

CONVALESCENT PLASMA THERAPY IN COVID-19 PATIENTS WITH ACUTE RESPIRATORY DISTRESS SYNDROME (ARDS)

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Abstract

COVID-19 is caused by SARS-CoV-1, an RNA virus of the betacoronavirus genus and making it the seventh coronavirus that infect humans. Because particular therapies are still in the research stage, there is currently no confirmed treatment for this illness that has been agreed by the World Health Organization (WHO) or other clinical institutes. The reason for this is because there are many different potential remedies. Antiviral treatments like pavirovir, oseltamivir, and remdesivir have been investigated and tested. On the other hand, the outcomes of the replies of patients who were given these medications are still quite inconsistent. Furthermore, the mortality rate accompanying with COVID-19 has remained at a level of less than 5.21 percent of all cases that have been documented. Patients suffering with COVID-19 may be treated with convalescent plasma, which is a therapeutic option that utilizes a mix of neutralizing antibodies and other immunological component. Activation of body-dependent cellular cytotoxicity (ADCC) and phagocytic activity against COVID-19 will occur as a result of this immunological component. This medication also has the potential to reduce the systemic inflammatory response brought on by COVID-19. Clinical improvement was different after 28 days when convalescent plasma was used as a treatment for patients with severe COVID-19 symptoms and emergency conditions compared to patients treated with conventional therapy alone. however it is not very significant.

Keywords: COVID-19, ARDS, Convalescence Plasma Therapy

TERAPI PLASMA KONVALESEN PADA PASIEN ACUTE RESPIRATORY DISTRESS SYNDROME ARDS COVID-19

Abstrak

Sindrom pernafasan akut yang parah koronavirus 2 (SARS-CoV-1), yang bertanggung jawab untuk COVID-19, adalah virus RNA yang termasuk dalam genus betacoronavirus dan merupakan salah satu spesiesnya. Virus khusus ini adalah koronavirus, menjadikannya yang ketujuh yang diketahui menginfeksi manusia. Karena terapi tertentu masih dalam tahap penelitian, Organisasi Kesehatan Dunia (WHO) dan lembaga klinis lainnya belum mengembangkan terapi yang pasti untuk mengatasi kondisi ini. Alasan untuk ini adalah karena ada banyak solusi potensial yang berbeda. Perawatan antivirus seperti pavirovir, oseltamivir, dan remdesivir telah diselidiki dan diuji. Di sisi lain, hasil jawaban pasien yang diberi obat tersebut masih cukup tidak konsisten. Selain itu, angka kematian terkait COVID-19 tetap berada pada level kurang dari 5,21 persen dari semua kasus yang telah didokumentasikan. Pasien yang menderita COVID-19 dapat diobati dengan plasma pemulihan, yang merupakan pilihan terapeutik yang menggunakan campuran antibodi penawar dan komponen imunologi lainnya. Aktivasi sitotoksitas seluler yang bergantung pada tubuh (ADCC) dan aktivitas fagositik terhadap COVID-19 akan terjadi sebagai akibat dari komponen imunologi ini. Obat ini juga berpotensi mengurangi respons peradangan sistemik yang ditimbulkan oleh COVID-19. Penggunaan plasma konvalence sebagai pengobatan pasien bergejala COVID-19 yang parah dan keadaan darurat menunjukkan perbedaan perbaikan klinis setelah 28 hari jika dibandingkan dengan terapi konvensional saja. meskipun kenyataannya tidak substansial.

Kata kunci: COVID-19, ARDS, Terapi Plasma Konvalence

INTRODUCTION

In 2019, the Wuhan Province in China was the location where the sickness known as COVID-19 (coronavirus disease 2019) was first detected. It soon spread around the planet, and as of the 20th of April, 2020, 2.4 million individuals all over the world had been impacted by it. As a direct result of this, the WHO classified it as a pandemic.¹ COVID-19 was infected with SARS-CoV-2, which stands for severe acute respiratory syndrome coronavirus-2. This virus is a species that is a part of the betacoronavirus genus that causes severe acute respiratory syndrome. This virus is the 7th specimen of a coronavirus that has been identified as being able to infect humans.²

There is not yet a clear therapy for the illness, and neither the WHO nor any other clinical institutions have developed one. The research phase of the illness is still under progress. This is due to the fact that the disease is still in its early stages.³ Antiviral medications such as favipiravir, oseltamivir, and remdesivir have been used in conjunction with one another to treat this condition. Despite this, patients' responses to the medication may run the gamut from positive to negative and anywhere in between. In addition, COVID-19 is still responsible with the deaths of 5.21 percent of all patients.⁴

The use of convalescent plasma (CP) as an antiviral model is one of the neutralizing antibody-containing antiviral models. The therapeutic effects of COVID-19 are thought to arise from three distinct immunological pathways: phagocytosis, complement activation, and antibody-dependent cellular cytotoxicity (ADCC). It is believed that the therapeutic action against COVID-19 is mediated via these immunological mechanisms. Defensins, pentraxins, and anti-inflammatory cytokines are thought to reduce severe inflammatory response syndrome (SIRS). SIRS is the underlying pathophysiology of acute respiratory distress syndrome (ARDS) and death.⁵

The CP treatment was revealed to be effective in treating a variety of viral conditions, for instance Middle East respiratory syndrome (MERS) and Ebola infection, according to the findings of a number of studies.⁶

In treating COVID-19, CP is not only useful when used in conjunction with other therapies. The reduction of viral load, improvement in cytokine response, and reduction in mortality are the goals of convalescent plasma treatment. In addition to this, the CP treatment transfers antibodies from patients who have successfully in good health caused by a specific agent to patients who are also afflicted by that agent. These forms of passive immunity may provide patients with assistance in combating their

sickness and slowing the rate at which it is progressing.⁷

Because it includes antibodies with certain receptor domains, CP attained from COVID-19 patients who have recovered is thought, according to the findings of some research, to have the potential to possess antiviral characteristics.⁸ When implementing CP, some elements must be taken into consideration despite the fact that their exact nature is still unclear. Considerations include the severity of patients who would benefit from CP, the availability of plasma donors, the timing of CP's delivery, and the severity of patients who would not benefit from CP.

