Kinetics of SARS-CoV-2 antibody responses pre-
COVID-19 and post-COVID-19 convalescent plasma
transfusion in patients with severe respiratory failure:
an observational case–control study

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ABSTRACT

Aims While the SARS-CoV-2 pandemic may be contained through vaccination, transfusion of convalescent plasma (CCP) from individuals who recovered from COVID-19 (CCP) is considered an alternative treatment. We investigate if CCP transfusion in patients with severe respiratory failure increases plasma titres of SARS-CoV-2 antibodies and improves clinical outcomes.

Methods Patients with COVID-19 (n=34) were consented for CCP transfusion and serial blood draws pretransfusion and post-transfusion. Plasma SARS-CoV-2 antireceptor binding domain (RBD) IgG and IgM titres were measured by ELISA serially, and compared with serial plasma titre levels from control patients (n=68). The primary outcome was survival at 30 days, and secondary outcomes were length of ventilator and/or extracorporeal membrane oxygenation (ECMO) support, and length of stay (LOS) in the hospital and in the intensive care unit (ICU). Outcomes were compared with matched control patients (n=34). Kinetics of antibodies and clinical outcomes were compared using LOess regression and ORs, respectively.

Results Prior to CCP transfusion, 74% of patients were anti-RBD seropositive for IgG (median 1:3200), and 81% were anti-RBD IgM seropositive (median 1:320), while 16% were seronegative. The kinetics of antibody titres in CCP recipients were similar to controls. CCP recipients presented with similar survival, duration on ventilatory and/or ECMO support, as well as ICU and hospital LOS compared with controls.

Conclusions CCP transfusion did not increase the kinetics of SARS-CoV2 antibodies and did not result in improved clinical outcomes in patients with COVID-19 with severe respiratory failure, suggesting that CCP may not be indicated in this category of patients.

INTRODUCTION

The current global health crisis posed by the SARS-CoV-2 pandemic demands urgent containment through vaccine development and distribution. The management of COVID-19 has nonetheless improved given refined supportive therapies including hyperoxygenation, steroids, remdesivir and anticoagulation.1 Another therapy that has been investigated is passive antibody administration through transfusion of convalescent plasma (CCP) (ie, plasma collected from individuals who have recovered from COVID-19) to prevent the development of severe COVID-19.2 Historically, CCP has been transfused successfully as postexposure prophylaxis and/or treatment for various pathogens, including other coronaviruses (eg, SARS-1, Middle East Respiratory Syndrome).3 Administration of CCP was first attempted during the early stages of the COVID-19 pandemic in China, where it was reported to confer clinical benefit as reflected by faster viral clearance and improved survival.4, 5 Today, over 250 000 patients have been transfused with CCP safely in the USA. The Mayo Clinic published preliminary results, citing that the CCP was associated with reduced mortality in recipients early after symptom onset compared with recipients hospitalised for at least 7 days in the intensive care unit (ICU).6 However, conflicting studies have shown no survival benefit when CCP is transfused to critically ill patients with advanced respiratory symptoms.7

The sponsoring institution of the current study, the University of Maryland Medical Center, has one of the highest acute level care and ICU capacities in the USA and has been uniquely prepared to treat COVID-19 with different emerging therapies, including CCP. We evaluated the longitudinal profiles of SARS-CoV-2 antibody titres in plasma from critically ill patients with COVID-19 before and after CCP transfusion and compared them to those measured in patients not transfused with CCP. Additionally, clinical outcomes of CCP recipients were compared with those from a matched control group.

METHODS

Study design

This is an observational retrospective control study to investigate the development of the humoral immune response to SARS-CoV-2 in CCP recipients (n=34) and compare it to the humoral response in a group of patients not treated with CCP (n=68, control A). A separate comparison of clinical outcomes is performed between CCP recipients and a matched control group of patients untreated with CCP (n=34, control B).
CCP treated subjects
Patients considered for enrollment in the study presented with severe COVID-19 and were hospitalised in the ICU at University of Maryland Medical Center and at two other sister hospitals. Patients were evaluated by an infectious disease clinician based on Federal Drug Administration (FDA) recommended guidance.2 An institutional ethics committee reviewed the indication of each CCP transfusion. Patients less than 18 years old were excluded. Informed consent was obtained, and CCP was transfused following FDA authorisation. All CCP transfusions occurred between 17 April 2020 and 19 July 2020. CCP units with a SARS-CoV-2 antibody titre >1:160, per FDA guidance, were procured by the regional blood centre.2 Patients received a single unit of ABO compatible CCP of approximately 250 mL. Following transfusion, CCP recipients were closely monitored for a minimum of 4 hours for possible transfusion-related adverse events. Blood samples for SARS-CoV-2 antibody titre measurements were collected immediately pre-transfusion (day 0) and on days 3, 7 and 14 post-transfusion. Data from three of the CCP recipients were excluded from the kinetics analysis due to insufficient plasma sample quantity; these were still included in the clinical outcome analysis.

