Early but not late convalescent plasma is associated with better survival in moderate-to-severe COVID-19

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Abstract

Background

Limited therapeutic options exist for coronavirus disease 2019 (COVID-19). COVID-19 convalescent plasma (CCP) is a potential therapeutic, but there is limited data for patients with moderate-to-severe disease.

Research question

What are outcomes associated with administration of CCP in patients with moderate-to-severe COVID-19 infection?

Study design and methods

We conducted a propensity score-matched analysis of patients with moderate-to-severe COVID-19. The primary endpoints were in-hospital mortality. Secondary endpoints were number of days alive and ventilator-free at 30 days; length of hospital stay; and change in WHO scores from CCP administration (or index date) to discharge. Of 151 patients who received CCP, 132 had complete follow-up data. Patients were transfused after a median of 6 hospital days; thus, we investigated the effect of convalescent plasma before and after this timepoint with 77 early (within 6 days) and 55 late (after 6 days) recipients. Among 3,217 inpatients who did not receive CCP, 2,551 were available for matching.

Results

Early CCP recipients, of whom 31 (40%) were on mechanical ventilation, had lower 14-day (15% vs 23%) and 30-day (38% vs 49%) mortality compared to a matched unexposed
cohort, with nearly 50% lower likelihood of in-hospital mortality (HR 0.52, [95% CI 0.28–0.96]; \(P = 0.036\)). Early plasma recipients had more days alive and ventilator-free at 30 days (+3.3 days, [95% CI 0.2 to 6.3 days]; \(P = 0.04\)) and improved WHO scores at 7 days (-0.8, [95% CI: -1.2 to -0.4]; \(P = 0.0003\)) and hospital discharge (-0.9, [95% CI: -1.5 to -0.3]; \(P = 0.004\)) compared to the matched unexposed cohort. No clinical differences were observed in late plasma recipients.

**Interpretation**

Early administration of CCP improves outcomes in patients with moderate-to-severe COVID-19, while improvement was not observed with late CCP administration. The importance of timing of administration should be addressed in specifically designed trials.

**Introduction**

Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), a positive-sense, single-stranded RNA virus of the *Coronaviridae* family, is the etiologic agent of Coronavirus disease 2019 (COVID-19) and is responsible for greater than 126 million global infections and approximately 2.7 million deaths since December 2019 [1–3]. During the first 6 months of the COVID-19 pandemic, treatment was largely supportive. In June 2020 the Randomized Evaluation of Covid-19 Therapy (RECOVERY) trial reported that dexamethasone reduced mortality in patients on respiratory support; thus, corticosteroids have become standard of care for patients on supplementary oxygen [4, 5]. Furthermore, remdesivir, a nucleotide analogue that disrupts viral replication, is the only other pharmacologic approved for COVID-19 in the United States [6], although with little to no effect on mortality or clinical course in hospitalized patients [7]. No additional therapeutics have yet to show efficacy in hospitalized patients with COVID-19.

Convalescent plasma containing antibodies generated following pathogens’ exposure has been employed in past epidemics as a means of passively transferring immunity from individuals with resolved infection. In addition to antibody-mediated protection via neutralization, therapeutic antibodies can directly induce cellular cytotoxicity, complement activation, and phagocytosis. Furthermore, convalescent plasma may contain beneficial anti-inflammatory cytokines, defensins, and pentraxins that quell a severe inflammatory response [8]. During the SARS [9] epidemic in 2003 and more recently during the H1N1 [10] pandemic of 2009, treatment with convalescent plasma resulted in significant mortality reduction.

In the current pandemic, single-arm observational studies have reported administration of COVID-19 convalescent plasma (CCP) to patients with mild to severe COVID-19 with variable results and limited cohort sizes [11–15]. Two randomized open-label trials were terminated prior to full enrollment; one due to drop in local COVID-19 incidence and a second due to concerns that the recruited cohort had preexisting anti-SARS-CoV-2 antibodies prior to enrollment [16, 17]. A third, randomized open-label trial of 464 hospitalized adults with moderate COVID-19 failed to show benefit in 28-day mortality or progression to severe disease; however, the study was limited by low or undetectable antibody titers in the donor plasma [18]. Most recently, a randomized, double-blinded, placebo-controlled trial found that administration of higher-titer CCP within 72 hours after symptom onset among mildly infected older adults reduced risk of progression to severe respiratory disease by 48% [19].
There is increasing evidence that clinical response to CCP might depend on the timing of administration. We conducted a propensity score matched analysis in one of the largest reported cohort of hospitalized patients with moderate-to-severe COVID-19 to investigate outcomes associated with early versus late transfusion of CCP.