The treatment of CP was reported to generate good consequences in a limited sample of patients suffering with severe COVID-19 symptoms, according to the research conducted by Shen et al. (2020).⁹ In order to determine whether or not this treatment was effective, which was a task that had become extremely challenging in pandemic conditions before the introduction of the vaccine, the research needed to involve a greater number of patients and should have been designed in a more effective manner.¹⁰ This article discusses the studies conducted on the use of CP therapy for ARDS patients carrying the COVID-19 virus.

ACUTE RESPIRATORY DISTRESS SYNDROME (ARDS)

The early onset of pulmonary oedema, bilateral pulmonary infiltration, and impaired respiratory system compliance are the characteristics of ARDS, which is not caused by a cardiac etiology. ARDS is an abbreviation for acute respiratory distress syndrome.¹¹ As this description describes it, ARDS is a severe kind of diffuse lung damage that occurs when all of the following criteria are satisfied:¹²

- (1) A new clinical cause or a worsening of respiratory symptoms within a week of the onset;
- (2) A bilateral X-ray opacity in the chest that has nothing to do with fluid accumulation, lung collapse, or nodules;
- (3) Recognizing breathing failure caused by fluid overload or cardiovascular collapse; and
- (4) Hypoxemia is defined as having a PaO₂/FiO₂ ratio that falls into one of these three categories:
 - a. low (200 < PaO₂/FiO₂ < 300 mmHg).
 - b. moderate (100 < PaO₂/FiO₂ ≤ 200 mmHg).
 - c. severe (PaO₂/FiO₂ ≤ 100 mmHg).

A feature of the heterogeneous illness known as ARDS is an intensification in the pulmonary capillary endothelial cell permeability.¹³

ARDS used to be commonly known as non-cardiogenic pulmonary edema. ARDS causes the alveoli to become filled with fluid exudate, increasing

the permeability of the alveolar-capillary barrier so that fluid containing protein can enter the alveoli. This is in contrast to congestive heart failure, which results in pulmonary edema as a result of elevated left heart pressure-related hydrostatic pressure. In congestive heart failure, this leads to pulmonary edema. The presence of fluid in the alveoli may lead to hypoxemia, shunting from right to left, and a reduction in respiratory compliance. Although arterial PCO₂ levels are normally within normal limits, there is an increase in ventilation dead space, which is reflected in increased minute ventilation. A common complication of ARDS is pulmonary hypertension, which may be triggered by a variety of factors, including the accumulation of fibrin inside blood vessels and the narrowing of blood vessels in response to low oxygen levels. This condition may be addressed by using techniques such as positive pressure ventilation and vessel compression.¹⁴

The pathological phases of ARDS are often described using a standard format that includes three stages that follow one another and overlap. During the initial stage of lung injury, known as the exudative phase, the pathological abnormalities that were seen were referred to as diffuse alveolar destruction. The alveolar gap is filled with an edematous fluid that contains protein, and the alveolar walls themselves are lined by hyaline membranes. In addition, the epithelium is disturbed, and neutrophils enter the interstitial space and alveoli, which results in the accumulation of macrophages and, in some cases, bleeding. This stage, which lasts for between five and seven days, is followed, in some individuals, by the proliferative phase. Fibrosis and the organization of the hyaline membrane have both occurred at this point in the process. Decreased neutrophil count and the severity of pulmonary edema are diagnostic of the proliferative phase, which is marked by pulmonary capillary occlusion, interstitial collagen buildup, and alveolar collagen deposition. Proliferative stage is characterized by pulmonary capillary obliteration, interstitial and alveolar collagen accumulation. The fibrotic phase may be seen on radiographs of patients with chronic ARDS (ARDS that has lasted for more than two weeks) after this phase has passed.¹⁴

At first, it was thought that either direct or indirect lung damage caused an excess production of inflammatory mediators in the pulmonary microcirculation. As inflammatory mediators like neutrophils activate and migrate across the surfaces of the alveolar epithelium and vascular endothelium, they produce proteases, cytokines, and reactive oxygen species (ROS). Pathological vascular permeability, a breach in the alveolar epithelial cell barrier, and necrosis of type I and type II alveolar cells are the results of the migration and release of these mediators. The result is a buildup of fluid in the lungs, known as pulmonary edema. The reduction in

lung compliance and the increase in gas exchange difficulty caused by hyaline membrane development and surfactant depletion. Collagen deposition, fibrosis, and the progression of illness are also the outcomes of fibroblast infiltration. During the recovery phase, a number of processes occur simultaneously. Activated neutrophils are slowed down by anti-inflammatory cytokines. Proliferation and differentiation of type II alveolar cells into type I alveolar cells strengthens the epithelial lining of the alveoli, allowing for the drainage of fluid from the alveoli into the microcirculation and pulmonary lymphatic system through an osmotic gradient. Alveolar cells and macrophages collaborate throughout the healing process to clear the alveoli of protein debris.¹⁴

DEFINITION THERAPY OF CONVALESCENT PLASMA

Patients who have overcome an illness and gained humoral immunity are used in convalescent plasma treatment. Humoral immunity is a kind of tolerance to the bacteria that cause sickness. In most cases, donors' plasma is obtained after they have completely recovered, making it suitable for use in the convalescent period. Water, proteins, and inorganic salts make up the bulk of convalescent plasma. Antibodies and immunoglobulins directed against an infectious pathogen may inhibit viral replication and lower viremia in those who are already infected. It is possible that these antibodies may kill viruses by doing two things: blocking the attachment of viruses to endosomes and halting the discharge of virions from infected cells. Third, preventing viral protein cleavage through extracellular proteolysis, and fourth, preventing viral protein entrance into human cells.¹⁵