Non-transfused control subjects
Non-transfused patients (control A) were used for comparison of antibody titres. Remnant plasma samples from non-transfused control A patients were aliquoted 1–3 days following collection and stored at −70°C prior to antibody titre measurement. Sample draws ranged from 0 to 48 days after the onset of symptoms, which varied in severity.

Non-transfused patients (control B), used for the clinical outcome analysis, were matched to CCP recipients based on sex, age, and on three levels of respiratory support requirement (non-ventilated, mechanically ventilated and ventilated with extracorporeal membrane oxygenation (ECMO)) and were admitted in the same hospital. Patients who were administered CCP at an outside institution prior to their admission, pregnant, or had instructions not to escalate care (do not intubate (DNI)/do not resuscitate (DNRI)) were excluded as controls. Seven non-transfused-patients were included in both control A and control B.

Clinical data collection and outcomes
After enrolment, the following clinical variables were collected from electronic medical records: symptoms at presentation, level of respiratory support (mechanical ventilation/ECMO status), comorbidities, other SARS-CoV-2 infections served as controls, patients with PCR confirmed SARS-CoV-2 infections were tested on the ELISA and the Ortho VITROS platform. On the ELISA, 22/24 samples were seropositive for anti-RBD IgG with a median titre of 1:6400 (figure 1A) while 23/24 were seropositive for anti-RBD IgM with a median titre of 1:240 (figure 1B). The Ortho VITROS platform detected SARS-CoV-2 antibodies in the majority of samples (22/24) at a median signal to cut-off ratio (S/C) of 490 (figure 1C). Two samples with IgM titres of 1:40 and undetectable IgG were negative by the Ortho VITROS method.

Characteristics of CCP recipients compared with non-transfused patients
CCP transfusion was considered and reviewed by an infectious disease expert, for 41 patients with COVID-19, of whom 34 patients were transfused with CCP on obtaining consent. Reasons for non-transfusion included patients or legally authorised proxy changing their mind about the treatment. The anti-RBD IgG and IgM responses of CCP recipients were compared with those of 68 non-transfused control patients (control A); CCP recipients presented with more severe disease requiring ECMO support, but both groups were similar in terms of sex and age (table 1). CCP recipients and matched non-transfused patient (control B) had similar frequencies of comorbidities, symptoms at presentation and other COVID-19 directed therapies administered during hospitalisation (table 1). ABO type distribution was different between the groups, although it was not available on eight of the 34 (23.5%) matched control patients (table 1).
Kinetics of anti-RBD IgG and IgM responses in CCP recipients and non-transfused patients

Anti-RBD IgG and IgM responses were examined based on titre levels measured in plasma samples drawn on successive days post-onset of symptoms (POS), starting on the day of transfusion in CCP recipients, which was a median of 12 days POS, and a median of 10 days POS in non-transfused control A patients.

The frequency of patients who generated an anti-RBD IgG and/or an IgM response was similar in CCP recipients compared with controls ((frequency of IgG response: 100% (31/31) and 100% (68/68)) (frequency of IgM response: 96.8% (30/31) and 100% (68/68)). Furthermore, the seroconversion rate for both anti-RBD IgG and IgM, analysed using a cumulative frequency plot, was similar in CCP recipients compared with controls (figure 2A,B). The longitudinal profiles of anti-RBD IgG responses showed a peak between 20 and 30 days POS and slowly decayed thereafter for both CCP recipients and controls (figure 2C). The anti-RBD IgM response peaked between 15 and 25 days POS and rapidly decayed thereafter for both CCP recipients and controls (figure 2D).

