<table>
<thead>
<tr>
<th><strong>Title</strong></th>
<th>Clinical significance of hepatic derangement in severe acute respiratory syndrome</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Author(s)</strong></td>
<td>Chan, HLY; Kwan, ACP; To, KF; Lai, ST; Chan, PKS; Leung, WK; Lee, N; Wu, A; Sung, JJY</td>
</tr>
<tr>
<td><strong>Citation</strong></td>
<td>World Journal Of Gastroenterology, 2005, v. 11 n. 14, p. 2148-2153</td>
</tr>
<tr>
<td><strong>Issued Date</strong></td>
<td>2005</td>
</tr>
<tr>
<td><strong>URL</strong></td>
<td><a href="http://hdl.handle.net/10722/162824">http://hdl.handle.net/10722/162824</a></td>
</tr>
<tr>
<td><strong>Rights</strong></td>
<td>This work is licensed under a Creative Commons Attribution-NonCommercial-NoDerivatives 4.0 International License.</td>
</tr>
</tbody>
</table>
Clinical significance of hepatic derangement in severe acute respiratory syndrome

Henry Lik-Yuen Chan, Ambrose Chi-Pong Kwan, Ka-Fai To, Sik-To Lai, Paul Kay-Sheung Chan, Wai-Keung Leung, Nelson Lee, Alan Wu, Joseph Jao-Yiu Sung

INTRODUCTION

Severe acute respiratory syndrome (SARS) is caused by a novel coronavirus (SARS-coronavirus, SARS-CoV) infecting primarily the lung and the enteric tract[1-9]. Up to August 2003, there were 8,422 reported cases worldwide and 916 cases died of this condition[10]. Although the outbreak of SARS is currently under control, the source of SARS-CoV has not been identified and the threat of SARS returning in winter persists.

Case series in Hong Kong and Toronto indicated that SARS is not merely a respiratory disease. Diarrhea and bleeding diathesis had also been reported in patients infected by SARS-CoV[11,12]. In our previous report, SARS-CoV was found in biopsy of small intestine and colon of patients with diarrhea[13]. Deranged liver functions were reported in 22-56% of patients at the time of hospital admission[13-17]. In a previous study, it has also been suggested that co-infection with hepatitis B virus (HBV) is associated with more severe respiratory disease[18]. The cause of impaired liver function, its clinical implication and association with HBV co-infection have not been fully explored.

In this study, we follow the natural course of hepatic involvement in SARS. The impact of liver derangement and chronic HBV infection on the clinical outcome of SARS patients is revisited.

MATERIALS AND METHODS

Patients

Patients in the present study were collected from a university medical center (Prince of Wales Hospital) and a community hospital (Princess Margaret Hospital) designated to look after SARS patients during the outbreak in Hong Kong. All patients fulfilled the case definitions of SARS by the World Health Organization, i.e., temperature above 38 ℃, difficulty in breathing and cough, pneumatic changes on chest X-ray or high-resolution computed tomography, and contact history with SARS patients[19]. All patients were initially treated with empirical antibiotics including cefotaxime and
clarithromycin (or levoﬂoxacin) to cover common pathogens causing community-acquired pneumonia. Oseltamivir (Tamiflu) was also given to treat possible inﬂuenza infection when little was known about SARS during the early phase of the outbreak. If fever persisted for more than 48 h, all patients received corticosteroids and ribavirin treatment. The choice of corticosteroids was intravenous hydrocortisone 100 mg every 8 h or oral prednisolone 1 mg/kg body weight per day. Ribavirin was given at 400 mg every 8 h intravenously or 1 200 mg thrice a day orally. Pulse intravenous methylprednisolone (500-1 000 mg/d), up to a maximum dose of 3 g, was given when there were signs of radiological or clinical deteoration. Patients, with oxygen saturation that fell below 90% at room air, were offered supplementary oxygen through nasal cannula. Those who required more than 4 L/min oxygen would be transferred to intensive care unit (ICU) for close monitoring. Mechanical ventilation by CPAP was implemented, when patients could not achieve 90% oxygen saturation despite receiving 5 L/min oxygen or more. All patients were kept in hospital for monitoring for at least 3 wk before discharge. Liver enzymes were checked every 1-3 d during hospital stay, on discharge and on follow-up visits. Clinical outcomes were assessed at least 8 wk after the admission of patients.

