Elucidating the pivotal role of convalescent plasma therapy in critically ill COVID-19 patients: A review

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Abstract

World Health Organization (WHO) declared coronavirus disease (COVID-19) a pandemic in March 2020. Currently almost every country in the world has reported cases with high mortality rates. The European Union (EU), the United States of America (USA) and the United Kingdom (UK) are the severely affected countries. Nevertheless, the WHO is very much concerned about countries with weak health systems. The clinical characteristics of COVID-19 varies extensively, ranging from asymptomatic infections to severe as well as critical pneumonia with high mortality rates in the elderly and patients with co-morbid medical illness. Convalescent Plasma Therapy (CPT) has been successfully used in treating various viral disease outbreaks such as 1918 influenza pneumonia pandemic, poliomyelitis, measles, mumps, Machupo virus, Junin virus, Lassa virus, Ebola etc. High-titer specific antibodies maybe capable of binding to Coronavirus-19 (CoV-19) and neutralize the viral particles, inhibit entry to uninfected cells, and trigger potent effector mechanisms such as complement activation as well as phagocytosis. Therefore, in most countries with very weak health systems with no Intensive Care Units (ICUs) or trained ICU physicians, earlier diagnosis/CTT for severely COVID-19 patients may be rewarding. Therefore, solidarity control trials on CPT for COVID-19 patients involving large number of patients are urgently needed.

Introduction

In December 2019 a novel strain of coronavirus, officially named Coronavirus-19 (CoV-19) was first isolated from three patients with coronavirus disease and communicated to the Chinese Center for Disease Control and Prevention.1,2 COVID-19 was linked to a cluster of acute respiratory illness cases from Wuhan, China.3 The World Health Organization (WHO) declared COVID-19 a pandemic in March 2020.4 Currently almost every country in the world has reported cases with high to moderate mortality rates. The European Union (EU), the United States of America (USA) and the United Kingdom (UK) are the severely affected countries. Nevertheless, WHO is very much concerned about countries with weak health systems.

The clinical presentation of COVID-19 varies extensively, ranging from asymptomatic infections to severe, including critical pneumonia with extreme mortality rates.5 Currently, there are no established antiviral medication for the treatment of COVID-19. Also, no vaccines are available for the prevention of COVID-19. Recently, the USA Food and Drug Administration (FDA) has approved the use of Convalescent Plasma Therapy (CPT) for the treatment of critically ill COVID-19 patients on conditions that doctors get approval over the telephone before initiating treatment.5 Nevertheless, the disease continues to kill many patients even in countries with the most robust health systems, because most of these countries overlook the use of CPT for severe COVID-19 patients at the Intensive Care Units (ICUs). In countries with very weak health systems with no ICUs or trained ICU physicians, early initiation of CPT for high risk COVID-19 patients maybe rewarding as well cost reduction.

Overview of convalescent plasma therapy for viral disease outbreaks

The use of CPT or immunoglobulins for viral infectious disease date to the 1918 Influenza Pneumonia Pandemic (IPP) (Figure 1).5,6 Luke et al. with in a meta-analysis involving eight studies with a total number of 1703 influenza pneumonia patients from 1918-1925 who received an infusion of influenza-convalescent human blood products revealed a pooled complete decline of 21% (95% CL 15–27; p<0.001) in the overall crude case mortality rate at low risk of bias.5 Also, CPT was used in the past to treat diseases such as poliomyelitis, measles, as well as mumps (Figure 1) during outbreaks similar to COVID-19.5 Furthermore, CPT was used to successfully treat patients with Machupo virus infections, which caused the Bolivian Hemorrhagic Fever (BHF) in Bolivia (Figure 1).7 Moreover, during Junin virus infection in Argentina, CPT was used to efficiently treat patients with the Argentinian Hemorrhagic Fever (AHF) (Figure 1).8,9 Similarly, in Nigeria, CPT was successfully used to treat Lassa Fever (LF) during the Lassa virus outbreak (Figure 1).10 It is wealth noting that almost these outbreaks occurred in countries with very weak health system at time of the disease outbreaks.