Methods

Study design

This was a cohort study that included 3,368 patients hospitalized with moderate (supplemental oxygen flow rate of $\geq 3$L/min) to severe or life-threatening (respiratory failure, shock, or multi-organ failure) COVID-19 managed within the Yale New Haven Health system (YNHHS) from March 8, 2020 to July 25, 2020. Of those, a total of 151 patients were transfused with CCP. We dichotomized the cohort based on the median time to CCP administration (6 days; interquartile range: 3, 11 days): The early CCP cohort received treatment within 6 days of hospitalization, while the late CCP cohort received treatment after 6 days of hospitalization. This cut-off was consistent with transfusion time used in another study \[15\]. For the early or late CCP cohorts, we conducted distinct propensity score matchings to generate two 1:1 matched unexposed cohorts, as previously described \[20\]. Index dates for the unexposed patients were assigned corresponding to the day of CCP administration within each matched pair.

This study was approved by the Yale University Institutional Review Board (HIC#: 2000027871) with a waiver of informed consent.

Eligibility and selection of convalescent plasma recipients

In April 2020 the U.S. Food and Drug Administration (FDA) provided physicians access to convalescent plasma through the US National Expanded Access Program (EAP), which was led by the Mayo Clinic (IRB# 20–003312). Patients hospitalized with COVID-19 within YNHHS were screened for eligibility to receive CCP through the EAP based on shared decision making between the Convalescent Plasma Clinical Team, primary medical team, and the patient or Legally Authorized Representative. Patients were eligible to receive CCP if they were 18 years of age or older, had a laboratory confirmed diagnosis of SARS-CoV-2 infection, were hospitalized within YNHHS with COVID-19 complications, and had moderate-to-severe disease as indicated by the following characteristics: 1. Supplementary oxygen requirements ($\geq 3$L/min nasal cannula) with pulmonary infiltrates on chest imaging; 2. Refractory acute respiratory failure; 3. Septic shock; or 4. Multisystem organ dysfunction. Absolute contraindication to the administration of CCP was confirmed new thromboembolic event. Relative contraindications were: 1. Confirmed or high suspicion for bacterial or fungal infection; 2. Suspicion of a new thromboembolic event; 3. Recent significant hemorrhage; 4. High risk for hemorrhage and on therapeutic anticoagulation; and 5. Severe IgA deficiency. At least two physicians on the Convalescent Plasma Clinical Team reviewed each CCP request and approved them based on the adherence to the inclusion criteria and absolute or relative contraindications.

Convalescent plasma donors

CCP was obtained from the hospital system’s blood suppliers, including New York Blood Center (New York, NY), Rhode Island Blood Center (Providence, RI), and the American Red Cross (Farmington, CT). Over the course of the study period the criteria for eligibility for CCP donation was modified by the FDA, though generally required PCR-confirmed diagnosis of
COVID-19 and complete resolution of symptoms for greater than 14 days prior to donation. The units transfused were not labeled by the blood suppliers with an anti-SARS-CoV-2 IgG titer.

Preparation and administration of convalescent plasma
Each patient received one unit of ABO compatible CCP, with a typical unit being approximately 200 mL in volume. Units were thawed on demand once requested and the patient was confirmed to have met all eligibility criteria.

Outcome measures
The primary endpoint was in-hospital mortality. Time was censored on the date of discharge or administratively on August 3, 2020, the date of closure of data extraction. Secondary endpoints included days alive and free from mechanical ventilation 30 days post-index date, length of hospital stay post CCP or index date in the unexposed cohort, and change in 8-point WHO ordinal scale at 7 days post CCP or index date and at hospital discharge (S1 Table in S1 File).