**Serological assays**

The level of anti-coronavirus IgG antibody was measured by immunonoﬂuorescence assay. Paired sera from acute (taken within 7 d after the onset of fever) and convalescent (taken 14-21 d after the onset of fever) blood samples were tested at serial two-fold dilutions starting from 1:40. Positive serological evidence of coronavirus infection was deﬁned as either having a seroconversion or ≥four-fold rise in antibody titer. Hepatitis B surface antigen (HBsAg) and hepatitis C antibodies (anti-HCV) were tested by commercially available enzyme-linked immunosorbent assay kits (Abbott GmBH Diagnostika, Wiesbaden-Delkenheim, Germany). Hepatitis B e antigen (HBeAg) and antibodies to hepatitis B e antigen (anti-HBe) were measured by ELISA (Sanofi Diagnostics, Pasteur, France).

**Data analysis**

Continuous variables were expressed as mean±SD for normal distribution data and median (range) if the distribution was skewed. Statistical analysis was performed by SPSS (version 11.0, Chicago). Categorical variables were compared by χ² test and continuous variables by Student’s t test or Mann-Whitney U test as appropriate. Adverse clinical outcomes were deﬁned as need of oxygen desaturation requiring oxygen supplementation, ICU admission, mechanical ventilation, liver decompensation and mortality. Liver decompensation was deﬁned as development of hepatic encephalopathy associated with elevated serum bilirubin (>51 mmol/L) and prolonged prothrombin time (>16 s). As the reference ranges of alanine aminotransferase (ALT) levels were different between the two hospitals, ALT levels were expressed as folds of increase above the upper limit of normal (ULN) in individual laboratories. The relationships of peak ALT levels and various adverse clinical outcomes were compared by receiver operator characteristic curve. Baseline clinical characteristics with a P value <0.1 for adverse clinical outcomes on comparing patients with high peak ALT levels vs. those with lower ALT levels were adjusted by multivariate logistic regression analysis. All statistical tests were two-sided. P value <0.05 was statistically significant.

**RESULTS**

**Clinical characteristics**

Two hundred and ninety-four patients including 126 male (43%) and 168 female were included in this study. The median age of this cohort was 36 years, range 12-83 years. Two hundred and forty-three patients had paired blood samples checked for SARS-CoV serology and all had positive results. These patients were admitted on the third (range 0-11) day after the onset of fever. Thirty (10%) patients were found to have positive HBsAg and 214 patients had negative HBsAg. In 50 cases, HBsAg status was not checked during hospitalization. All patients in the Prince of Wales cohort had negative anti-HCV antibodies, and 7 of 12 HBV-infected patients had positive HBeAg. Anti-HCV and HBeAg status were not routinely monitored in the Princess Margaret Hospital cohort. Lamivudine (100 mg/d) was commenced in 20 of the 30 chronic hepatitis B patients on or before the commencement of corticosteroid treatment and was continued thereafter. Two chronic hepatitis B infected patients had co-existing liver cirrhosis. One of them had inoperable multi-focal hepatocellular carcinoma and the other was admitted for bleeding esophageal varicale. Forty-one (14%) patients had other co-morbid illnesses including hypertension (12), diabetes mellitus (5), end-stage renal failure (2), chronic rheumatic heart disease (2), ischemia heart disease (2), sick sinus syndrome (1), atrial fibrillation (1), asthma (1), chronic obstructive airway disease (1), bronchiectasis (1), old pulmonary tuberculosis (1), previous cerebrovascular accident (1), autism (1) and pregnancy (1).

Overall, 141 (48%) patients had oxygen desaturation, 50 (17%) required admission to ICU, 33 (11%) patients required mechanical ventilation, and 27 (9%) patients died. None of the patients developed liver decompensation after contracting SARS. All mortalities were due to respiratory failure related to SARS with or without sepsis and multi-organ failure. The outcome of the studied cohort is summarized in Figure 1.