SEVERE CATEGORY CLINICAL SYMPTOMS IN COVID-19 PATIENTS

Dyspnea distinct as a respiratory rate above 30 breaths per minute, a blood oxygen saturation below 93%, a PaO₂:FIO₂ portion of 300 mmHg or less, and a percentage of air infiltration into the lungs more than 50% are all indicative of severe COVID-19 symptoms.¹⁶

CYTOKINE STORM MECHANISM IN COVID-19

Similar in appearance to SARS-CoV is SARS-CoV-2, a dissimilar betacoronavirus. In order to infect cells, both viruses rely on a protein known as angiotensin-converting enzyme 2 (ACE2). ACE2 receptors are another name for these receptors. These receptors may be found not just in cardiovascular tissue but also in hematological cells such as monocytes and macrophages. Lymphopenia is an essential component of a COVID-19 infection and is linked to the clinical severity of the condition. MERS-CoV uses dipeptidase 4 to infect monocytes and T cells, while

SARS-CoV uses dipeptide peptidase 4 to infect primary monocytes and dendritic cells. The possibility of dendritic cell infection by SARS-CoV-2 has also been aimed. It's possible that apoptosis of T cells, which happens when dendritic cells don't work right, also contributes to COVID-19's immunopathology.¹⁷

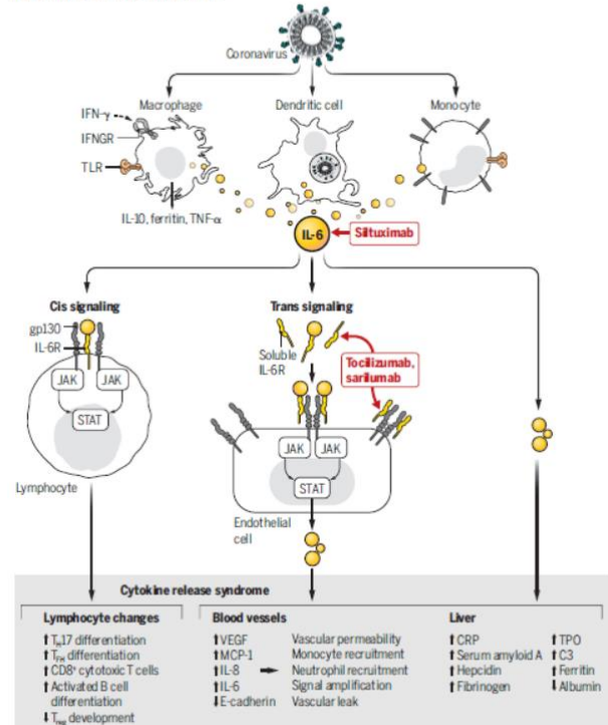
In both SARS and MERS infections, cytokine release syndrome is a leading cause of illness (CRS). After contracting MERS, a person's blood will show elevated levels of cytokines such as interleukin-6 (IL-6) and others involved in inflammation. Clinical symptoms such as ARDS and respiratory failure, as well as CAS, are often reported in individuals with COVID-19, and are linked to an increase in serum IL-6. When IL-6 is present, the inflammatory protein C-reactive protein (CRP) rises, serving as a biomarker for severe betacoronavirus infection.¹⁸

As a result of infection with the betacoronavirus, innate immune cells such as monocytes, macrophages, and dendritic cells mature and release inflammatory cytokines like interleukin 6. (IL-6). Both cis signaling and transsignaling are considered to be the primary traditional ways via which In the context of cis communication, IL-6 may serve as a messenger. At a complex containing gp130, IL-6 binds to the membrane-bound IL-6 receptor (mIL-6R); subsequent signals are translated by STAT3 and JAKS (Janus kinases) (signal transducer and activator of transcription 3). While gp130 is widely distributed, mIL-6R is exclusively found on immune cells. CRS is induced by the activation of cis signaling, which has pleiotropic effects, meaning it may influence both the adaptive and innate immune systems (B and T cells, neutrophils, macrophages, and natural killer (NK) cells).¹⁹

To activate trans-signaling, circulating IL-6 binds to the soluble IL-6R (sIL-6R), creating a complex with gp130 dimer potential on the cell surface. Signaling involving IL-6, sIL-6R, JAK, and STAT3 activates endothelial cells despite the fact that these cells do not express mIL-6R. Along with IL-8 and IL-6, the production of monocyte chemoattractant protein-1 (also known as MCP-1) and vascular endothelial growth factor (also known as VEGF) also occurs as a consequence of this process. On the other hand, the amount of E-cadherin produced by endothelial cells is lower. In ARDS, VEGF plays an important part in the pathophysiology of vascular permeability and leaky hypophysiology, in addition to pulmonary dysfunction, by lowering E-cadherin expression, which can be shown in Figure 1.²⁰

Pathways leading to cytokine release syndrome

Coronavirus infection results in monocyte, macrophage, and dendritic cell activation. IL-6 release then instigates an amplification cascade that results in cis signaling with T_H17 differentiation, among other lymphocytic changes, and trans signaling in many cell types, such as endothelial cells. The resulting increased systemic cytokine production contributes to the pathophysiology of severe COVID-19, including hypotension and acute respiratory distress syndrome (ARDS), which might be treated with IL-6 antagonists such as tocilizumab, sarilumab, and siltuximab.



CRP, C-reactive protein; IFN-γ, interferon-γ; IFNGR, IFN-γ receptor; IL, interleukin; IL-6R, IL-6 receptor; JAK, Janus kinase; MCP-1, monocyte chemoattractant protein-1; STAT3, signal transducer and activator of transcription 3; T_H, T follicular helper cell; T_H17, helper 17 cell; TNF-α, tumor necrosis factor-α; TLR, Toll-like receptor; TPO, thrombopoietin; T_H2, T regulatory cell; VEGF, vascular endothelial growth factor.