The kinetics of individuals’ anti-RBD IgG and IgM response were also compared between CCP recipients (figure 3A,B) and non-transfused control A patients (figure 3C,D). As was observed at the overall population level in figure 2, the kinetics of the anti-RBD IgG and IgM responses in CCP recipients and control groups at the individual patient level were similar (figure 3A–D).

### Table 1  Demographics and clinical characteristics

|                      | Overall (n=129) | CCP group (n=34) | Control A (n=68) | Control B (n=34) | P value CCP vs A | P value CCP vs B |
|----------------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| Male sex, n (%)*     | 88 (68.2)       | 23 (67.6)       | 46 (67.6)       | 23 (67.6)       | 1               | 1               |
| Age (years), mean (SD)† | 57.4 (16.4)   | 55.4 (16.6)     | 59.0 (16.8)     | 57.2 (15.3)     | 0.31            | 0.65            |
| Comorbidities, n (%)*|                 |                 |                 |                 |                 |                 |
| BMI >30              | 25 (19.4)       | 6 (17.6)        | 18 (26.5)       | 4 (11.8)        | 0.46            | 0.73            |
| Diabetes             | 50 (38.8)       | 14 (41.2)       | 28 (41.2)       | 12 (35.3)       | 1.00            | 0.80            |
| Hypertension         | 67 (51.9)       | 16 (47.1)       | 38 (55.9)       | 17 (50.0)       | 0.41            | 1.00            |
| Chronic obstructive pulmonary disease | 10 (7.8) | 3 (8.8)         | 7 (10.3)        | 2 (5.9)         | 1.00            | 1.00            |
| Chronic kidney disease | 14 (10.9)   | 2 (5.9)         | 11 (16.2)       | 1 (2.9)         | 0.21            | 1.00            |
| Hyperlipidaemia      | 36 (27.9)       | 7 (20.6)        | 27 (39.7)       | 4 (11.8)        | 0.07            | 0.51            |
| Coronary artery disease | 8 (6.2)     | 1 (2.9)         | 6 (8.8)         | 3 (8.8)         | 0.42            | 0.61            |
| Other therapies      | 2 (1.6)         | 2 (5.9)         | 0 (0.0)         | 0 (0.0)         | 0.11            | 0.49            |
| Symptoms, n (%)*     |                 |                 |                 |                 |                 |                 |
| Dyspnoea             | 118 (91.5)      | 32 (94.1)       | 59 (86.8)       | 33 (97.1)       | 0.33            | 1.00            |
| SpO2 <93%            | 107 (82.9)      | 28 (82.4)       | 51 (75.0)       | 34 (100.0)      | 0.46            | 0.03            |
| Respiratory rate >30 | 74 (57.4)       | 19 (55.9)       | 33 (48.5)       | 24 (70.6)       | 0.53            | 0.31            |
| Arterial O2/FiO2 <300| 107 (82.9)      | 34 (100.0)      | 45 (66.2)       | 33 (97.1)       | <0.001          | 1.00            |
| Respiratory failure  | 109 (84.5)      | 34 (100.0)      | 47 (69.1)       | 34 (100.0)      | <0.001          | 1.00            |
| Septic shock         | 48 (37.2)       | 15 (44.1)       | 25 (36.8)       | 13 (38.2)       | 0.52            | 0.81            |
| Disease severity n (%)* | 0.001          | 0.001           | 0.001           | 0.001           | 0.002           | 0.002           |
| Non-ventilated       | 39 (30.2)       | 6 (17.6)        | 29 (42.6)       | 6 (17.6)        |                 |                 |
| Mechanical ventilation| 65 (50.4)     | 17 (50.0)       | 35 (51.5)       | 17 (50.0)       |                 |                 |
| ECMO                 | 25 (19.4)       | 11 (32.4)       | 4 (5.9)         | 11 (32.4)       |                 |                 |
| ABO, n (%)*          |                 |                 |                 |                 | <0.001          | 0.002           |
| A neg                | 4 (3.1)         | 0 (0.0)         | 3 (4.4)         | 2 (5.9)         |                 |                 |
| A pos                | 30 (23.3)       | 6 (17.6)        | 17 (25.0)       | 9 (26.5)        |                 |                 |
| AB pos               | 2 (1.6)         | 0 (0.0)         | 2 (2.9)         | 0 (0.0)         |                 |                 |
| B neg                | 1 (0.8)         | 0 (0.0)         | 1 (1.5)         | 0 (0.0)         |                 |                 |
| B pos                | 20 (15.5)       | 8 (23.5)        | 10 (14.7)       | 2 (5.9)         |                 |                 |
| O neg                | 1 (0.8)         | 0 (0.0)         | 1 (1.5)         | 0 (0.0)         |                 |                 |
| O pos                | 45 (34.9)       | 20 (58.8)       | 14 (20.6)       | 13 (38.2)       |                 |                 |
| N/A                  | 26 (20.2)       | 0 (0.0)         | 20 (29.4)       | 8 (23.5)        |                 |                 |
| Other therapies, n (%)* |                |                 |                 |                 |                 |                 |
| Hydroxychloroquine   | 70 (54.3)       | 18 (52.9)       | 39 (57.4)       | 19 (55.9)       | 0.68            | 1.00            |
| Azithromycin         | 54 (41.9)       | 9 (26.5)        | 38 (55.9)       | 12 (35.3)       | 0.01            | 0.60            |
| Steroids             | 21 (16.3)       | 11 (32.4)       | 2 (2.9)         | 8 (23.5)        | <0.001          | 0.59            |
| Tocilizumab          | 30 (23.3)       | 14 (41.2)       | 10 (14.7)       | 7 (20.6)        | 0.01            | 0.11            |
| Remdesivir           | 25 (19.4)       | 7 (20.6)        | 11 (16.2)       | 7 (20.6)        | 0.59            | 1.00            |
| Stem cells           | 4 (3.1)         | 0 (0.0)         | 3 (4.4)         | 2 (5.9)         | 0.55            | 0.49            |