Statistical analysis
Patient demographics and pre-existing comorbidities were collected at admission. Vital signs, oxygen therapy, concomitant medications, and laboratory tests were collected longitudinally during the hospitalization. Descriptive statistics were used to summarize patient demographics and clinical characteristics. In bivariate analysis, two-sample Student’s t-test, Wilcoxon rank-sum test, Chi-square or Fisher’s exact tests were used for comparisons of exposed and unexposed cohorts, as appropriate.

Two separate propensity-score matches were carried out to select unexposed patients for the early CCP cohort or unexposed patients for the late CCP cohort (S2 and S3 Figs in S1 File). CCP recipients were matched with unexposed patients using both propensity scores and exact matching on the worst WHO scores preceding CCP administration or index date, and on administration of remdesivir. No replacement was allowed for matching, and matched unexposed patients in the early cohort were allowed in the late cohort. The standardized mean difference between matched groups was set at \( \leq 0.25 \). Because of the stringent exact matching criteria, the optimal algorithm that minimized differences between the matching pairs was only able to yield 1:1 unexposed match. Group comparisons were performed to confirm expected balance between matched groups. For each matched cohort, a multivariate logistic regression model was fitted to estimate the probability of receiving CCP within the specified time frame and create propensity for each individual (S2 Table in S1 File). The models included demographic and clinical characteristics at hospital admission; concomitant medications, laboratory results prior to CCP administration, and supplemental oxygen status as covariates. The list of covariates is presented in S2 Table in S1 File.

For the primary analysis, we used the Kaplan-Meier product-limit estimator to estimate the hospital mortality function. We compared CCP exposed and unexposed patients for the early and for the late cohorts using the log-rank test. Mortality at 14 days and 30 days post CCP or index date was estimated and reported with 95% confidence intervals. Secondary outcomes were analyzed using mixed effects models or ordinal logistic regression with robust variance estimation to compare treatment groups with their respective matched unexposed cohort. The effect sizes are presented as mean difference with 95% confidence interval. A two-sided alpha level of 0.05 was required for statistical significance. All statistical analyses were performed using SAS 9.4 (Cary, NC) software.
Results

Of 3,368 patients with confirmed SARS-CoV-2 infection within YNHHS, 151 eligible patients received CCP under the US National Expanded Access Program (Fig 1). Among the CCP recipients, 80 received CCP within 6 days after admission, and 71 received CCP after 6 days of hospitalization. Of those who received CCP, 3 early and 16 late recipients were excluded from propensity score matching due to missing laboratory values and/or supplemental oxygen information during the first week of hospitalization, leaving a total of 77 CCP patients in the early cohort and 55 CCP patients in the late cohort available for inclusion in the analyses. One of three excluded early recipients died (33%), and 9 out of 16 excluded late recipients died (56.3%). Among 3,217 SARS-CoV-2 positive hospitalized patients who did not receive CCP transfusion, 666 were excluded from the unexposed cohort due to incomplete clinical data, leaving 2,551 patients eligible for propensity score matching.

Baseline demographic characteristics and clinical factors for the early and late cohorts are summarized in Table 1. Overall, age, sex, race, ethnicity, BMI, and Charlson Comorbidity Index were equally distributed between the CCP and the unexposed group. Likewise, the mean maximum WHO score prior to the index date and the level of supplemental oxygen were comparable between CCP and unexposed group. Baseline laboratory values obtained prior to CCP

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**Fig 1. Consort selection tree.** Patient disposition in the observational cohort analysis.