Figure 1. CRS lines on COVID-19²⁰

SARS-Cov-2 infection-related acute respiratory distress syndrome (ARDS) was fatally established in the clinical data obtained from COVID-19 patients with severe symptoms. This condition was commonly associated by organ failure and lung alveolar damage. Additional investigation found that the condition was associated with cytokine storm, commonly referred to as CRS, which is an increase in the synthesis of cytokines in the body. Cytokines are a class of tiny proteins that have a substantial function in the immunological response of the body, both to infections and to inflammation. But as a counterweight to overactive immune responses, excessive cytokine synthesis harms tissue.²¹

According to the findings of a number of research, the first phases of an infection are characterized by a pause in the release of cytokines and chemokines. After this, a limited amount of interferons (IFNs) are created, which is subsequently tracked by a rapid rise in pro-inflammatory of immune cell-attracting cytokines and chemokines. This leads to an excessive infiltration of lung tissue, which in turn causes damage to the lung tissue. Additionally, infected cells produce more chemokines to entice inflammatory mononuclear macrophages after activation of other signals (IMM). That leads to an abnormal increase in pro-inflammatory cytokine production, which only makes

things worse. IFN- γ and other proinflammatory cytokines drive T cells to commit suicide during the latter stages of the infection, which prevents the virus from being eliminated.²²

IFN- α and IFN- β , both of which are subtypes of IFN- γ , role as initial line of guard against viral infections when they are activated through the JAK-STAT pathway. An infection caused by a coronavirus may induce a strong but delayed response from the immune system's IFNs, which might foster the development of cytokine storms under certain conditions.²³

One of the proteins produced by the coronavirus, known as the NSP1 protein (non-structural protein 1), inhibits the phosphorylation of STAT1, which in turn stops the production of IFNs in the host cell. STAT1 is the transcription factor that is responsible for the expression of interferon stimulated genes (ISGs), which have a responsibility to produce antiviral defensive mechanisms. The composition of M (membrane) and N (nucleocapsid) proteins in coronaviruses allows them to block IFN signaling. One possible approach to getting target is to disrupt the activity of IRF3, a transcription factor for the IFN gene. Additionally, irregularities or inhibition of induction IFN brought on by the aging host and TRAF3's proteolytic degradation may contribute to the pathophysiology of the disease by creating an imbalance between pro-inflammatory cytokines and responses in aged individuals. These two elements may work together to cause an imbalance in pro-inflammatory cytokines and reactions in the aged.²⁴ It is solely for research purposes that leukocyte and cytokine counts are measured in patients with COVID-19.

RISKS OF CONVALESCENT PLASMA THERAPY

Similar to SARS, studies have revealed that viremia peaks during the first week after infection. It is more common for patients to develop an immunological response, which may lead to a potentially catastrophic cytokine storm, during the second week following the beginning of symptoms. Given that passive immunity via the administration of pathogen-specific antibodies is the basis for CP treatment, there are hidden dangers associated with CP infusion, such as the exacerbation of hyperimmune reactions. Although one research found that CP treatment could lessen serum cytokine responses depending on when the medication was administered. This has been corroborated by studies of SARS, lending more credence to the idea that treating CP at an earlier stage is preferable. Therefore, it is crucial to carefully time the administration of CP in COVID-19.²⁵

In reality, the titer of SARS-CoV2 neutralizing antibody (NAT) determines the therapeutic efficacy of CP on COVID-19. Finding of a study of people with SARS show that levels of a certain IgG started to rise in the third week after

symptoms began and peaked twelve weeks later. Additional studies have shown that CP with a NAT of 1:160 may reduce mortality from influenza. The CP isolated from patients who are still improving 12 weeks after symptom start with NAT of at least 1:160 is thought to be more potent. The capacity to get CP is limited, however, by factors such as the health of the donor, the availability of suitable donors, and the presence of informed permission.²⁵

All risks associated with transfusions must be taken into account. Transfusion of CP may lead to a variety of unwanted side effects, including fever, anaphylaxis, chills, hemolysis, circulatory overload, and transfusion-associated acute lung damage. When considering the safety of a CP transfusion, it is important to remember the risk of transmitting diseases like hepatitis and HIV.²⁵

CONVALESCENT PLASMA THERAPY PROCEDURES

The first step was identifying a suitable CP donor from among the COVID-19 patients who had been declared clinically negative with the PCR test twice in the span of more than a day, indicating that they had fully recovered and been released from the hospital at least two weeks before. Plasmapheresis is used to remove plasma from a patient throughout the healing process. Fresh frozen plasma (FFP) is used to create various plasma products.²⁶

Titer IgG antibodies are measured and reported in relation to S-RBD (Spike-Receptor Binding Protein). Only plasma units with IgG titers of at least 1:640 should be used to ensure therapeutic potential, as the Food and Drug Administration (FDA) recommendation.²⁶

The CP transfusion dosage for COVID-19 is around 4–13 milliliters per kilogram of the recipient's body weight. It is of the utmost importance that the ABO the patient blood type and the ABO type of the dispersed plasma be same. A convalescent plasma transfusion begins with 10 milliliters given over the course of the first 15 minutes, and the rate of administration is subsequently raised to 100 milliliters per hour while the patient is carefully monitored. When determining an appropriate transfusion rate, it is possible to take a patient's risk of fluid overload as well as their tolerance into consideration.²⁸

THE EFFECT MECHANISM OF CONVALESCENT PLASMA THERAPY

Antivirus mechanism

When it comes to getting rid of viruses, NAbs are crucial since they can defend against viral infections. Viruses and bacteria may be fought off with the help of passive immunity, which is powered by antibodies. Keep in mind that the plasma concentration of NAbs from the recovered donor has an influence on how well the treatment works. NAbs have been shown to bind to the S1RBD, S1-N, and

S2 terminal domains of the SARS and MERS viruses. The entry of these viral proteins is thereby inhibited, and viral multiplication is stymied. Activation of the complement system, antibody-dependent cellular cytotoxicity, and phagocytosis are all antibody-mediated processes that may improve the therapeutic efficiency of CP.²⁷

