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procedures were implemented. 17 (17%) of these patients came from Wuhan, the city hardest hit by the outbreak (appendix 2, p 3). One patient died after 19 days in hospital because of multiple organ failure.

At follow-up, none of our participants had clinical symptoms suggestive of SARS-CoV-2 infection. By contrast, five (2%) of 250 patients without cirrhosis and six (16%) of 38 health-care workers were diagnosed with COVID-19 by casual testing in our ward. Several outpatients complained about mild gastrointestinal and respiratory symptoms, which were resolved by rest, proton pump inhibitors, and probiotics (appendix 2, p 2).

As an additional comparator, we calculated the incidence of COVID-19 among 101 inpatients with decompensated cirrhosis at five other hospitals in Wuhan over the same period, where our approach had not been implemented. 17 (17%) of these 101 patients were diagnosed with COVID-19 (p=0·018 vs our group; appendix 2, p 4). This simple approach could be an effective means of preventing COVID-19 in patients with decompensated cirrhosis. However, our sample size is small and larger studies are needed.

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COVID-19 and the liver: little cause for concern

The largest study on COVID-19 to date showed that the prevalence of elevated aminotransferases and bilirubin in people faring worst was at least double that of others. Although clinically significant liver dysfunction was not quantified, this and other studies have led some to suggest that this finding might present clinical challenges. Close inspection of the available data supports a higher prevalence of abnormal aminotransferase levels in severe COVID-19 disease, but these studies actually suggest that clinically significant liver injury is uncommon, even when data for the most severely ill patients are selected (table). Although high levels of positive end expiratory pressure can contribute to hepatic congestion by increasing right atrial pressure and impeding venous return, data suggest that many patients admitted to hospital with COVID-19 have liver blood test abnormalities without mechanical ventilation. Furthermore, the distribution of aminotransferase levels among patients with COVID-19 do not support hypoxic hepatitis being a common phenomenon, according to the published literature. Drug-induced liver injury is a possible contributing factor to the observed abnormal liver blood test abnormalities after therapeutics begin and should be considered by clinicians, but mild liver test derangement is present at baseline in many patients with COVID-19 before significant medication use. Several studies have reported elevated levels of creatinine kinase and lactate dehydrogenase or myoglobin in association with COVID-19 severity (table). It is therefore possible that aminotransferase elevations do not necessarily arise from the liver alone and that COVID-19 infection might induce a myositis similar to that observed in severe influenza infections.

It has been proposed that COVID-19 causes direct liver injury via a viral hepatitis, but we believe that there are alternative explanations. First, the derangement of liver function is clearly mild. Second, when liver function tests for patients with different durations of symptoms are examined, there is no evidence that later presentation is associated with greater liver function derangement. The only post-mortem liver biopsy from a patient with COVID-19 showed only microvesicular steatosis, a common finding in sepsis. Most
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importantly, other respiratory viruses produce similar elevations of liver function biomarkers, which is thought to relate to hepatic damage from immune interactions involving intrahepatic cytotoxic T cells and Kupffer cells.3 This phenomenon waxes and wanes in parallel with respiratory viral disease and in the absence of hepatic viral replication, which might explain why worse outcomes were not seen in the 42 patients with chronic liver disease and COVID-19 who had outcome data (table).

Hepatic dysfunction in severe COVID-19 is accompanied by greater activation of coagulative and fibrinolytic pathways, relatively depressed platelet counts, climbing neutrophil counts and neutrophil to lymphocyte ratios, and high ferritin levels.4 Although these markers are seen as non-specific markers of inflammation, we believe that they fit the paradigm of disease severity coinciding with a failure of innate immune regulation.5 Such unbalanced immunity favours NETosis and coagulation activation and possibly also alters systemic iron metabolism secondary to macrophage activation.6 Notably, this alteration of immune balance occurs with increased age, and older patients might therefore be expected to fare worse, with a greater reliance on this pathway.7

Clinicians cannot be complacent about the risks of COVID-19 in patients with chronic liver diseases and cirrhosis, because these patients have poor immune function and worse outcomes from acute respiratory distress syndrome than the rest of the critically ill population.8 However, we believe that collateral liver damage from virally induced cytotoxic T cells and the induction of a dysregulated innate immune response is a more probable explanation for the association between deranged liver markers and COVID-19 disease severity. Furthermore, we suspect that what is termed COVID-19-induced hepatic damage is predominantly a clinical distraction. We urge clinicians and the scientific community to focus attention towards viral control and modulating innate immune dysfunction in COVID-19.

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