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Table 1. Baseline demographic characteristics and clinical factors.

|                              | All Patients (N = 3368) | CCP Early (n = 77) | Unexposed Early (n = 77) | P value | CCP Late (n = 55) | Unexposed Late (n = 55) | P value |
|------------------------------|-------------------------|--------------------|--------------------------|---------|------------------|--------------------------|---------|
| **Demographics**             |                         |                    |                          |         |                  |                          |         |
| Age, years                   | 64 ± 20                 | 60 ± 14            | 62 ± 17                  | 0.49    | 66 ± 13          | 66 ± 15                  | 0.95    |
| Gender, n (%)                |                         |                    |                          |         |                  |                          |         |
| Female                       | 1718 (51)               | 34 (44)            | 35 (46)                  | 17 (31) | 25 (46)          |                          |         |
| Male                         | 1650 (49)               | 43 (56)            | 42 (55)                  | 38 (69) | 30 (55)          |                          |         |
| Race, n (%)                  |                         |                    |                          |         |                  |                          |         |
| White                        | 1610 (48)               | 24 (31)            | 24 (31)                  | 24 (44) | 27 (49)          |                          |         |
| Black or AA                  | 870 (26)                | 18 (23)            | 17 (22)                  | 10 (18) | 14 (26)          |                          |         |
| Other                        | 888 (26)                | 35 (46)            | 36 (47)                  | 21 (38) | 14 (26)          |                          |         |
| Ethnicity, n (%)             |                         |                    |                          |         |                  |                          |         |
| Hispanic or Latino           | 873 (256)               | 33 (43)            | 31 (40)                  | 19 (35) | 17 (31)          |                          |         |
| Other                        | 2495 (74)               | 44 (57)            | 46 (60)                  | 36 (66) | 38 (69)          |                          |         |
| BMI                          | 30 (8)                  | 33 (11)            | 33 (10)                  | 0.97    | 31 (7)           | 32 (8)                   | 0.73    |
| **Comorbidities**            |                         |                    |                          |         |                  |                          |         |
| Charlson Comorbidity Index   | 5 ± 4                   | 4 ± 3              | 4 ± 3.4                  | 0.43    | 5 ± 3            | 5 ± 3                    | 0.82    |
| Hypertension, n (%)          | 2322 (69)               | 43 (55)            | 45 (58)                  | 0.74    | 43 (78)          | 44 (80)                  | 0.81    |
| Diabetes mellitus, n (%)     | 1354 (40)               | 27 (35)            | 30 (39)                  | 0.62    | 28 (51)          | 28 (51)                  | 1.00    |
| Chronic kidney disease, n (%)| 851 (32)                | 11 (14)            | 18 (23)                  | 0.15    | 23 (42)          | 14 (26)                  | 0.07    |
| Chronic lung disease, n (%)  | 1095 (33)               | 24 (31)            | 26 (34)                  | 0.73    | 19 (35)          | 21 (38)                  | 0.69    |
| Heart disease (CVD), n (%)   | 792 (24)                | 11 (14)            | 12 (16)                  | 0.82    | 9 (16)           | 9 (16)                   | 1.00    |
| Smoke, n (%)                 | 1482 (44)               | 29 (38)            | 28 (36)                  | 19 (35) | 17 (31)          |                          |         |
| Never smoke                  | 1886 (56)               | 48 (62)            | 49 (64)                  | 36 (66) | 38 (70)          |                          |         |
| O2 by face mask or nasal cannula, n (%) | 1265 (39) | 4 (5.2%)          | 4 (5.2%)                 | 1.00    | 2 (4)           | 2 (4)                    | 1.00    |
| Non-Invasive ventilation or high-flow O2, n (%) | 810 (24)   | 41 (53)           | 41 (53)                  | 1.00    | 24 (44)         | 24 (44)                  | 1.00    |
| Intubation and mechanical ventilation, n (%) | 386 (12)   | 31 (40)           | 31 (40)                  | 1.00    | 29 (53)         | 29 (53)                  | 1.00    |
| Max FiO2                     | 35 ± 27                 | 76 ± 35            | 65 ± 37                  | 0.08    | 76 ± 34          | 68 ± 36                  | 0.23    |
| ECMO, n (%)                  | 10 (1)                  | 0                 | 2 (3)                    | 0.50    | 1 (2)           | 1 (2)                    | 1.00    |
| **Lab value (maximum value prior to index treatment day)** | | | | | | | |
| ALC, x 1000/μL               | 1.7 ± 5.9               | 1.4 ± 0.6          | 1.4 ± 0.7                | 0.70    | 1.7 ± 3.8       | 1.3 ± 0.5                 | 0.45    |
| D-dimer, mg/L                | 4.7 ± 5.9               | 9.4 ± 10.8         | 9.3 ± 10.2               | 0.96    | 8.2 ± 8.6       | 6.2 ± 5.5                 | 0.17    |
| Creatinine, mg/dL            | 36.9 (92.1)             | 67.5 (176.7)       | 72.5 (186.1)             | 0.86    | 53.0 (90.1)     | 44.9 (80.2)               | 0.62    |
| AST, U/L                     | 114 ± 503               | 354 ± 1352         | 113 ± 128                | 0.12    | 139 ± 162       | 136 ± 157                 | 0.92    |
| ALT, U/L                     | 76 ± 190                | 185 ± 445          | 92 ± 109                 | 0.08    | 81.2 ± 86.1     | 101 ± 141                 | 0.38    |
| ALP, U/L                     | 104 ± 79                | 117 ± 88           | 98 ± 41                  | 0.09    | 107 ± 66        | 102 ± 69                  | 0.70    |
| NT-proBNP, pg/mL             | 3.3 ± 9.6               | 1.3 ± 2.3          | 1.8 ± 5.9                | 0.50    | 4.1 ± 11.2      | 3.0 ± 9.1                 | 0.58    |
| Fibrinogen, mg/dL            | 575 ± 157               | 654 ± 143          | 643 ± 120                | 0.62    | 674 ± 139       | 597 ± 139                 | 0.005  |
| C-reactive protein, mg/dL    | 12.5 ± 8.7              | 17.7 ± 8.0         | 17.7 ± 8.1               | 0.96    | 17.8 ± 8.4      | 15.8 ± 7.2                | 0.17    |
| **Medications, n (%)**       |                         |                    |                          |         |                  |                          |         |
| Hydroxychloroquine           | 2229 (66)               | 53 (69)            | 53 (69)                  | 1.00    | 53 (96)         | 48 (87)                  | 0.16    |
| Dexamethasone                | 67 (2)                  | 4 (5)              | 4 (5)                    | 1.00    | 3 (5)           | 1 (2)                    | 0.62    |
| Remdesivir                   | 305 (9)                 | 38 (49)            | 38 (49)                  | 1.00    | 13 (24)         | 13 (24)                  | 1.00    |
| Received within 6 days       | 226 (7)                 | 30 (39)            | 30 (39)                  | 10 (18) | 10 (18)         |                          |         |
| Received after 6 days        | 74 (2)                  | 8 (10)             | 8 (10)                   | 3 (6)   | 3 (6)           |                          |         |

