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# Liver Damage in Pediatric Critically Ill COVID-19 Patients: Brazilian Case-Series

# Werther Brunow De Carvalho<sup>1</sup>, Michele Luglio<sup>2</sup>, Uenis Tannuri<sup>3</sup>, Maria Fernanda Badue Pereira<sup>4</sup>, Isadora Souza Rodrigues<sup>5</sup> and Cintia Johnston<sup>6</sup>

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<sup>1</sup> Faculdade de Medicina da Universidade de São Paulo

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# 8 Abstract

- 9 Coronavirus disease 2019 (COVID-19) Coronavirus disease 2019 (COVID-19) has become an
- <sup>10</sup> important cause critical care admission worldwide. In the context of newly described
- <sup>11</sup> multisystem inflammatory syndrome temporally related to SARS-CoV-2 (PIM-TS), the
- <sup>12</sup> question of liver compromise came into evidence. Our group summarized a case series of 6
- <sup>13</sup> critically ill COVID-19 pediatric patients that presented some degree of liver damage, as
- <sup>14</sup> demonstrated by liver and/or canalicular enzymes elevation, a yet not fully explored
- <sup>15</sup> characteristic of the infection in the pediatric patient, that may indicate a more severe
- <sup>16</sup> progression. Observations regarding the role of systemic inflammatory response can be taken
- <sup>17</sup> from the described cases.
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19 Index terms— hepatic, intensive care, coronavirus.

# 20 1 Introduction

oronavirus disease 2019 (COVID-19) has increasingly become an important cause of critical care admission.
 Some adult studies and case series have focused on the important aspect of extra-pulmonary commitment by

23 Severe Acute Respiratory Syndrome Coronavirus 2 (SARS-CoV-2), with attention to potential liver damage 1.

Between 14 and 53% of a dult patients with COVID-19 showed alanine aminotransferase (ALT) and/or a spartate aminotransferase elevations 1 .

In the context of the recently described multisystem inflammatory syndrome temporally related to SARS-CoV-2 infection (PIM-TS), Whittaker et al 2 compiled data of 58 individuals, showing that ALT median levels on the different phenotypic groups varied from 26 (12-141) to 86 (34-129) U/L.

Our group summarized a series of critically ill COVID-19 pediatric patients with hepatic damage (as demonstrated by liver and/or canalicular enzymes elevation), admitted to a Brazilian tertiary hospital Pediatric Intensive Care Unit (PICU), dedicated to cases of SARS-CoV-2 infection.

# 32 **2** II.

# 33 **3** Methods

From March to June 2020, 35 patients were admitted to Pediatric COVID-19 dedicated wards and PICU in a single tertiary center in São Paulo. Of those patients, 15 needed intensive care support. All the patients had a confirmed diagnosis of SARS-CoV-2 infection performed by nasopharyngeal reverse transcriptase polymerase chain reaction (RT-PCR), serological tests (IgM and IgG) and/or diagnosis of PIM-TS, following the World Health Organization (WHO) criteria 2.

<sup>39</sup> Demographic, clinical and laboratory data were obtained from medical records by two independent investiga-

tors (ML and ISR). After retrospective analysis of 15 critical patients' records, 6 patients without previous hepatic illnesses showed some degree of liver damage and were included in the case series, after thorough discussion among experts and all authors' agreement. As COVID-19 is a new disease, consensus towards the precise definition
of liver damage is still lacking 3. On this case series, the authors defined liver damage by the presence of
new elevations of alanine aminotransferase (ALT), aspartate aminotransferase (AST), gama-glutamyl transferase
(GGT), alkaline phosphatase (ALP) and/or Total Bilirubin in relation to the patient's baseline values (when
available) or the institution's laboratory references, through the hospitalization for critical COVID-19.

Patients consented at the time of hospital admission for the inclusion in a case series, and approval was obtained from the hospital ethics committee for the report of these cases.