A single SARS-CoV-specific antibody, CA3022, was shown by Tian et al. to bind to the COVID-19 RBD and to not compete with ACE-2 for this binding. This was proved by the fact that the SARS-CoV-specific antibody bound to the COVID-19 RBD. Significant differences may be found between COVID-19 RBD and SARS-CoV at the C-terminus residue. Despite the fact that COVID-19 is unable to form a binding with the ACE-2 receptor, this variation has an effect on the cross-reactivity of NAbs.²⁷

Plasma includes NABs as well as the protective antibodies IgG and IgM. Improved prevention and/or treatment might be possible with the use of non-NAb antibodies that bind to the virus. IgG antibodies against N are initial seen after the first four days of symptoms in a person with SARS-CoV infection, and seroconversion occurs 14 days after infection. Up to 89% of cured SARS patients showed detectable levels of specific IgG and NABs 2 years after infection. IgM levels peaked 9 days after sickness onset, while IgG production took over after 2 weeks.²⁷

Donors who had previously been infected with COVID-19 but had recovered showed SARS-CoV2-specific antibody titers ranging from 1,800 to 16,200 and NABs titers from 80 to 480, as reported by Shen et al. Infections were reduced in those who got plasma that had been donated and administered on the same day. After receiving a CP transfusion, the recipient's IgG and IgM titers rose steadily. Defending against viral infection is a key function for NABs. In another study, researchers analyzed the temporal dynamics of the emergence of NABs that specifically target SARS-CoV-2. In SARS-CoV2 infect patients, NABs titers were low before day 10, rose with a peak 10–15 days following the beginning of disease, and remained constant thereafter in all patients.²⁷

Immunomodulating mechanism

According to F(ab')₂'s method of action, activated complement contributes significantly to inflammation throughout the body, neutrophils moving to the pulmonary, and tissue damage. Some antibodies block complement cascades (C3a and C5a), lending credence to this theory. Additional research has shown that plasma IgG inhibits the effects of cytokines including IL-1 β and TNF- α that can be seen on Figure 2.²⁷

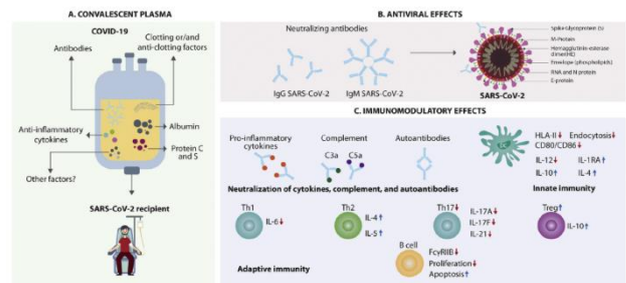


Fig. 1. Schematic representation of convalescent plasma components and its mechanisms of action. A. Main convalescent plasma components. B. Antiviral effects of NABs. IgG and IgM are the main isotypes, although IgA may be also important, particularly in mucosal viral infections. Other non-NABs may exert a protective effect. The humoral immune response is mainly directed towards spike (S) protein. C. Anti-inflammatory effects of CP include network of autoantibodies and control of an overactive immune system (i.e., cytokine storm, Th1/Th17 ratio, complement activation and regulation of a hypercoagulable state) (see text for details). N: Nucleoprotein; M: Membrane; E: Envelope.

Figure 2. Convalescent plasma effect mechanism²⁷

The interaction of NABs with Fc and/or complement receptors is the mechanism behind antibody-dependent enhancement (ADE), whereby infection severity is increased despite the relatively low levels of NABs present. In macrophages and other cells is beneficial for virus reproduction. Given the potential for ADE to have a detrimental impact affecting individuals with latent infections, it is important to keep this occurrence in mind while treating COVID-19 patients by promptly administering convalescent plasma.²⁷

Part-time regulation of IgG function is performed by FcRn. Antibody trafficking within the cell and subsequent excretion on the cell's rear surface is made possible by these receptors by blocking the breakdown and removal of IgG through a pinocytosis process. When FcRn is fully saturated, immunomodulatory pathways may be activated, which might benefit patients receiving convalescent plasma.²⁷

Fc- γ receptors are found on almost every kind of immune cell. These receptors are essential for the regulation and suppression of lymphocyte and other immune cell activities. When the Fc receptor is activated by IgG, the immune system is inhibited as a consequence of enhanced regulation of FcRIIB.²⁷

IgG plays a critical role in reducing inflammation by inhibiting dendritic cell maturation and activating B-catenin on its own. Research suggests that IgG may boost production of Th2 cytokines including IL-4 and IL-10 while reducing the number of Th1 cells, as well as FN- γ and TNF- α synthesis and TNF- α levels. By suppressing dendritic cells, which in turn stops signals to B cells, IgG limits the expansion of Th17 cells and reduces antigen presentation on T cells.²⁷

Efficacy of Convalescent Plasma Therapy in COVID-19 Patients with ARDS

Individuals diagnosed with ARDS who also met the criteria for severe pneumonia with rapid viral load progression, mechanical ventilation, and antiviral and methylprednisolone treatment were included in the COVID-19 case series published by Shen et al. (2020). Patients having a neutralization titer of higher than 40 and an antibody specificity

(IgG) titer of greater than 1:1000 (final dilution with ELISA) against SARSCoV-2 are given convalescent plasma transfusions.²⁸

Table 1. Analyzing Pre and Post Convalescent Plasma Transfusion Viral Load, Clinical Index, and Laboratory Results²⁸