**Liver enzyme derangement on admission**

Seventy (24%) patients, including 15 chronic hepatitis B-infected patients, had elevated ALT levels on admission. The median ALT levels on admission was 0.55 (0.16-26.09) times upper limit of laboratory normal. The proportion of patients with different ALT levels, serum bilirubin and prothrombin time on admission are shown in Figure 2. Two chronic hepatitis B infected patients were admitted with icteric flare-up of chronic hepatitis B on lamivudine treatment and contracted SARS during their hospital stay. They had elevated serum bilirubin to more than 150 mmol/L and one of them had prolonged prothrombin time to 17 s. All other patients who had elevated ALT levels had normal
serum bilirubin levels and had no evidence of hepatic decompensation.

**Progression of liver derangement on follow-up**

During the course of illness, 204 (69%) patients, including 23 HBsAg-positive patients, had elevated ALT levels. Majority of patients had ALT levels elevated at day 5-7 from fever onset and ALT peaked at the end of second week (Figure 3). The median peak ALT levels was 1.53 (0.28-316.25) times ULN. The proportion of patients with different peak ALT levels, serum bilirubin and prothrombin time are shown in Figure 2. Twenty-eight (9.5%) patients had ALT raised to over 5× ULN; among them 7 patients had elevated serum bilirubin (median 83 mmol/L, range 48-231 mmol/L) and 10 patients had elevated prothrombin time (median 20 s, range 14-56 s). Among the 138 patients in the Prince of Wales Hospital cohort, the median peak alkaline phosphatase (ALP) level was 0.78 (range 0.31-19.38) times ULN and 40 (29%) patients had experienced elevated ALP during the course of illness. None of these patients developed hepatic encephalopathy. Seven of the 28 (25%) patients died of SARS and multi-organ failure. On the other hand, only 5 of 89 (6%) patients who had persistently normal ALT died.

The area under the receiver operator characteristic curve of peak ALT for oxygen desaturation, need of ICU care, mechanical ventilation and mortality were 0.70 (95%CI 0.64-0.75; *P* <0.001), 0.72 (95%CI 0.65-0.79; *P* <0.001), 0.71 (95%CI 0.62-0.80, *P* <0.001) and 0.65 (95%CI 0.54-0.76, *P* <0.001) respectively.

---

**Figure 1** Clinical outcomes of patients included in the study. ALT, alanine aminotransferase. *One patient did not have serial ALT results. Two patients did not have follow-up ALT results after discharge.*

---

**Figure 2** A: Proportion of patients with different ALT levels at initial visit (*n* = 294), at peak ALT (*n* = 293; 1 missing data) and on last follow-up (*n* = 264; 27 patients died, 3 missing data). ALT normal, 1-2× ULN, 2-5× ULN, >5× ULN. B: Proportion of patients with different serum bilirubin levels at initial visit (*n* = 294) and at peak bilirubin (*n* = 293, 1 missing data). <35 µmol/L, 35–51 µmol/L, >51 µmol/L. C: Proportion of patients with different prothrombin time at initial visit (*n* = 293, one missing data) and at peak prothrombin time (*n* = 283, 11 missing data). <12 s, 12–16 s, >16 s.
P = 0.011), respectively. Using the coordinate of peak ALT >5× ULN, the sensitivity and specificity for any adverse outcomes were 15-27% and 93-95% respectively. Patients who had peak ALT over 5× ULN had significant male predominance, more co-existing comorbid conditions, more chronic hepatitis B patients and marginally higher serum creatinine levels as compared to those who had lower peak ALT levels (Table 1). Using peak ALT >5× ULN as a cutoff and after adjusting these potential confounding factors, the odds ratio of peak ALT >5×ULN for oxygen desaturation was 3.24 (95%CI 1.23-8.59, P = 0.018), ICU care was 3.70 (95%CI 1.38-9.89, P = 0.009), mechanical ventilation was 6.64 (95%CI 2.22-19.81, P = 0.001) and death was 7.34 (95%CI 2.28-24.89, P = 0.001).

Excluding 27 patients who died and 3 patients who had no follow-up ALT results, 84 of 264 (32%) patients had persistently normal ALT levels during the entire course of illness. Among the 180 patients who had elevated ALT levels, 128 (71%) had ALT subsequently normalized and 37 (21%) patients had ALT on downward trend at the last follow-up visit. The remaining 8% of patients still had elevated ALT levels on the last follow-up.