*Fisher’s exact test.
†Welch’s one-way ANOVA.
ANOVA, analysis of variance; BMI, body mass index; CCP, convalescent plasma; ECMO, extracorporeal membrane oxygenation; FiO₂, fractional inspired oxygen; N/A, not available; SpO₂, oxygen saturation.
Furthermore, IgG and IgM titres increased with higher respiratory support requirement in both CCP recipients and controls (figure 3A–D).

Immediately prior to CCP transfusion, 23/31 (74.2%) patients were anti-RBD IgG seropositive (median titre 1:3200, range 1:50–1:9600) (figure 3A), and 25/31 (80.6%) patients were anti-RBD IgM seropositive (median titre 1:320, range 1:40–1:640), (figure 3B), while 5/31 (16%) patients were seronegative for both. Interestingly, three out of these five CCP recipients died within 30 days of transfusion, one of whom was a recent kidney transplant recipient on immunosuppressive therapy. The highest anti-RBD IgG and IgM titres achieved by these three CCP recipients were similar to titres from non-transfused patients, measured at about the same number of days POS (figure 2C,D).

Anti-RBD IgG and IgM titre distribution in CCP recipients and in non-transfused patients

Next, we compared the distribution of anti-RBD IgG (figure 4A) and IgM (figure 4B) titres between CCP recipients and non-transfused patients (controls A) depending on the level of respiratory support needed. In non-ventilated patients, anti-RBD IgG and IgM titres were similar in CCP recipients compared with controls (IgG Median titres: 1:6400 and 1:3200 (figure 4A)), (IgM median titres: 1:480 and 1:160 (figure 4B)). In mechanically ventilated patients anti-RBD IgG and IgM titres were similar in CCP recipients compared with controls (IgG median titres: 1:12800 and 1:6400), (IgM median titres: 1:320 and 1:320 (figure 4B)). CCP recipients on ECMO had similar anti-RBD IgG titres compared with patients on ECMO (IgG median titres: 1:12800 and 1:6400). In contrast, IgM titre levels were higher in CCP recipients vs controls on ECMO (1:640 and 1:80, respectively) due to a difference in the number of days POS at which samples were drawn. Indeed CCP recipients and control A samples were not matched for POS for comparison. IgM measurements for CCP recipients on ECMO were taken at a median of 18 days POS compared with 28 days POS for control A patients on ECMO.

Clinical outcomes of CCP recipients and non-transfused (matched control B) patients

CCP recipients and matched control patients with COVID-19 (control B) presented with similar 30-day in-hospital mortality sample collection 52/68 (76.5%) patients were anti-RBD IgG seropositive (median titre 1:3200, range 1:100–1:6400), and 63/68 (92.6%) patients were anti-RBD IgM seropositive (median titre 1:160, range 1:70–1:640), while 4/68 (5.9%) patients were seronegative for both.
When stratifying the two groups based on disease severity, no difference in 30-day in-hospital mortality was observed (table 2). Additionally, CCP recipients and matched controls were similar in their respective median ICU LOS and median hospital LOS (table 2). The subgroups of CCP recipients also had similar ICU LOS and hospital LOS when compared with their respective matched control subgroups (table 2). CCP recipients and matched controls also had a similar median number of days on mechanical ventilation and median duration on ECMO (table 2).

