AST) and alanine aminotransferase (ALT) levels as markers of liver cell necrosis. Statistical analysis: Mann–Whitney U after Kruskal–Wallis. n=15 per group (n=7 for scram-i1 experiment).
of Cxcl1, Tnf and Il1β was increased in LDCEtOH compared with LDCCtrl mice (figure 3B–D). Increased Il1β expression in LDCEtOH mice was prevented by x1/2pal-i1 administration (figure 3C), and even when administration was delayed until disease had established (figure 3C). CXCL1 serum levels were similarly reduced (figure 3A) by x1/2pal-i1, while Cxcl1 and Tnf mRNA levels only trended lower (figure 3B, C). These data show that x1/2pal-i1 treatment of experimental ASH normalises hepatic expression of inflammatory cytokines, which are characteristically induced in human ASH.

Patients suffering from ASH exhibit increased caspase-1 expression.36 37 Similarly, chronic ethanol exposure in mice induces pro-caspase-1 expression.37 To test whether blocking CXCR2 receptors by x1/2pal-i1 decreases inflammasome activation, we probed liver tissue lysates for caspase-1, a hallmark of inflammasome activity. Our results indicate that the blockade of CXCR2 reduces the amount of pro-caspase-1 in x1/2pal-i1-treated mice, whereas DMSO and scram-1 treatment had no effect. Interestingly, no differences in the amount of active caspase-1 (p20) were detected, which could, however, be a result of a relatively short half-life of cleaved caspase-1 (figure 3E).37 38

x1/2pal-i1 pepducin abrogates neutrophil accumulation in experimental ASH

Hepatic neutrophil accumulation is a characteristic feature of ASH and is similarly observed in experimental ASH (figure 4A, B), as revealed through staining with naphthol AS-D chloroacetate, a chromogenic substrate of the specific esterases of neutrophilic granules. Administration of x1/2pal-i1 to LDCEtOH mice reduced the numbers of neutrophils per high-power field to those observed in LDCCtrl mice (figure 4A). Activated neutrophils have a short half-life and high turnover in inflamed tissue. Delayed CXCR1/2 blockade by x1/2pal-i1 likewise resulted in a similar complete abrogation of neutrophil accumulation in LDC3O4 mice (figure 4B), whereas scram-1 had no effect on neutrophil numbers. Finally, liver MPO levels were decreased to baseline levels in x1/2pal-i1-treated LDC3O4 mice (figure 4C).

Hepatocyte Hep3b and HepG2 cells secrete CXCL8 upon alcohol exposure, attracting neutrophils in vitro

In addition to liver Kupffer cells, hepatocytes also contribute profoundly to inflammatory cytokine secretion in the liver. We
chose the human Hep3b and HepG2 cell lines to test the hypothesis that ethanol exposure directly induces CXCL1 secretion in hepatocytes. As shown in figure 4D, 1.5 and 15 mM ethanol in culture medium resulted in a substantial increase in CXCL8 release from Hep3b and HepG2 cells. Supernatants from ethanol-exposed Hep3b and HepG2 (data not shown) cells exhibited chemotactic activity to neutrophils in vitro, and this activity was completely blocked either by a neutralising anti-CXCL8 monoclonal antibody and x1/2pal-i1, whereas a blocking anti-CXCL17 antibody had no effect (figure 4E). In another set of experiments, human neutrophil chemotaxis towards supernatants of ethanol-treated HepG2 cells was performed in the presence of scram-i1. Scram-i1 had no effect on the induction of chemotaxis towards the supernatants. CXCL8 (IL-8) was neutralised by adding antibodies directed against CXCL8. CXCR1 and CXCR2 receptor signalling was inhibited by x1/2pal-i1. The nonsense pepducin scram-i1 had no effect on neutrophil chemotaxis induced by HepG2 cell supernatants. Chemotaxis experiments were performed in modified Boyden chambers. Statistical analysis: Mann–Whitney U after Kruskal–Wallis, n=3. LDC, Lieber–DeCarli diet.