**Co-infection with hepatitis B virus**

Two of the 30 HBsAg-positive patients died despite lamivudine treatment. One patient was admitted for icteric flare-up of chronic hepatitis B and the other had liver cirrhosis admitted for esophageal variceal bleeding. Both patients acquired SARS during their hospital stay. Seven patients had persistently normal ALT levels throughout the admission and follow-up visits (six were on lamivudine). Twenty-one HBsAg-positive patients had elevated ALT during the SARS illness. Among them, 10 patients (6 on lamivudine) had transient elevation of ALT which subsequently returned to normal levels, 8 (5 on lamivudine) had declining levels of ALT and 3 patients (1 on lamivudine) had persistently elevated ALT levels. In this series, co-infection with viral hepatitis B was not associated with higher peak ALT level, increased risk of oxygen desaturation, ICU admission, mechanical ventilation or mortality (Table 2).

**DISCUSSION**

Although SARS is primarily a pulmonary disease, liver derangement was commonly observed[13-17]. In this study, approximately a quarter of patients who had elevated ALT on admission, and a further 45% of patients who had normal ALT on admission had ALT elevation during the course of illness. In a majority of patients ALT levels started to elevate towards the end of first week and peak at the end of second week. High peak ALT level appears to be an independent predictor of more severe illness and worse clinical outcome. Most patients however had transient elevation of ALT, which normalize spontaneously with the recovery of SARS.

The underlying cause for ALT elevation was uncertain but several mechanisms were worth considering. Direct pathogenic effect of SARS-CoV on the liver is unlikely due to the failure of identification of SARS-CoV or specific hepatitis features in the liver at autopsy in previous reports[20-23]. Elevated ALT might be related to the prescribed medications including antibiotics and high dose ribavirin. As a quarter of patients who had elevated ALT levels on admission well before the prescription of drugs, drug treatment should not be the cause of liver enzyme derangement in these patients. Furthermore, many patients had ALT >5× ULN. This is exceedingly rare with antibiotic and ribavirin given at these dosages[20]. Although chronic hepatitis C virus infection has only been excluded in about half of the cases in this series (the Prince of Wales Hospital cohort), the prevalence of chronic hepatitis C is very low among the general population in Hong Kong.
We believe that the elevation of liver enzyme is a reactive response towards SARS-CoV infection[6]. The hepatic acute phase response involving cytokine release from inflammatory cells is a defense reaction of the body against the causative agent to protect the vital functions of the liver[26]. The liver enzyme elevation in SARS is not typical of cholestasis secondary to sepsis as majority of patients did not have accompanied elevation of ALP. This is usually a transient reaction and therefore majority of patients have ALT levels returned to normal after recovery.

Although SARS-CoV is not a direct cause of liver injury, results from this study indicated that gross liver enzyme elevation, as indicated by high peak ALT levels, is an independent factor associated with poor clinical outcome. Peak ALT over 5×ULN increased the risk of mortality by seven-folds. Although age and co-morbid illnesses had been found to have significant negative impact on prognosis in SARS[27,2], in this study, the age of patients did not differ significantly between those with high or low peak ALT levels. Baseline ALT levels have not been found to be associated with any adverse clinical outcome in the past probably because of the delayed elevation of ALT levels in most patients. In fact most immunological damage of SARS in the lungs occurred in the second week of the illness[12,28]. The level of ALT elevation may reflect the severity of acute phase response, which in turn may reflect the severity of tissue damage in SARS[7]. We believe that the elevated ALT can serve as a surrogate marker to predict the clinical outcome of SARS.

We did not find any difference in various adverse clinical outcomes among chronic hepatitis B patients as compared to the HBsAg-negative patients. In our cohort, 16 (53%) chronic hepatitis B patients had elevated ALT levels upon admission and two of them suffered from icteric flare-up of hepatitis. Perhaps those patients who suffered from severe flare-up of hepatitis or decompensated liver cirrhosis might have a higher risk of mortality[29]. Lamivudine was not prescribed to one-third of the patients but none of them died or developed hepatic decompensation.

In conclusion, we found that elevation of transaminase is a very common feature in SARS. The ALT elevation is usually transient and likely to be reactive in nature. Co-infection with HBV in the absence of liver cirrhosis or reactivated hepatitis does not affect the normal course of the disease.

REFERENCES


23 Zhang JZ. Severe acute respiratory syndrome and its lesions in digestive system. World J Gastroenterol 2003; 9: 1135-1138


Science Editor Guo SY  Language Editor Elsevier HK