**DISCUSSION**

The kinetics of SARS-CoV-2 IgG and IgM antibodies from plasma of patients with COVID-19 transfused with CCP were comparable to those from a cohort of patients with COVID-19 who did not receive CCP. Furthermore, most CCP recipients already had detectable SARS-CoV-2 antibodies in their plasma prior to transfusion. The highest SARS-CoV-2 antibody titres were observed in the plasma of the sickest subgroup of patients requiring both ventilatory and ECMO support. CCP recipients compared with matched control patients did not show any mortality benefit at 30 days post-transfusion, nor a reduction in either ICU or hospital LOS, or duration of mechanical ventilation/ECMO support; similarly subgroups comparisons based on disease severity, showed no difference in outcomes.

While some of the current findings corroborate results from earlier studies, they contradict others. Hegerova et al reported a modest survival benefit in a matched control study of patients with severe COVID-19 following CCP transfusion within 7 days of hospitalisation. A prospective, propensity score-matched study showed that patients transfused with CCP within 72 hours of admission experienced the most benefit compared with the control group. By contrast, in an open-label, randomised controlled trial (PLACID TRIAL), CCP was not associated with a reduction in overall mortality or progression to severe COVID-19, even when administered within 72 hours of symptoms onset. A retrospective study from March 2020 with 10 patients showed improved oxygenation and better patient survival following CCP transfusion. However, 4 of the 10 patients had high (≥1:640) SARS-CoV-2 neutralising antibody titres prior to CCP transfusion, suggesting that the patients' own immunity may have been responsible for the recovery rather than CCP transfusion. Nevertheless, these data suggest the need to prioritise CCP transfusion to patients with COVID-19 within 3–5 days of symptom onset when antibody production is still in the fledgling stages, or in those patients who are immunosuppressed (e.g., hypogammaglobulinaemia). However, in the current study, three of the CCP recipients who were seronegative prior to transfusion died within 30 days, one of these was a kidney transplant recipient who was receiving T-cell...
immunosuppression prior to COVID-19 diagnosis, suggesting that T cell response may also be important for controlling SARS-CoV-2 during the acute phase of the infection. T-cell immune responses were not assessed in the current study, but further flow cytometry analyses characterising the profile of lymphocyte subsets in patients with COVID-19 are underway in our laboratory to confirm results from other studies. Indeed a study by Peng et al showed that both SARS-CoV-2 antibody and memory T cell responses were stronger in patients who had recovered from severe disease.14 But patients with COVID-19 with severe disease and who died showed a significant quantitative and functional reduction in CD4 and CD8 T cells.15

The rise of antibody titres in the present study was expected to be higher in CCP recipients on days 2–3 post-CCP transfusion, but it remained similar to the rise observed in control patients. An increase in antibody titres may have been observed if the patients had been drawn for a plasma titre within 24 hours post-CCP transfusion. In a randomised control trial, PlasmAr Study, of 215 patients with severe pneumonia, total SARS-CoV-2 antibody titres were higher in the CCP treated group at day two post-transfusion. Still, no effect on 30-day clinical outcome and mortality between treated vs placebo groups was observed.15 Similarly, in the current study, most patients treated with CCP presented with severe COVID-19, requiring ventilatory support and/or ECMO. The antibody response increased proportionally with the severity of COVID-19, which was also previously reported.16–18

While the exact SARS-CoV-2 antibody titres in CCP units were unknown, these titres should have been greater than 1:160. Exogenous IgG antibodies typically have a half-life of 21 days and should last in the peripheral blood for at least 3 months.19 20 But the kinetics of the individual patient antibody response to SARS-CoV-2 make it difficult to determine the impact of CCP on the titres following CCP administration. Additionally, individuals that received CCP already had high antibody titres of 1:3200 on average and the addition of CCP was not likely to have a measurable impact on antibody titres given the dilutional effect. For example, if a patient received CCP with a 1:200 titre and the volume of plasma given accounted for less than 1/10 of the patient’s plasma volume, the end result would be a ~1:20 titre, which would be negligible for a patient with a 1:3200 titre.