# 49 **4** III.

#### 50 5 Results

51 On table 1, main demographic, clinical and liver enzymes characteristics of those included patients are 52 summarized. On Figure 1, the temporal evolution of AST/ALT and GGT levels, during PICU stay, is shown:

#### 53 6 Patient 2:

Admitted to PED with 2 days of weakness and inappetence, followed by cough and progressive respiratory 54 distress. The only gastrointestinal symptoms present during hospitalization were sporadic vomits. The patient 55 showed signs of hypoperfusion and hypotension, with point-of-care ultrasound (POCUS) showing turgid inferior 56 vena cava, right atrium dilation and compromised global systolic function, and Dobutamine infusion was started. 57 SARS-CoV-2 nasopharyngeal PCR was positive, with all other etiologic exams (blood cultures, respiratory virus 58 panel) negative. CRP was elevated on admission (111.0 mg/dL). The patient was transferred to PICU, where 59 evolved with worsening respiratory failure and the need of invasive mechanical ventilation. Echocardiography 60 showed Left Coronary Artery dilation (z-score = +2.7) and left ventricular systolic and diastolic dysfunction 61 (LVEF = 24% with Milrinone and Adrenaline). Due to the diagnosis of incomplete Kawasaki disease, two doses 62 of IVIG (total 2 g/kg) were administered and high dose aspirin was started. PT and aPTT showed elevations, 63 with values of 34.9s and 34.2s (INR = 2.86 and R = 1.19), respectively. After 5 days in critical care, the patient 64 died. 65

#### 66 7 Patient 3:

Admitted to PED with 4 days of progressive respiratory distress with no fever or other symptoms. No gastrointestinal symptoms were present during hospitalization. Patient was intubated and mechanically ventilated due to hypoxic respiratory failure. SARS-CoV-2 nasopharyngeal RT-PCR was positive, with all other etiologic exams (blood cultures) negative. Echocardiography was normal. CRP (48.2 mg/L), Troponin-t (0.059 ng/mL), d-dimer (4,157 ng/mL) and LDH (892 U/L) were elevated at admission. PT (12.3s, INR = 1.01) and aPTT (26.7s, R = 0.98) were normal through PICU stay. The patient showed mild ALT/AST elevations, that resolved after hospital discharge. After 11 days on critical care, the patient was discharged to general pediatric ward.

# 74 8 Patient 4:

After 4 months of hospitalization following chemotherapy complications and acute renal failure, the patient began 75 intermittent fever without other symptoms. Infectious screening exams were performed, including SARS-CoV-2 76 nasopharyngeal RT-PCR that came out positive. Galactomannan test was the only other etiologic test to be 77 78 positive. Echocardiography, thoracic and cranio-facial CT-scans showed no new alterations. Abdominal CT-scan 79 showed liver dimensions on the upper limit with regular shape and contour, without biliary dilation and mild colonic parietal thickening. In spite of the colonic findings, no gastrointestinal symptoms were present. Patient 80 developed severe hypokalemia and worsening renal function, needing PICU transfer. During the course of SARS-81 CoV-2 infection, the patient showed a nearly 6-fold ALT/AST elevation, associated with moderate GGT/ALP 82 elevations (in relation to the previous basal values, shown in table 1) and stable values of PT and aPTT, 14.4s 83 (INR = 1.18) and 29.2s (R = 1.07) respectively. Alterations on liver enzymes resolved after discharge. 84

#### 85 9 Patient 5:

Admitted to PICU after 4 days of odynophagia, fever and headache, evolving with vomits and episodes of 86 convulsion. The diagnosis of refractory status epilepticus was made, and continuous midazolam and pentobarbital 87 were started after orotracheal intubation. Multiple antimicrobial schemes were used through hospitalization 88 89 (Table 1). Cerebral Spinal Fluid (CSF) showed mild alterations (3 cells, with normal glucose and protein levels). 90 Cranial Computed Tomography scan (CT-scan) was normal and thoracic CT-scan showed bilateral ground-glass 91 opacifications. SARS-CoV-2 PCR was negative, as were all other etiologic tests (CSF culture and viral PCRs, 92 blood cultures, respiratory virus panel). Echocardiography showed right (z-score: +3.0) and anterior descendent (z-score: +2.6) coronary arteries dilation. CRP (318.1 mg/L) and Ddimer (19,514 ng/mL) were elevated at 93 admission and during intensive care. IVIG and high-dose methylprednisolone pulse-therapy were administered, 94 due to the diagnosis of PIM-TS. The patient presented episodes of melena and was submitted to endoscopic 95 evaluation that showed two ulcerations on duodenal superior wall. PT also showed mild elevations, with a peak 96 of 15. 97