(Continued)
administration or index date were similar among the CCP and unexposed group, with the exception of mean fibrinogen level which was higher in the CCP treated group compared with the unexposed group in the late cohort. Concomitant medications, including hydroxychloroquine, remdesivir, tocilizumab, and dexamethasone were well balanced in the early cohort. Tocilizumab was administered more frequently in the CCP group compared with the unexposed group in the late cohort (Table 1).

**Primary endpoint**

In the propensity score matched analysis, the estimated in-hospital mortality was 15% by day 14 among early CCP recipients, compared with 23% in their matched unexposed cohort. At day 30, estimated in-hospital mortality was 38% in early CCP recipients compared to 49% in their matched unexposed cohort (Table 2). Overall, early CCP recipients were nearly 50% less likely to die in the hospital compared to patients in the matched unexposed cohort (HR 0.52, [95% CI 0.28 to 0.96]; P = 0.036) (Fig 2A). In contrast, there were no differences in mortality

Table 1. (Continued)

|                      | All Patients (N = 3368) | CCP Early (n = 77) | Unexposed Early (n = 77) | P value | CCP Late (n = 55) | Unexposed Late (n = 55) | P value |
|----------------------|-------------------------|--------------------|--------------------------|---------|------------------|-------------------------|---------|
| Tocilizumab          | 1304 (39)               | 69 (90)            | 67 (87)                  | 0.38    | 53 (96)          | 46 (84)                 | 0.03*   |
| Pressor use, n (%)   | 528 (16)                | 35 (46)            | 31 (40)                  | 0.51    | 42 (76)          | 33 (60)                 | 0.07    |