**Figure 4** Pepducin treatment inhibits neutrophil infiltration and activation. (A and B) x1/2pal-i1 prevents accumulation of neutrophils in liver tissue sections of alcoholic mice. To compare the numbers of neutrophils, histological sections were stained with naphthol AS-D chloroacetate. Neutrophil numbers were counted in four representative microscopic fields from 10 individual mice per group. (C) Ethanol-induced neutrophil chemotaxis was CXCL8 dependent. CXCL8 (IL-8) was neutralised by adding antibodies directed against CXCL8. CXCR1 and CXCR2 receptor signalling was inhibited by x1/2pal-i1. The nonsense pepducin scram-i1 had no effect on neutrophil chemotaxis induced by HepG2 cell supernatants. Chemotaxis experiments were performed in modified Boyden chambers. Statistical analysis: Mann–Whitney U after Kruskal–Wallis, n=3. LDC, Lieber–DeCarli diet.

x1/2pal-i1 treatment prevents lipid accumulation in experimental ASH

Hepatocyte lipid accumulation is a cardinal feature of ASH.\(^3\) LDC\(^{Ctrl}\) mice exhibited little fat accumulation in their livers as determined by Oil-red-O staining, whereas the number of hepatic lipid vesicles was drastically increased in LDC\(^{EtOH}\) mice (figure 5A). Administration of x1/2pal-i1 completely prevented this increase in lipid vesicles in LDC\(^{EtOH}\) mice (figure 5A). Triglycerides are thought to represent the primary lipid species in ballooned hepatocytes in ASH.\(^3\) We, therefore, quantified triglycerides within the lipophilic fraction extracted from whole murine livers via the Folch method. Triglyceride content in LDC\(^{EtOH}\) mice was significantly increased compared with LDC\(^{Ctrl}\) mice (figure 5B). x1/2pal-i1 completely prevented this increase in LDC\(^{EtOH}\) mice to levels below those observed in LDC\(^{Ctrl}\) mice (figure 5B). Compared with vehicle-treated LDC\(^{Ctrl}\) mice, both Oil-red-O+ lipid vesicles and liver triglyceride content trended higher upon x1/2pal-i1 treatment (figure 5A, B). However, this was not associated with increased liver/body weight ratio (figure 2A). To exclude potential toxicity of x1/2pal-i1, higher doses (5 mg/kg, every other day) were administered in mice on a normal diet. We did not observe an increase in ALT (figure 5D) or AST (figure 5E) levels in serum nor was the liver-to-body weight ratio altered compared with vehicle-treated mice (figure 5C).

**CONCLUSION**

Here, we demonstrate that CXCR1/2 signalling plays a central role in an experimental model of ASH, which phenocopies core features of human ASH. Remarkably, late therapeutic blockade of CXCR1/2 signalling with the pepducin x1/2pal-i1 in established, severe disease, was effective in preventing further mortality and reversing liver inflammation (figure 6).
Neutrophil infiltration in the liver is a classic hallmark of human ASH, and correspondingly observed in experimental ASH. Efficacy of x1/2pal-i1 in experimental ASH was associated with a profound reduction of neutrophil granulocytes infiltrating the liver, signifying a critical role of neutrophils in this disease. More specifically, neutrophils are recruited to sites of invading pathogens or tissue injury within minutes. Apart from removing pathogens by phagocytosis, neutrophils, however, release an array of mediators such as MPO, matrix metalloproteases, serine proteases, elastases, cathepsin G and nicotinamide adenine dinucleotide phosphate oxidase, which can promote tissue damage. Hepatic neutrophil infiltration has also been demonstrated in other experimental models of liver disease such as ischaemia-reperfusion, ConA-induced and CCl4-induced hepatitis. Interestingly, in a murine sepsis model we have previously shown that CXCR1/2 blockade results in suppression of the systemic inflammatory response syndrome without yielding higher bacterial loads. Blocking CXCR1 and CXCR2 receptors with pepducins did not suppress chemotaxis induced by other neutrophil chemoattractants; therefore, x1/2pal-i1 may be rather considered immunomodulatory rather than immunosuppressive. Blocking neutrophil emigration from the blood stream into liver tissue and preventing their overzealous activation via x1/2pal-i1 may represent an intervention at a critical checkpoint that shuts off organ destruction, resulting in protection from acute liver failure, and may also prevent further consequences such as fibrosis.