	Patient				
	1	2	3	4	5
Clinical characteristics					
Body temperature, °C					
Just before transfusion	38.6	39.0	37.6	38.3	39.0
Day 1 posttransfusion	38.5	36.8	37.7	37.9	39.0
Day 3 posttransfusion	38.1	36.6	37.0	36.6	36.8
Day 7 posttransfusion	37.8	37.2	36.5	37.9	36.8
Day 12 posttransfusion	37.0	36.8	36.6	36.8	37.9
SOFA score ^a					
Just before transfusion	5	10	3	3	2
Day 1 posttransfusion	4	12	4	3	2
Day 3 posttransfusion	6	10	3	2	2
Day 5 posttransfusion	5	11	2	2	2
Day 7 posttransfusion	3	7	2	2	1
Day 12 posttransfusion	2	4	2	1	1
PaO ₂ /FIO ₂ ^b					
Just before transfusion	276	209	172	188	205
Day 1 posttransfusion	300	134	184	242	292
Day 3 posttransfusion	220	230	164	233	304
Day 7 posttransfusion	245	206	220	290	230
Day 12 posttransfusion	284	316	342	322	366
Ct value ^c (viral load proxy)					
On admission to hospital	23.0	19.7	18.9	38.0	28.0
Lowest value during hospitalization ^d (highest viral load)	19.2	19.7	18.9	26.6	26.5
Just before plasma transfusion	28.5	22.0	33.0	26.6	35.9
Day 1 posttransfusion	30.0	23.7	38.5	28.0	Negative
Day 3 posttransfusion	34.4	25.0	Negative	Negative	Negative
Day 7 posttransfusion	38.0	32.0	Negative	Negative	Negative
Day 12 posttransfusion	Negative	Negative	Negative	Negative	Negative
Mechanical ventilation					
Onset, days before transfusion	11	2	12	9	2
Extubated, days posttransfusion	Intubated	Intubated	2	9	9
ECMO					
Onset, days before transfusion	Not received	1	Not received	Not received	Not received
Removal, days posttransfusion	NA	5	NA	NA	NA
Laboratory findings					
C-reactive protein, mg/L (normal range, <8)					
Before transfusion	163.4	242.8	65.	156.0	173.1
Day 1 posttransfusion	146.2	223.0	108.3	NT	186.8
Day 3 posttransfusion	115.1	75.2	78.7	160.8	233.7
Day 5 posttransfusion	31.3	10.4	74.7	NT	260.4
Day 7 posttransfusion	31.2	13.9	6.2	9.6	5.5
Day 12 posttransfusion	5.3	33.1	NT	5.8	3.2
Procalcitonin, ng/mL (normal range, <0.1)					
Before transfusion	1.2	7.3	0.1	0.2	0.2
Day 1 posttransfusion	1.3	19.7	0.1	0.08	0.4
Day 3 posttransfusion	1.6	13.9	0.09	0.07	1.5
Day 5 posttransfusion	0.9	1.8	0.08	NT	0.9
Day 7 posttransfusion	1.1	0.1	0.04	0.04	0.09
Day 12 posttransfusion	0.4	0.2	NT	0.04	0.07
IL-6, pg/mL (normal range, 0-7)					
Before transfusion	70.5	438.2	63.9	79.1	87.8
Day 1 posttransfusion	74.9	NT	118.5	39.3	NT
Day 3 posttransfusion	34.5	1045.0	67.0	25.8	797.9
Day 5 posttransfusion	24.1	334.1	590.5	NT	NT
Day 7 posttransfusion	30.8	29.8	174.3	34.0	69.9
Day 12 posttransfusion	6.1	31.8	NT	2.7	54.9
Length of hospital stay, d					
	Remains hospitalized	Remains hospitalized	53	51	55
Current status as of March 25, 2020					
	Stable, still receiving mechanical ventilation	Stable, still receiving mechanical ventilation	Discharged home	Discharged home	Discharged home

Abbreviations: Ct, cycle threshold; ECMO, extracorporeal membrane oxygenation; NT, not tested.

^a The SOFA score is calculated using 6 systems: respiratory, coagulation, hepatic, cardiovascular, central nervous system, and kidney. A score of 0 is given for normal function through to 4 for most abnormal for each system. The worst values on each day are recorded, and the final SOFA score is the sum of the scores of each system.

^b PaO₂/FIO₂ ratio was defined as the ratio of the partial pressure of arterial oxygen to the percentage of inspired oxygen.

^c Cycle threshold is the number of polymerase chain reaction cycles required for gene amplification. A higher Ct value is correlated with a lower viral load.

^d Lowest value (highest viral load) between hospital admission and plasma transfusion.

Within three days of receiving plasma transfusion, four out of five patients had normalized body temperatures, and in the 12 days that followed, PaO₂/FIO₂ rose (before 172-276, after 284-366).

Antibody titers and neutralization levels improve following a transfusion (80-320 before 40-60 days), and virus loads drop and become negative within 12 days. Four patients with ARDS showed improvement by the 12th day after transfusion, and three patients were able to discontinue mechanical breathing after the second week of treatment. After 53, 51, and 55 days in the hospital, respectively, three of the five patients have been discharged, and 37 days after getting the blood transfusion, both of the remaining patients are doing well. Table 1 shows the variations in parameters before and after a CP transfusion.

An experiment was conducted and recorded by Simonovic et al. (2020) in which people with severe pneumonia cause of COVID-19 were randomly allocated to receive either CP or a placebo. 30 days after the intervention, the patient's clinical state was evaluated using a six-point ordinal scale to determine the study's result.²⁹

One hundred and five of the 228 convalescent plasma patients were given a placebo. The median antibody titer for SARS-CoV-2 in recovered individuals is 1:3200. (at now, between 1:800 and 3:1200) A severe study is defined by the presence of hypoxemia. Every patient is followed up with regularly. At day 30, there was no statistically significant difference between the two groups (control and convalescent plasma; odds ratio, 0.83; 95% CI, 0.52-1.35; P=0.46). Those who were given convalescent plasma had a 10% death rate, whereas those who were given a placebo had an 11% mortality rate. Both groups had the same drawbacks.²⁹

A comparison of the clinical outcomes of patients treated with convalescent plasma and those treated with placebo is shown in Figure 3.