Comparative baseline demographics, comorbidities, laboratory values, and medications among all COVID-19 patients, early CCP recipients, late CCP recipients and the respective early/late unexposed cohorts. P-values represent significance of bivariate analyses using Student’s t-test, Wilcoxon rank-sum test, Chi-square or Fisher’s exact tests to compare exposed and unexposed cohorts as appropriate. Abbreviations: ALC = absolute lymphocyte count, ALT = alanine transaminase, AST = aspartate aminotransferase, BMI = body mass index, CCP = COVID-19 convalescent plasma, ECMO = extracorporeal membrane oxygenation, NT-proBNP = N-terminal-pro b-type natriuretic peptide.

* *, **, and *** correspond to significant p-values for comparison between groups of <0.05, <0.01, and <0.001 respectively.

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|                      | CCP Early | Unexposed Early | Difference or HR (95% CI) | p-value | CCP Late | Unexposed Late | Difference or HR (95% CI) | p-value |
|----------------------|-----------|-----------------|---------------------------|---------|----------|----------------|---------------------------|---------|
| **Primary outcomes** |           |                 |                           |         |          |                |                           |         |
| 14-day mortality     | 15%       | 23%             | 0.52 (0.28, 0.96)         | 0.036*  | 28%      | 29%            | 0.98 (0.53, 1.83)         | 0.95    |
| 30-day mortality     | 38%       | 49%             |                           |         | 42%      | 47%            |                           |         |
| **Secondary outcomes** |         |                 |                           |         |          |                |                           |         |
| Alive and mechanical ventilation-free, days (SD) | 21.4 (11.6) | 18.1 (12.6) | 3.3 (0.2, 6.3) | 0.04*  | 13.4 (12.3) | 16.9 (12.8) | -3.5 (-8.6, 1.7) | 0.18    |
| Length of hospitalization, days (95% CI) | 18.4 (16.3, 20.7) | 17.5 (14.9, 20.1) | 1.0 (-2.4, 4.4) | 0.57    | 24.7 (22.6, 26.8) | 19.9 (16.9, 22.9) | 4.9 (1.2, 8.5) | 0.01*   |
| WHO score on index date, mean (SD) | 5.2 (0.7) | 4.9 (0.9) | 0.3 (0.2,0.5) | 0.0002*** | 5.7 (0.5) | 5.3 (0.9) | 0.4 (0.1, 0.6) | 0.0001*** |
| WHO score change at 7-days, mean (SD) | -0.5 (1.1) | 0.3 (1.6) | -0.8 (-1.2, -0.4) | 0.0003*** | 0 (1.3) | -0.2 (1.6) | 0.2 (-0.3, 0.8) | 0.41    |
| Change of WHO score at discharge, mean (SD) | -0.7 (2.0) | 0.2 (2.2) | -0.9 (-1.5, -0.3) | 0.004** | -0.1 (2.3) | -0.4 (2.1) | 0.6 (-0.4, 1.5) | 0.23    |

Table reports primary and secondary outcomes for the two separate propensity score matched cohorts of early and late transfused patients. CCP effect on mortality is reported as a hazard ratio and secondary outcomes report differences in days or WHO score, each with 95% confidence intervals and p-values shown. Differences in alive and mechanical ventilation free days are measured at 30-days post index. Length of hospitalization is measured from index day.

* *, **, and *** correspond to significant p-values for comparison between groups of <0.05, <0.01, and <0.001 respectively.

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In the propensity score matched analysis, the estimated in-hospital mortality was 15% by day 14 among early CCP recipients, compared with 23% in their matched unexposed cohort. At day 30, estimated in-hospital mortality was 38% in early CCP recipients compared to 49% in their matched unexposed cohort (Table 2). Overall, early CCP recipients were nearly 50% less likely to die in the hospital compared to patients in the matched unexposed cohort (HR 0.52, [95% CI 0.28 to 0.96]; P = 0.036) (Fig 2A). In contrast, there were no differences in mortality
among late CCP recipients compared to their matched unexposed cohort (HR 0.98, [95% CI 0.53 to 1.83]; P = 0.95) (Fig 2B, Table 2). In the late cohort, the estimates of 14-day and 30-day in-hospital mortality were 28% vs 29% and 42% vs 47%, comparing CCP recipients vs matched unexposed patients, respectively.