#### 98 10 Patient 6:

Patient on the seventh day after chemotherapy (vincristine and doxorubicin) was admitted to the PED with 2 99 days of fever, cough and progressive respiratory distress. The patient was diagnosed with neutropenia and sepsis, 100 starting empiric antimicrobial therapy, with association of Amphotericin B through the course of hospitalization. 101 SARS-CoV-2 nasopharyngeal PCR was positive, with all other etiologic exams (blood cultures, respiratory virus 102 panel) negative. The patient evolved with respiratory failure and hemodynamic instability, needing invasive 103 ventilatory and inotropic support on the second day of hospital admission. No gastrointestinal symptoms 104 appeared during PICU stay. D-dimer (1,932 ng/mL) and ferritin (3,295 ng/mL) showed elevations through 105 hospitalization, with normal PT (13s, INR = 1.0) and aPTT (30s, R = 1.11). Echocardiography showed small 106 pericardial effusion and thoracic CT-scan showed diffuse bilateral ground-glass opacifications. The patients 107 108 deceased after 5 days in ICU due to refractory shock.

#### 109 **11 IV.**

#### 110 **12** Discussion

III Important variations are found when evaluating ALT/AST levels in patients with COVID-19 3,4. This case series corroborates previous findings, with AST elevations ranging from 65 to 5908 U/L.

Transaminase elevations seen in this series may be related to four mechanisms: (1) Drug induced liver injury (DILI); (2) Direct biliary injury by coronavirus; (3) inflammatory response in the context of cytokine storm; (4) Ischemia/Reperfusion and microthrombosis 7.

Abnormalities on liver enzymes seen can occur on either the initial viremia phase or during the subsequent inflammatory phase 7. It was already reported that high ALT and bilirubin can be considered biomarkers of a more severe clinical course 7,8.

The potential for DILI in the context of critical COVID-19 cannot be neglected 9. All patients included in our case series received at least one Category A or B hepatotoxic drug, as described by LiverTox® 10. Drug induced liver damage may be an important contributing factor to a multifactorial condition.

Different from previous reports 4,5, patients included showed moderate elevations on GGT levels, consistent with experimental observations that cholangiocytes express ACE-2 receptors, a target for direct viral invasion and damage 6.

Three patients had features consistent with PIM-TS (patients 1, 2 and 5) 2, as defined by the WHO criteria. SARS-CoV-2 can be considered the trigger of an uncontrolled systemic inflammatory condition or cytokine "storm". In this context, cellular apoptosis and necrosis and the release of damage-related patterns may induce injuries to multiple organs, the liver included. Hepatic endothelial involvement in the inflammatory process, with consequent neutrophil extracellular traps (NETS) stimulation and microthrombi formation, in a process similar to the one happening in the lungs 11, needs to be further studied.

Effenbergeret al 12 explored the connection between systemic inflammation and liver injury in COVID-19 hospitalized patients. IL-6 and CRP levels positively correlated with AST elevations (respectively, r 2 = 0.481, p<0.001 and r 2 = 0.38, p<0.001) in all 96 included patients with pronounced effects on critically ill patients. Those findings correlate with this case series, as high levels of CRP and PCT were seen in patients with liver enzyme elevations. This cytokine "storm" may play a vital role in the hepatic damage.