The strength of this study is based on the characterisation of the kinetics of SARS-CoV-2 antibodies following CCP transfusion, which has not been previously described longitudinally in comparison to control plasma samples from non-transfused patients with COVID-19. Times of seroconversion and antibody kinetics in patients transfused with CCP were similar to that of non-transfused patients. These suggest that the antibodies measured are mostly the ones naturally produced by the patients’ immune system rather than those from the CCP treatment. These data on kinetics of SARS-CoV-2 antibodies are consistent with reports showing patients with COVID-19 in general with detectable IgG and IgM in plasma between four and 7 days POS.21 To strengthen the study, we compared the antibody titre measurements by ELISA to those obtained on a commercially available instrument, the Ortho VITROS total anti-SARS-CoV-2 Ig platform, which had been previously validated against a SARS-CoV-2 neutralising live-cell assay.22 23 The median IgG titres prior to CCP transfusion were high (1:3200). Interestingly, Salazar et al showed that anti-RBD IgG titres greater than 1:1350 correlated with SARS-CoV-2 neutralisation (VN) titres greater than 1:160 at 80% probability.22 The FDA (https://www.fda.gov/media/141477/download) recommended that an IgG titre detected on the Ortho VITROS platform at a S/C of 12 or on the Mount Sinai ELISA at 1:2880 may be considered equivalent to a VN titre >1:250 (https://www.fda.gov/media/141477/download). Furthermore, Luchsinger et al showed that both the Ortho VITROS total Ig and IgG platforms, set at a median S/C values of 101 and 11.7, respectively, correlated well to neutralising antibody results and gold-standard ELISAs.24 Our validation showed that the median anti-RBD IgG titre of 1:6400 in ELISA positive control samples, also tested by the Ortho VITROS total Ig method, showed a median S/C of 490 for total anti-SARS-CoV-2 Ig, suggesting that titres of 1:6400 and 1:3200 on the ELISA used in the present study are much higher than the recommended S/C of 12 and are indicative of high neutralising antibody titres.

There are limitations associated with this study. Although the blood supplier originally qualified the CCP donations as high titre (>1:160), in April 2020,2 the exact titres were not provided. Additionally, this is an observational study, thus the reliability in examining clinical outcomes compared with a prospective, randomised, control trial is not as robust; but at the advent of the first surge of the pandemic, a randomised trial was not practical at our institution. Lastly, the numbers of patients enrolled in each group are small, but the clinical outcomes of CCP recipients were compared with matched control patients.

Figure 4 Anti-RBD IgG and IgM response distributions in patients with COVID-19 stratified by disease severity and respiratory support needed. Distribution of anti-RBD IgG (A) and IgM (B) titres in CCP patients compared with controls depending on the level of respiratory support needed no ventilator (vent), ventilator only. Seronegative samples were excluded (A, B). Statistical analysis was performed using a Kruskal-Wallis test. An alpha value of 0.05 or less was considered statistically significant. All titre levels were converted to a log 10 scale. CCP, convalescent plasma; RBD, receptor binding domain.
Table 2  Comparison of secondary clinical outcomes in subgroups of COVID-19 severity