Secondary endpoints
The early CCP cohort had more days “alive and ventilator free” by 30 days post-index date compared with their matched unexposed cohort (mean 3.3 days, [95% CI 0.2 to 6.3]; P = 0.04, Table 2). There were no differences comparing CCP recipients and matched unexposed patients in the late cohort (mean -3.5 days, [95% CI -8.6 to 1.7]; P = 0.18).

In the early CCP cohort, WHO scores were significantly lower at 7 days post administration or post index date (difference in means -0.8, [95% CI -1.2 to -0.4]; P = 0.0003) and at hospital discharge (difference in means -0.9, [95% CI -1.5 to -0.3]; P = 0.004) compared to their matched unexposed cohort. Early CCP recipients were nearly twice as likely to demonstrate an improvement in their WHO scores at discharge from baseline index day scores (OR 1.9, [95% CI 1.1 to 3.3]; P = 0.02, Fig 3A, Table 2). 42.9% of CCP recipients on mechanical ventilation at baseline (WHO score of 6) no longer required supplementary oxygen at the time of discharge (WHO score of 3) compared to 36.4% of unexposed matches (S1 Fig in S1 File). There were no differences in length of hospital stay post treatment or index date (mean, 18.4 days vs 17.5 days, P = 0.57) or likelihood of discharge (HR for discharge 0.9, 95% CI 0.6 to 1.3; P = 0.46).

In the late cohort, there were no differences in WHO scores at day 7 (mean +0.2, [95% CI -0.3 to 0.8]; P = 0.41) or at discharge (mean +0.6, [95% CI -0.4 to 1.5]; P = 0.23) between CCP recipients and their matched unexposed cohort (Fig 3B, Table 2). The change from baseline (CCP transfusion or index date) to discharge in WHO scores was not different either (OR 0.6 [95% CI 0.3 to 1.3]; P = 0.22). Hospital length of stay was longer in CCP recipients compared with unexposed matches, (mean, 24.7 days vs 19.9 days, P = 0.01), although the likelihood of discharge was not different (HR for discharge 0.6, 95% CI 0.3 to 1.1) (Table 2).

Adverse outcomes
No immediate transfusion-related adverse events were observed after CCP administration in any of the cohorts and no transfusion reactions were reported to the blood bank.

Discussion
In this cohort study of CCP transfusion in patients with moderate-to-severe COVID-19, we found a nearly 50% lower in-hospital mortality, fewer days spent on mechanical ventilation, and greater improvement in WHO scores after early administration of CCP (within 6 days of hospitalization) when compared to a propensity score-matched unexposed cohort. When administered after 6 days of hospitalization, CCP transfusion was not associated with improved mortality or clinical outcomes.

On August 23, 2020, the FDA granted Emergency Use Authorization for the use of CCP in the management of COVID-19 based on preliminary analysis of the EAP which found that CCP administered within 3 days of admission produced clinical benefit. Specifically, the observational study reported by Joyner et al. found lower 30-day mortality in patients who received plasma within 72 hours of first positive SARS-CoV-2 PCR compared to patients transfused after 72 hours [14]. There is a growing body of literature supporting a biological rationale for the early administration of CCP as a means to enhance response to this treatment [15, 21–24]. Active viral shedding is highest within the first week of symptoms, with peak replication occurring within the first 5 days [21, 22]. Moreover, antibody responses to SARS-CoV-2 start to
develop in the first week, with antiviral immunoglobulin peaks by day 22 of symptoms [23].

With the SARS epidemic of 2003, Cheng et al. described higher hospital discharge and lower mortality in patients who received anti-SARS convalescent plasma prior to 14 days after the onset of illness [9]. This is also consistent with data observed by Libster and colleagues, who found a 48% relative risk reduction in developing severe respiratory disease in older adults receiving CCP within 72 hours after the onset of mild COVID-19 symptoms [19]. However, this aforementioned study failed to detect any benefit in mortality due to very few deaths.