In the beginning of the pandemic, the main focus of intensivists was on the viral potential to induce hypoxia. 136 Hypoxia-reperfusion injury to the liver can stimulate hepatocyte cell death and inflammation, marked by oxygen 137 reactive species accumulation 13, another potential causative mechanism to liver damage. PICU mortality 138 among the described patients was 50% (3/6) with a length of stay of 12.5 (6.5-20) days, while the remainder of 139 the pediatric COVID-19 critically ill patients experienced a mortality rate of 33.3% (3/9) and length of stay of 7 140 (3-10) days. Presence of liver enzyme alterations indicates a more severe disease course, with all patients but one 141 (patient 4) needing ventilatory, hemodynamic support or both. Given the tertiary condition of our center, the 142 population included is mainly composed of patients with chronic conditions, what have impacts on the outcomes 143 144 seen. In regard of the liver enzyme elevations, special care was taken to compare previous individual baseline levels to the highest values seen towards disease course. 145

This study has limitations of a small case series, which needs confirmation on larger groups. Due to the retrospective nature of the study and to conditions inherent of a pandemic in a developing country, a complete evaluation of radiological and histological aspects of the hepatic compromise may be lacking. As the focus was on the clinical description of patients with liver abnormalities, comparison with the global data of all COVID-19 patients admitted to the hospital was not made and can be an important future step.

Reports from over the world 13 show slightly different outcomes and evolutions of clinical conditions associated with COVID-19 in children. In a recent report by Sadiq et al 14, Pakistani children with PIM-TS showed an incidence of coronary artery aneurism (62.5%), higher than European and North American numbers (9-36%). Some of the findings in our case series can be justified by regional differences, that may be better identified in future studies. Knowledge of those disparities are relevant to deepen the understanding of the clinical potential of SARS-CoV-2 infection.

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# 12 DISCUSSION

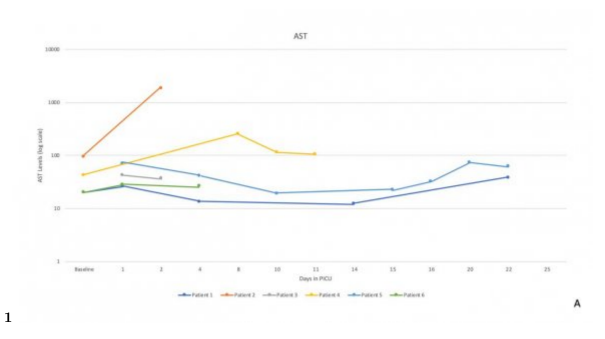


Figure 1: Figure 1 :

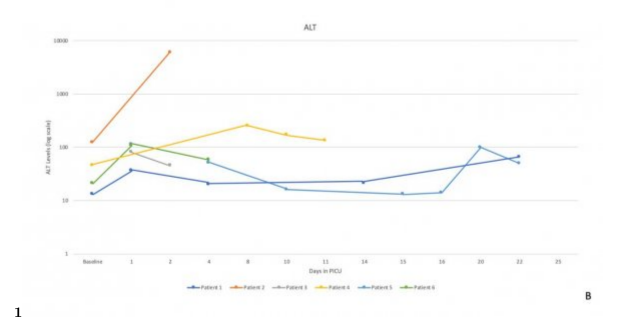


Figure 2: Figure 1 :

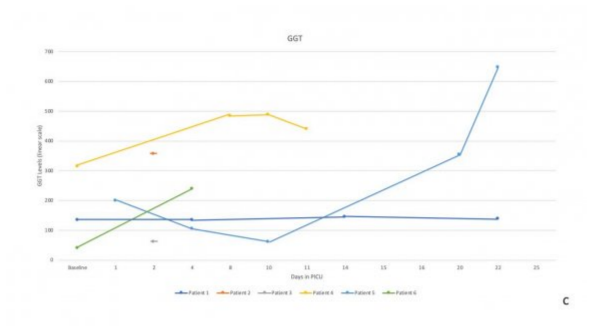


Figure 3:

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Figure 4: Table 1 :

#### 12 DISCUSSION

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