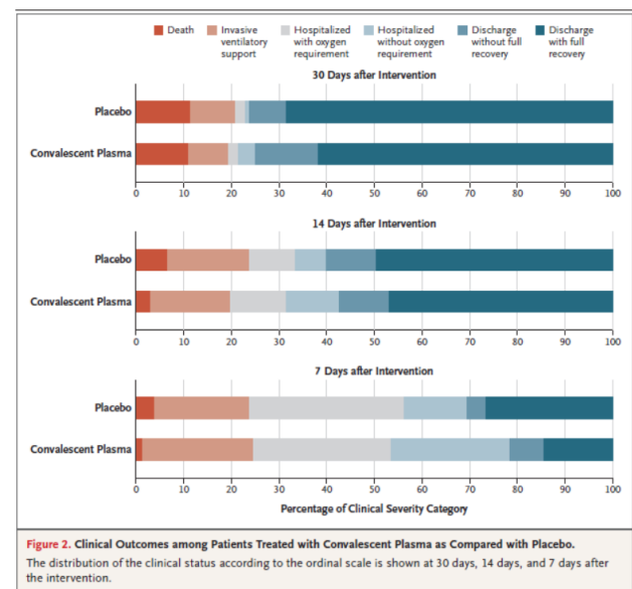


Figure 2. Clinical Outcomes among Patients Treated with Convalescent Plasma as Compared with Placebo.

The distribution of the clinical status according to the ordinal scale is shown at 30 days, 14 days, and 7 days after the intervention.

Figure 3. Comparing the Clinical Outcomes of Patients Treated with Convalescent Plasma with a Placebo²⁹

A review of the literature by Rajendran et al. Using electronic databases (PubMed, Embase, and Medline), a 2020 study of convalescent plasma reviewed five papers reporting 27 individuals.³⁰

The amount of recovered plasma that was utilized in each study had a different dosage. There was a single 200 mL CP dose utilized in a Chinese study, and the antibody titer was more than 1:640, while another Chinese study utilized a dose of 2400 mL in a male patient who was 73 years old. Convalescent plasma was administered to every patient between the days of 6 and 50 following the commencement of their symptoms or hospitalization.³⁰

Seventeen of the 21 patients receiving convalescent plasma treatment required mechanical ventilation due to the severity of their symptoms. Seventeen ARDS patients were given ECMO therapy. While hospitalization duration data was lacking, hospital discharge times were documented in almost all trials (15 total).³⁰

A considerable and unfavorable decline in viral load was seen between day 1 and day 30 following plasma injection in all 5 investigations. In a very short amount of time, almost all patients returned to baseline clinically, with improvements in temperature regulation, absorption of lung lesions, ARDS, and the ability to wean off of mechanical ventilation. After receiving a blood transfusion, the recovery time might range from 1 to 35 days.³⁰

CONVALESCENT PLASMA THERAPY IN PREGNANT WOMEN WITH COVID-19

Particular attention should be paid to controlling COVID-19 in pregnant women because of the risk of teratogenic consequences from antiviral drugs and immunosuppression caused by pregnancy. Recovery plasma EBM was studied by Franchini et al. (2021) in a clinical setting during pregnancy (see Table 2). Twelve pregnant women were reported as case reports in the research. Preeclampsia affected two women, but six mothers reported feeling OK. The future outlook for each mom is laid forth in detail. Survival, clinical progress, oxygenation, and recuperation are all parts of a full prognosis. Two infants were described as healthy, four as experiencing mild sickness, two as being in severe condition, one as having passed away, and three as not being noticed at all. All three pregnancies were healthy, and the baby is expected to do well.³¹

Eleven women and one mother experienced severe ARDS before beginning CP treatment. No co-morbid conditions were reported in any of the five patients. In our sample, five mothers had several chronic diseases, whereas the other two had simply obesity as a chronic illness. In CP treatment, gestational age may be anywhere from 21 weeks and 36 weeks and 2 days. Steroids (n = 8), heparin (n = 7), hydroxychloroquine (n = 5), human

monoclonal antibodies (tocilizumab, n = 2), and antivirals from the analog family of nucleotides (redeliver, n = 5) were all used throughout the patients' hospital stays. Of these nine patients, three had tracheostomies and three had extracorporeal membrane oxygenation/ECMO. Six patients required invasive mechanical ventilation.³¹

Table 2. A Synopsis of Convalescent Plasma Treatment for COVID-19 in Expectant Mothers.³¹