In 2003, during the peak of Severe Acute Respiratory Syndrome-coronavirus 1 (SARS-CoV-1) outbreak in China, CPT was an effective treatment option in severe cases (Figure 1).11–13 This treatment modality was the last alternative for the survival rate of patients with SARS-CoV-1 whose illness continuous to worsen notwithstanding treatment with pulsed methylprednisolone.14 Cheng et al. in 2005 demonstrated that in their study involving 80 SARS-CoV-1 patients that CPT correlated with better rate of hospital discharge approximately 3 weeks after the onset symptom compared with patients who did not receive CPT.15 The long hospital stay could be due to the delay in initiation of treatment because patient where put on conventional therapies before the initiation of CPT. Nevertheless, several studies exhibited a shorter hospital stay as well as lower death rate in patients treated with CPT than patients who were
not on CPT.11,14

Hung et al., in a multicenter, prospective, double-blind, randomized controlled trial involving 20 patients in 2009, demonstrated that convalescent plasma obtained from patients who recovered from influenza A H1N1 pdm09 virus infection was an effective and efficient treatment modality for severe influenza A (H1N1) infection (Figure 1).16 They indicated that CPT correlated well with a lower viral load, serum cytokine response as well as a decreased death rate within 5 days of symptom onset.16 In 2014, WHO approved CPT as one the critical treatment option for patients with Ebola. The use of convalescent plasma obtained from patients who had recovered from Ebola virus disease was efficient and effective practical treatment during these outbreaks (Figure 1).17

In 2015, during the Middle East Respiratory Syndrome Coronavirus (MERS-CoV) outbreak, CPT was effective and efficient in treating severe patients (Figure 1).18 Based on the success story of CPT, a protocol for its use was established.18 Nevertheless, a practicality interventional study involving CPT during MERS-CoV infection treatment failed to detect acceptable high-titer plasma from patients with confirmed or suspected MERS, their close family members, or healthcare workers exposed to MERS.19

Mair-Jenkins et al. demonstrated in a meta-analysis that the viral load and death rate was low in patients with severe SARS-CoV and MERS-CoV infections who received various doses of convalescent plasma.20 The study further revealed that, CPT was associated with no adverse events or complications.20 It is very highly anticipated that, CPT may be a game changer in the fight against COVID-19 most especially in countries with very weak health systems without ICUs or ICU physicians. Thus, control trials are urgently needed to establish the potency of CPT for COVID-19.

Clinical characteristics and potential treatment regimes

The clinical characteristics of COVID-19 varies extensively, ranging from asymptomatic infections to severe, including critical pneumonia with high mortality rates in the elderly and patients with co-morbid medical illness.3 Chen et al. observed symptoms such as fever, cough, fatigue, anorexia, myalgia, and diarrhea.2 They indicated that these symptomatology’s did not differ significantly between dead patients and healed patients. Nevertheless, they observed that dyspnea, chest tightness, as well as disorders of consciousness were more frequent the deceased.3 The above symptoms could be indicative for the use of CPT in countries without ICUs. Huang et al. indicated that, limited number of patients with COVID-19 had noticeable upper respiratory tract signs and symptoms such as rhinorrhea, sneezing, or sore throat.21 They concluded that, the target cells might be located in the lower airway since majority of their patients presented with low respiratory signs and symptoms.21

A body of evidence indicates that most dead patients had tachycardia and/or tachypnoea with lower pulse oxygen saturation (<93%).3 The study further indicated that, in most of the deceased patients, chronic hypertension as well as other cardiovascular comorbidities accounted for mortality compared to recovered patients.3 Furthermore, hypoxemia related symptoms were more frequent in patients who eventually died as compared with recovered patients. Patient who eventually died usually developed systematic inflammation as well as multi-organ failure as compared to recovered patients.3 The above symptomatology point to the fact that patients who died from COVID-19 often aggravates into severe or critical condition on admission before dying.3 It is worth noting that the onset of certain symptoms may aid physicians detect patients at risk of a poor outcome and hence early initial of CPT.3