| Subgroup severity                  | Overall                  | CCP group                 | Control B                | OR (95% CI)   | P value |
|-----------------------------------|--------------------------|---------------------------|--------------------------|---------------|---------|
| Ventilatory support               | (n=68)                   | 25.0 (15.8, 44.5)         | 23.5 (16.0, 64.3)        | 28.0 (15.8, 44.0) | 1.01 (0.99 to 1.03) | 0.33 |
| ECMO support                      | (n=34)                   | 28.0 (18.0, 35.0)         | 21.0 (13.0, 32.8)        | 30.0 (21.0, 43.0) | 1.01 (0.98 to 1.05) | 0.52 |
| LOS in ICU                        | (n=34)                   | 28.0 (18.0, 35.0)         | 21.0 (13.0, 32.8)        | 30.0 (21.0, 43.0) | 1.01 (0.99 to 1.03) | 0.60 |
| LOS in hospital                   | (n=34)                   | 28.0 (18.0, 35.0)         | 21.0 (13.0, 32.8)        | 30.0 (21.0, 43.0) | 1.01 (0.99 to 1.03) | 0.60 |
| 30 days in-hospital mortality, n (%) | (n=22)                  | 17 (25.0)                 | 9 (26.5)                 | 8 (23.5)      | 1.12 (0.34 to 3.79) | 1 |
| Non-mechanically ventilated       | (n=22)                   | N/A                       | N/A                      | N/A           | N/A     |
| Mechanicaly ventilated            | (n=22)                   | 18.0 (13.0, 28.8)         | 18.0 (13.0, 25.0)        | 18.0 (7.0, 31.0) | 1.01 (0.97 to 1.05) | 0.64 |
| ECMO                              | (n=17)                   | N/A                       | N/A                      | N/A           | N/A     |
| LOS in ICU                        | (n=17)                   | 18.0 (13.0, 25.3)         | 18.0 (13.0, 26.0)        | 20.0 (10.0, 23.0) | 1.00 (0.96 to 1.05) | 0.98 |
| LOS in hospital                   | (n=17)                   | 27.0 (15.5, 63.9)         | 27.0 (17.0, 41.0)        | 27.0 (15.0, 37.0) | 1.01 (0.97 to 1.04) | 0.68 |
| 30 days in-hospital mortality, n (%) | (n=11)                  | 13 (38.2)                 | 7 (41.2)                 | 6 (35.3)      | 1.16 (0.27 to 5.17) | 1 |
| ECMO                              | (n=11)                   | 45.0 (31.0, 67.8)         | 65.0 (29.5, 70.5)        | 44.0 (23.5, 54.5) | 1.02 (0.99 to 1.07) | 0.26 |
| Mechanicaly ventilated            | (n=11)                   | 28.5 (19.3, 45.8)         | 28.0 (18.0, 55.0)        | 31.0 (21.0, 43.0) | 1.01 (0.98 to 1.05) | 0.52 |
| ECMO                              | (n=11)                   | 36.5 (25.0, 63.8)         | 38.0 (25.0, 70.0)        | 35.0 (25.5, 50.0) | 1.02 (0.99 to 1.06) | 0.29 |
| LOS in hospital                   | (n=11)                   | 50.5 (31.3, 69.8)         | 68.0 (29.0, 88.5)        | 48.0 (36.0, 59.0) | 1.02 (0.99 to 1.06) | 0.25 |
| 30 days in-hospital mortality, n (%) | (n=11)                  | 4 (18.2)                  | 2 (18.2)                 | 2 (18.2)      | 1.00 (0.06 to 16.14) | 1 |

Values are number of days as median (IQR).
Analysis of statistically significant differences between groups performed using an OR test with Wald 95% CIs.

CCP, convalescent plasma; ECMO, extracorporeal membrane oxygenation; ICU, intensive care unit; LOS, length of stay; N/A, not available.

hospitalised at the same hospital to decrease bias due to the clustering of enrollment.

In conclusion, the current data may further guide clinicians in defining eligibility criteria for CCP transfusion for the treatment of COVID-19. Indeed, these data do not support CCP transfusion to patients with severe COVID-19, especially if presenting with plasma SARS-CoV-2 IgG and IgM neutralising antibody levels at presentation. Taken together with the current literature, our findings confirm that CCP is probably most effective when administered to patients with low antibody titres, who are earlier in the disease course, and who do not yet have complicating COVID-19.

**Take home messages**

- Kinetics of SARS-CoV-2 antireceptor binding domain (RBD) IgG and IgM titres in convalescent plasma (CCP) recipients were similar to patients with COVID-19 not transfused.
- The majority of patients (74%) were seropositive for anti-RBD IgG and seropositive for IgM (81%), prior to transfusion with CCP.
- SARS-CoV2 antibodies were proportionally higher in patients with more severe COVID-19 requiring increased respiratory support.
- CCP recipients and matched controls showed similar 30-day survival and length of respiratory support as well as length of stay in the intensive care unit and in the hospital.