Author Year (Ref)	Design	Country	Age, y	Gestational Age	Severity of Disease	Comorbidity	Procedures	CP Treatment				Other Medications	Outcome	
								Units transfused	NAAT	Days from Hospitalization	AE		Maternal	Infant/Neonatal
Chen, 2020 (2)	CR	Italy	39	39w and 2 d	Mild ARDS	Class I obesity	VD	2	NR	<1, 44	None	Chloroquine, hydroxychloroquine, tocilizumab, remdesivir, lopinavir, ritonavir	Maternal well-being	Stillborn, well oxygenated with CP
Zhang, 2020 (3)	CR	China	35	39w and 2 d	Severe ARDS	-	ECMO, PE low flow (ECMO), ECMO	1	NR	<7	None	Logsdon's intravenous immunoglobulin, tocilizumab, remdesivir, lopinavir, ritonavir	Maternal well-being	Neonatal death due to placental abruption
Anderson, 2020 (4)	CR	USA	36	22w and 2 d	Severe ARDS	Type 2 DM, asthma, obesity, Class II obesity	ECMO	1	NR	<1	None	Remdesivir, hydroxychloroquine, tocilizumab, lopinavir, ritonavir, LMWH	Maternal well-being	Neonatal death due to congenital pneumonia
Uzunali, 2020 (5)	CR	Italy	36	27w and 4 d	Severe ARDS	-	BMV, PE low flow (ECMO), ECMO	2	NR	<1, 23	None	Chloroquine, hydroxychloroquine, tocilizumab, remdesivir, lopinavir, ritonavir, LMWH	Maternal well-being	Neonatal death due to congenital pneumonia
Luftman, 2020 (6)	CR	USA	40	36w	Severe ARDS	-	ECMO, PE low flow (ECMO), ECMO	1	NR	<5	None	Remdesivir, hydroxychloroquine, tocilizumab, lopinavir, ritonavir	Discharged with severe weight gain	Neonatal death due to congenital pneumonia
Maggiore, 2020 (7)	CR	Mexico	35	37w and 4 d	Severe ARDS	-	ECMO, PE low flow (ECMO), ECMO	2	NR	<1, 45	None	Logsdon's intravenous immunoglobulin, tocilizumab, remdesivir, lopinavir, ritonavir	Maternal well-being	Neonatal death due to congenital pneumonia
Moran, 2020 (8)	CR	USA	36	36w and 2 d	Severe ARDS, PE	Asthma, Class II obesity, Class II obesity, BMI 40	BMV, ECMO	1	NR	NR	NR	Hydroxychloroquine, tocilizumab, remdesivir, lopinavir, ritonavir, LMWH	Discharged to care of mother with severe weight gain	Neonatal death due to congenital pneumonia
Jahid, 2020 (9)	CR	Iran	36	36w and 1 d	Mild ARDS	-	ECMO	NR	NR	NR	NR	Remdesivir, hydroxychloroquine, tocilizumab, lopinavir, ritonavir	Maternal well-being	Neonatal well-being
Quaranta, 2020 (10)	CR	USA	32	22w and 4 d	Severe ARDS	-	ECMO, PE low flow (ECMO), ECMO	NR	NR	NR	NR	Hydroxychloroquine, tocilizumab, remdesivir, lopinavir, ritonavir, LMWH	Discharged to care of mother with severe weight gain	Neonatal death due to congenital pneumonia
Sakuma, 2020 (11)	CR	Iran	30	22w and 2 d	Severe ARDS	Class II obesity	-	2	NR	<10, 110	None	Logsdon's intravenous immunoglobulin, tocilizumab, remdesivir, lopinavir, ritonavir	Maternal well-being	Neonatal death due to congenital pneumonia
Lee, 2020 (12)	CR	USA	30	20w and 1 d	Severe ARDS	Type 2 DM, hypertension, obesity, Class II obesity	ECMO	NR	NR	<1	None	Remdesivir, hydroxychloroquine, tocilizumab, lopinavir, ritonavir	Discharged to care of mother with severe weight gain	Neonatal death due to congenital pneumonia
Nayfeh, 2020 (13)	CR	Qatar	35	32w	Severe ARDS	Asthma, gestational diabetes	ECMO, PE low flow (ECMO), ECMO	2	NR	<1	NR	Logsdon's intravenous immunoglobulin, tocilizumab, remdesivir, lopinavir, ritonavir	Clinical and discharged on CP	Neonatal death due to congenital pneumonia

Abbreviations: AE, adverse reactions to CP transfusion; ARDS, acute respiratory distress syndrome; BMI, body mass index; BMV, bilevel mechanical ventilation; Class I, low-molecular weight heparin; NAAT, nucleic acid testing; NR, not reported; PE, pulmonary embolism; PE, prone positioning; VD, vaginal delivery; y, years; w, weeks.

SAR-COV-2 seems to exacerbate clinical symptoms in both mothers and their babies. Premature delivery, maternal death, fetal death in utero, and newborn mortality are common outcomes of pregnancies interrupted by SAR-COV 2. There was a 5% maternal mortality rate and a 6% infant mortality rate, respectively. Although passive immunotherapy with CP transfusion is often deemed appropriate in the patient groups with such specific characteristics, to far only twelve cases of CP record in pregnant women were described. The average gestational age was 27.9 weeks, with a range of 22-36 weeks, and the average age of the patients was 32.0 years (range, 22-42 years). However, the majority of reported cases (i.e., in the third trimester of pregnancy) were in women younger than 35. Research shows that third-trimester SARS-CoV-2 infections are more dangerous.³³ Critically sick patients with moderate to severe ARDS have always been given CP. The high proportion of invasive operations (7/12.58.3%) needed to cure life-threatening hypoxia demonstrates the severity of respiratory disorders. These invasive procedures include invasive mechanical ventilation, tracheostomy, and oxygenation of the extracorporeal membrane.

According to the research that had been done in the past, pregnant women that positive COVID-19 and had an accompanying illness had a greater risk of having complications with their pregnancy. Seven of the twelve pregnant women polled had several medical problems (most commonly obesity, diabetes, and asthma). Six out of nine doctors agree that two CP units are necessary

for clinical improvement (56%). There is a wide range (1-17 days) between hospital admission and the first transfusion in a CP unit, although typically it takes 2 days. The anti-viral advantages of plasma hyperimmune are maximized when it is infused as soon as possible after hospitalization (ideally within 72 hours). 34-36 Unfortunately, only two CP units have the anti-SARSCov-2 neutralizing titer, a key metric for assessing CP effectiveness. However, CP transfusion has not been linked to any adverse effects, demonstrating its safety as a treatment option. In addition to hyperimmune plasma, a number of other medications, such as 1) antibiotics; 2) steroids; anticoagulants employing low molecular weight heparin (LMWH); 3) hydroxychloroquine; and 4) antiviral medicines employing lopinavir, ritonavir, or remdesivir, are utilized. These medications are used either in conjunction with hyperimmune plasma or as a second-line therapy after the initial one has been. In each and every case that was investigated and recorded, the only person who prevailed was the mother.

According to a review of the relevant data, convalescent plasma treatment while pregnancy with severe COVID-19 is beneficial for both mother and baby. Since current research are based on a single case report, they may be biased. This is why these studies are so low-quality. Well-designed and well-funded registries and research including pregnant women may help comprehend CP's role in treating COVID-19 throughout pregnancy.

CONCLUSION

When CP was utilized as a treatment for patients with severe COVID-19 symptoms and emergency situations, they got better faster (within 28 days) than when they just got regular treatment. Differences in the amount of plasma given and the number of antibodies in the convalescent plasma used may have a big effect on differences in clinical outcomes. This makes it hard to tell how well convalescent plasma treatment is working.

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