Huang et al. indicated that most patients presented with bilateral ground-glass opacities on chest CT scans.21 Countries with weak health systems do not have CT scans for the evaluation of patients with COVID-19. These countries only have X-Rays for the diagnosis of respiratory diseases. Thus, urgent X-Ray characteristics features of COVID-19 are needed to aid in the diagnosis. These clinical parameters maybe helpful to frontlines care gives for COVID-19 patients in arriving at optimum decision for the use of CPT for patients in countries with weak health systems. The current treatment of COVID-19 involves >85% of anti-viral agents, such as oral oseltamivir 75 mg every 12h, intravenous ganciclovir 0.25g every 12h, as well as oral lopinavir/ritonavir tablets 400/100mg twice daily.19,22 Also, Remdesivir is another promising anti-viral medication under clinical trials.19,22 An old
anti-malarial, chloroquine phosphate or hydroxychloroquine is a derivative of chloroquine, and is effective in inhibiting COVID-19 cellular invasion and thus inhibits exacerbation of pneumonia as well as lung inflammation. Hydroxychloroquine has been approved by the USA FDA for the treatment of COVID-19. 

Based on the mechanism of action of hydroxychloroquine, we stipulate it will be best preventive medication as a post-exposure prophylaxis as well as treatment for asymptomatic carriers who are likely to spread the virus widely. Clinical trials are urgently needed in this direction. It has been stipulated that 80%-90% of medication used in treating COVID 19 in China were Traditional Chinese Medicine (TCM). Glycyrhizae Radix Et Rhizoma (Gancao), Astragali Radix (Huangqi), Atractylodis Macrocephalae Rhizoma (Baizhu), Saposhnikoviae Radix (Fangfeng), as well as Lonicerae Japonicae Flo were the five most commonly used herbs. 

**Intensive care management**

The elderly patients (>60 years), male sex, as well as comorbidities like hypertension, diabetes and immunosuppression are linked to higher morbidity and mortality in COVID-19 infection. These categories of patients are candidates for ICU as well as CPT. Murthy et al. indicated that ICU is very likely to have a substantial influence on patient outcomes. A body of evidence indicates that the death rate appears to be around 2%, and about 25.9% of the patients with COVID-19 pneumonia required ICU admission, mostly because 20.1% often develop Acute Respiratory Distress Syndrome (ARDS). 

There is currently no vaccine or approved anti-viral regime for the treatment of critically ill patients. ICU regimes essentially emphasize on supportive care such as oxygenation, ventilation, as well as fluid management. Severe COVID-19 patients may require extracorporeal membrane oxygenation which is usually managed by well-trained ICU doctors. Low income countries do not have these kinds of doctors. Studies have demonstrated that a combination of low-dose systematic corticosteroids, anti-virals and atomized inhalation of interferon have been incorporated in the regime of COVID-19 patients. Nevertheless, control trails on the usage of systematic corticosteroids anti-virals and atomized inhalation of interferon for treatment of COVID-19 is urgently needed.

**Criteria for convalescent plasma therapy and Mode of administration**

We advocate that CPT may be beneficial for critical ill COVID-19 patients in countries without ICUs or ICU physicians. In the current COVID-19 outbreak, severe disease is defined as dyspnea, respiratory frequency >30 breaths per minute, blood oxygen saturation <93%, ratio of arterial partial pressure of oxygen to fraction of inspired oxygen (PaO2/FiO2) <300, or lung infiltrates >50% within 24 to 48 hours. Life threatening disease is defined as respiratory failure, septic shock, or multiple organ dysfunction or failure. Most of these parameters are measured by the ventilator which are not forecoming in countries with weak health systems. Therefore, clinical parameter will be the critical determiner of CPT regime.

Eligible plasma donors are patients who have recovered from COVID-19 infection. Appropriate protocols such as written informed consent ought to be obtained from eligible patients. It is advocated that donors should also have tested negative for COVID-19 and other potential viruses such as hepatitis B virus, hepatitis C virus, Human Immune Deficiency Virus (HIV), as well as bacteria like syphilis at the time of blood donation. Also, laboratory investigation such as white blood cell count, lymphocyte count, chemistry parameter for the dictation of liver and kidney function, Cycle Threshold value (Ct) as well as inflammatory factors like C-Reactive Protein (CRP), procalcitonin, and IL-6, and serum antibody titer for IgG, IgM, and neutralizing antibodies need to be assessed. In most studies as well as protocols in various viral outbreaks, patients often received a single transfusion of convalescent plasma as the desired therapy. 