CXCR1/2 blockade via x1/2pal-i1 was associated with normalisation of increased hepatic IL-1β expression during...
experimental ASH. IL-1β secretion due to caspase-1 activation in Kupffer cells has previously been shown to induce hepatic steatosis, and blockade with recombinant IL-1 receptor antagonist (IL-1Ra, Anakinra) in experimental steatohepatitis attenuated liver inflammation. 35 In our model of experimental ASH, we show that blocking CXCR2 receptors downregulates the expression of pro-caspase 1. Interestingly, Marques et al 46 reported that DP2156a, an allosteric small peptide inhibitor of CXCR1/2, reduced the number of infiltrating neutrophils in paracetamol-induced liver injury, but this was not associated with a reduction in liver injury. The latter required co-blockade of the formyl peptide receptor 1, 48 which could either point to differences in pathogenesis of these liver injury models or the specific potency of x1/2-pal-i1 in blocking CXCR1/2 signalling. The IL-8 receptors CXCR1 and CXCR2 are not only expressed on leucocytes but can also be found on human hepatocytes under pathological conditions. 32 45 Upon ischaemia/reperfusion injury, CXCR1 is upregulated on murine hepatocytes and appears involved in hepatocyte proliferation. Genetic deletion of CXCR1 or its blockade by repertaxin resulted in a significant decrease of BrdU incorporation into proliferating hepatocytes. 47 In this model of ischaemia/reperfusion injury, which indeed causes one single assault to the liver tissue, immigrating neutrophils are a good prognostic factor and facilitate repair and regeneration. 45 In ASH, in contrast, with alcohol present as a constant noxa, repair and regeneration might result in hyperproliferation and consequent development of neoplasia. Liver cirrhosis bears a high risk for developing hepatocellular carcinoma, and it might be speculated that increased hepatocyte CXCR1 expression consequent to liver injury could promote pathological hyperproliferation. Indeed, CXCL5, another CXCR2 ligand, has been shown to be associated with a high neutrophil load and poor outcome in hepatocellular carcinoma. 48 CXCR1 and CXCR2 also promote neoangiogenesis in tumour tissue. 30 31 Hence, blockade of CXCR1/2 signalling may not only beneficially affect acute inflammation and the development of steatosis, but could potentially also decrease the risk of hepatocellular carcinoma.

Treatment for severe ASH includes supportive measures to reduce ascites, prevent infections, treat hepatic encephalopathy (antibiotics, lactulose), ascertain sufficient protein intake and maintain serum albumin levels, and supplementation of vitamins (antibiotics, lactulose), ascertain sufficient protein intake and maintain serum albumin levels, and supplementation of vitamins (antibiotics, lactulose), ascertain sufficient protein intake and maintain serum albumin levels, and supplementation of vitamins (antibiotics, lactulose), ascertain sufficient protein intake and maintain serum albumin levels, and supplementation of vitamins (antibiotics, lactulose), ascertain sufficient protein intake and maintain serum albumin levels, and supplementation of vitamins (antibiotics, lactulose), ascertain sufficient protein intake and maintain serum albumin levels, and supplementation of vitamins (antibiotics, lactulose), ascertain sufficient protein intake and maintain serum albumin levels, and supplementation of vitamins (antibiotics, lactulose), ascertain sufficient protein intake and maintain serum albumin levels, and supplementation of vitamins (antibiotics, lactulose), ascertain sufficient protein intake and maintain serum albumin levels, and supplementation of vitamins (antibiotics, lactulose), ascertain sufficient protein intake and maintain serum albumin levels, and supplementation of vitamins. 18 Wieser V, et al. Gut 2016;65:1–9. doi:10.1136/gutjnl-2015-310344


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