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REFERENCES
1. Sharma A, Tiwari S, Deb MK, et al. Severe acute respiratory syndrome coronavirus-2 (SARS-CoV-2): a global pandemic and treatment strategies. *Int J Antimicrob Agents* 2020;56:106054.
2. Bloch EM, Shoham S, Casadevall A, et al. Deployment of convalescent plasma for the prevention and treatment of COVID-19. *J Clin Invest* 2020;130:2757–65.
3. Casadevall A, Piccirillo L-A, L-a P. The convalescent sera option for containing COVID-19. *J Clin Invest* 2020;130:1545–8.
4. Li L, Zhang W, Hu Y, et al. Effect of convalescent plasma therapy on time to clinical improvement in patients with severe and life-threatening COVID-19: a randomized clinical trial. *JAMA* 2020;324:460–11.
5. Shen C, Wang Z, Zhao F, et al. Treatment of S critically ill patients with COVID-19 with convalescent plasma. *JAMA* 2020;323:1582–9.
6. Joyner MJ, Bruno KA, Klassen SA, et al. Safety update: COVID-19 convalescent plasma in 20,000 hospitalized patients. *Mayo Clin Proc* 2020;95:1888–97.
7. Simonovich VA, Burgos Pratx LD, Scibona P, et al. A randomized trial of convalescent plasma in Covid-19 severe pneumonia. *N Engl J Med* 2021;384:619–29.
8. Stadlbauer D, Amanat F, Chromikova V, et al. SARS-CoV-2 seroconversion in humans: a detailed protocol for a serological assay, antigen production, and test setup. *Curr Protoc Microbiol* 2020;57:e100.
9. Hegerova L, Gooley TA, Sweerus KA, et al. Use of convalescent plasma in hospitalized patients with COVID-19: case series. *Blood* 2020;136:759–62.
10. Salazar E, Christensen PA, Graviss EA, et al. Treatment of coronavirus disease 2019 patients with convalescent plasma reveals a signal of significantly decreased mortality. *Ann J Pathol* 2020;190:2289–303.
11. Agarwal A, Mukherjee A, Kumar G. Convalescent plasma in the management of moderate covid-19 in adults in India: open label phase II multicentre randomised controlled trial (PLACID trial). *BMJ* 2020;151:m3939.
12. Duan K, Liu B, Li C, et al. Effectiveness of convalescent plasma therapy in severe COVID-19 patients. *Proc Natl Acad Sci U S A* 2020;117:9490–6.
13. Luetkens T, Metcalf R, Planelles V, et al. Successful transfer of anti-SARS-CoV-2 immunity using convalescent plasma in an MM patient with hypogammaglobulinemia and COVID-19. *Blood Adv* 2020;4:4864–8.
14. Peng Y, Mentzer AJ, Liu G, et al. Broad and strong memory CD4+ and CD8+ T cells induced by SARS-CoV-2 in UK convalescent individuals following COVID-19. *Nat Immunol* 2020;21:1336–45.
15. Diao B, Wang C, Tan Y, et al. Reduction and functional exhaustion of T cells in patients with coronavirus disease 2019 (COVID-19). *Front Immunol* 2020;11:827.
16. Seow J, Graham C, Merrick B, et al. Longitudinal observation and decline of neutralizing antibody responses in the three months following SARS-CoV-2 infection in humans. *Nat Microbiol* 2020;5:1598–607.
17. Wang Y, Zhang L, Sang L, et al. Kinetics of viral load and antibody response in relation to COVID-19 severity. *J Clin Invest* 2020;130:5235–44.
18. Klein SL, Pekosz A, Park H-S, et al. Sex, age, and hospitalization drive antibody responses in a COVID-19 convalescent plasma donor population. *J Clin Invest* 2020;130:6141–50.
19. Mankański S, Lee M, Fischer S, et al. The half-lives of IgG subclasses and specific antibodies in patients with primary immunodeficiency who are receiving intravenously administered immunoglobulin. *J Lab Clin Med* 1988;112:634–40.
20. Waldmann TA, Strober W. Metabolism of immunoglobulins. *Prog Allergy* 1969;13:1–110.
21. Sethuraman N, Jeremiah SS, Ryo A. Interpreting diagnostic tests for SARS-CoV-2. *JAMA* 2020;323:2249–51.
22. Salazar E, Kuchipudi SV, Christensen PA, et al. Convalescent plasma anti-SARS-CoV-2 spike protein ectodomain and receptor-binding domain IgG correlate with virus neutralization. *J Clin Invest* 2020;130:6728–38.
23. Luchtinger LL, Ransengna BP, Jin DK, et al. Serological assays estimate highly variable SARS-CoV-2 neutralizing antibody activity in recovered COVID-19 patients. *J Clin Microbiol* 2020;58:e02005–20.