The preparation of CPT most strictly meets the WHO criteria for blood and blood products. ABO blood types of the critical ill COVID-19 patients must strictly be compatible with the convalescent plasma donor. Shen et al. advocates that the patients should be transfused 2 consecutive times with 200-250 mL of ABO-compatible convalescent plasma. In all a maximum of 400 mL of convalescent plasma ought to be transfused on the same day it was obtained from the donor. We advocate that all other plasma transfusion protocol should be strictly followed to avoid advice transfusion reactions.

**Antibodies detection in COVID-19 patients**

Long et al. demonstrated acute antibody responses to CoV-19 in 285 patients with COVID-19 using a Magnetic Chemiluminescence Enzyme Immunoassay (MCLIA) for virus-specific antibody detection. They indicated that, the percentage of patients with positive CoV-19-specific IgG was 100% from day 17-19 after the onset of symptom while the percentage of patients with positive CoV-19-specific IgM plateaued at 94.1% on day 20-22 after the onset of symptoms. They further indicated that, IgG as well as IgM titters were more elevated in severe group than the mild group. Nevertheless, they observed substantial difference in IgG titer 2-week after the onset of symptoms. Their study failed to establish how long the IgG and IgM titters last after the patient with COVID-19 has fully recovered. Their study was unable test samples for virus neutralization and thus the neutralizing actions of the detected IgG antibodies are unknown. Furthermore, they were unable to determine the association between antibody response as well as clinical outcomes and thereafter. 

Long et al. further indicated that serum from patients with COVID-19 exhibited no cross-binding to the S1 subunit of the CoV-19 spike antigen. They also did not observe cross-reactivity of serum samples from patients with COVID-19 to nucleocapsid antigens of CoV-19. Long et al. study however failed to explicitly explain the critical role of IgG as well as IgM in the patients with COVID-19 other than diagnosis of the infection. Their study did not explore the role of these antibodies in convalescent therapy. Thus, similar studies are warrant to affirm their finding on CoV-19 spike proteins and IgG and IgM in the patients with COVID-19. Studies have shown that, the function and viral structure of CoV-19 are modulated by the Envelope protein (E), Membrane protein (M), the Spike protein (S) and the Nucleocapsid protein (N). The N protein aid in the development of the viral capsid as well as arrangement of whole viral structure while the S protein aid in viral attachment to the host cell via the ACE2 and TMPRSS2. We are not convinced that antibodies generated by CoV-19 did not cross-bind with its binding receptors or proteins just like other viral infections.

Okba et al. outlined the elaboration of serologic assays for identification of virus neutralizing antibodies and antibodies to the N protein as well as various S protein domains, such as the S1 subunit, and the
Receptor-Binding Domain (RBD) of CoV-19 in an ELISA design. They assessed CoV-19-specific antibody reactions in severe and mild patients by using serum samples obtained at different times postonset of illness from 3 PCR-established COVID-19 patients from France. They further observed that, amongst the S antigens investigated, S1 was more specific than S2 in detecting CoV-19 antibodies. The also observed a high degree of conservation in the coronavirus S2 subunit in comparison to S1. The detected seroconversion of IgG during a second week of COVID-19. In their three in-house ELISAs investigation, the RBD as well as N protein ELISAs were more sensitive than S1 ELISA in identifying antibodies in mildly infected patients and demonstrated greater associations with PRNT50 titers.

A preprint published by Wu et al. using a pseudotyped-lentiviral-vector-based neutralization assay to measure Cov-19-specific NABS in plasma obtained from 175 recovered COVID-19 patients with mild symptoms showed higher titers of CoV-19 Receptor Binding Domain (RBD), S1, and S2-specific binding antibodies. Their study revealed that, the titers of NABS plateaued from day 10-15 after the onset COVID-19 and remained stable thereafter in patients. They also observed that, the NABS titers in patients correlated with their ages. Elderly patients had expressively higher titers of NABS than younger patients. They concluded that, antibodies targeting on distinctive domains of S protein, such as S1, RBD, and S2, may all contribute to the neutralization. Their study was limited because, the kinetics of viral loads were not measured and patients in severe as well as critical condition were excluded from their study because they received passive antibody treatment before sample collection.

Zhao et al. explored the dynamics of total Antibody (Ab), IgM as well as IgG antibody against Kindly fuse this line as one. CoV-19 in serial blood samples obtained from 173 patients with established COVID-19. They used enzyme linked immunosorbent assay (ELISA) to evaluate the antibodies. The ELISA for absolute antibodies recognition was developed based on double-antigens sandwich immunoassay (Ab-ELISA), utilizing mammalian cell secreted recombinant antigens possessing the Receptor Binding Domain (RBD) of the spike protein of CoV-19 as the immobilized as well as HRP-conjugated antigen. They detected that, the seroconversion rate of Ab, IgM and IgG were 93.1% (161/173), 82.7% (143/173) and 64.7% (112/173), respectively. They observed that the cumulative seroconversion rate for Ab and IgM plateaued at 100% approximately 1-month after onset of the COVID-19. They also observe a strong positive correlation between clinical severity and Ab titer since 2-week after illness onset, for the first time in COVID-19 patients.

Several studies have demonstrated that, in CoVs, the N protein has high immunogenic activity and is copiously secreted during infection. Grifoni et al. identified five domains on the S glycoprotein of CoV-19 predicted to be linked with a robust immune activity. Liu et al. demonstrated recombinant S-based ELISA was more sensitive than the recombinant N-based one in the detection of IgM antibodies when they performed a recombinant N-based as well as recombinant S-based ELISAs to detect IgM and IgG antibodies in human serum against CoV-19. This mean that convalescent plasm containing IgM and IgG are likely to bind to both S-spike protein and the N protein leading to lessen of viral load and disease burden (Figure 2).

**Convalvescent plasma therapy for COVID-19**

A body of evidence indicates that the efficacy and efficiency of CPT is founded on the idea that antibodies from convalescent plasma might suppress viremia. It is demonstrated that the donor plasma has perceptible IgG as well as IgM anti-COVID-19 antibodies and neutralized the virus in *in vitro* cultures (Figure 2). Zhen et al. in study involving 21 recovered COVID-19 patients demonstrated that, CPT can discontinue CoV-19 shedding but cannot reduce mortality in critically end-stage COVID-19 patients, therefore, CPT ought to initiated earlier in the course of the disease to achieve favorable outcomes. The key mechanism via which these immunoglobulins bind to COVID-19 antibodies to eliciting the desired effect still rigorous investigations. It was observed that, the patients typically develop a primary immune response by days 10-14, subsequent to virus clearance.

**Figure 2.** An illustration showing the mechanism via which the IgM and IgG from Convalescent plasma binds to coronavirus-19 (CoV-19) resulting in none binding of CoV-19 receptors as well as decrease coronavirus disease-19 (COVID-19).
Poh et al. identification of two (S14 and S21) immunodominant linear B-cell epitopes on the S glycoprotein of CoV-19 that are critical in controlling infection (Figure 2). They screened 25 convalescence serum samples at 1:1000 dilution for neutralizing antibodies against a pseudo-typed lentivirus secreting CoV-19 S glycoprotein tagged with a luciferase reporter during current COVID-19 pandemic in Singapore. Six convalescents out of the 25 patients exhibited a good neutralizing response and were used for functional characterization above. Furthermore, research should also focus on how long immunity last during the convalescent period. In countries with weak health systems, it should be more effective to administer the convalescent plasma at the early stage of disease.

Conclusions

High-titer specific antibodies maybe capable of binding to COVID-19 and neutralize the viral particles, inhibit entry to uninfected cells, and trigger potent effector mechanisms such as complement activation as well as phagocytosis. The disease continues to kill many patients, even in country with most robust health system, because most of these country outlook CPT for severe COVID-19 at the ICUs. Therefore, early initiation of CPT for severely COVID-19 patients in countries with very weak health systems, with no ICUs or trained ICU physicians, early initiation of CPT for severely COVID-19 may be rewarding. Therefore, solidarity control trials on plasma therapy for COVID-19 involving large number of patients are urgently needed.

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