Another small study using propensity score matched method reported benefit for patients with severe COVID receiving CCP within 7 days with a HR of 0.34 [15]. The results of our investigation support these findings utilizing a larger sample size and suggest that the administration of CCP after the first week of hospitalization does not attain the same results as earlier CCP administration.

In contrast, there have been multiple randomized controlled trials of CCP with negative findings, the most recent being the CCP arm of RECOVERY, which found no benefit in mortality, mechanical ventilator liberation or hospital discharge [25]. An important distinction from the present study is that the median time to CCP in RECOVERY was 9 days from symptoms. Zeng and colleagues reported that administration of CCP 21 days after initial symptom onset was not associated with mortality benefit [21].

**Fig 2.** Kaplan-Meier curves of survival in early (A) and late (B) convalescent plasma recipients vs respective matched unexposed patients. Shown are Kaplan-Meier estimates of survival from time of index in (A) early CCP recipients (solid line) vs matched unexposed patients (dashed line) (B) late CCP recipients (solid line) vs matched unexposed patients (dashed line). Survival improved for early CCP at 14-days (15% vs 23%) and 30-days (38% vs 49%) compared to matched unexposed patients (HR 0.52, [95% CI 0.28-0.96]; p = 0.0367). There was no difference in mortality at 14-days (28% vs 29%) or 30-days (42% vs 47%) among late CCP recipients compared to their matched unexposed patients (HR 0.98,[95% CI 0.53to 1.83]; p = 0.95). Censoring is indicated by the tick mark “+” with number by each ten-day interval marked below the number at risk.

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**Fig 3.** Histogram plots of WHO ordinal scale scores before and after intervention in the early (A) and late CCP (B) compared to matched unexposed cohorts. Shown is a histogram of the change is WHO COVID-19 severity score from time of index to discharge in (A) early CCP recipients (gray solid bars) vs matched unexposed patients (black solid bars) (B) late CCP recipients (gray dashed bars) vs matched unexposed patients (black solid bars). WHO scores in (A) early CCP recipients were significantly improved at discharge (difference in mean -0.9, [95% CI -1.5 to -0.3]; p = 0.004) when compared to unexposed patients. Relative to the unexposed patients, early CCP recipients were nearly twice as likely to demonstrate an improvement in their WHO scores at discharge from baseline index day scores (OR 1.9,[95% CI 1.1 to 3.3]; p = 0.02). Among (B) late CCP recipients, there was no effect of CCP on WHO scores at discharge (difference in mean +0.6, [95% CI -0.4 to 1.5]; p = 0.23). The odds ratio for improvement at discharge was 0.6 (95% CI 0.3 to 1.3; p = 0.22).

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SARS-CoV-2 detection failed to improve survival despite successful viral clearance [26]. Simo-
novich et al. also reported no benefit to CCP in patients who had a median time from symp-
toms onset to CCP transfusion of 8 days in a recent study [27]. Indeed, our findings in the late
cohort are consistent with this prior work and further emphasize the importance of early
administration of CCP to achieve clinical improvement.

It is important to note that 40% of early CCP recipients in our cohort were on mechanical
ventilation, suggesting that the clinical benefits of early CCP administration extend to even
patients who are critically ill. Although previous uncontrolled case series reported improve-
ment in mechanically ventilated patients who received CCP [28, 29], an open-label, random-
ized trial by Li et al. failed to demonstrate any significant clinical improvement in a small
subset of patients with life-threatening COVID-19 [16], although fewer patients in the CCP
group died compared to unexposed patients [16]. To our knowledge, our study is among the
first to report significant clinical benefit with early administration of CCP in hospitalized
patients with moderate-to-severe COVID-19.

Interestingly, in our study late CCP recipients had longer lengths of hospitalization than
their matched unexposed counterparts despite having similar likelihood of discharge, mortal-
ity, and WHO scores at day 7 and discharge. Future studies should help elucidate potential
explanations for these observations.

Limitations
The interpretation of our findings is limited by the observational nature of the study. However,
propensity score matching to parse electronic medical record data is a valid and commonplace
method in medical research [20, 30]. Matching characteristics were comparable between
exposed and unexposed patients. We acknowledge that it is possible that despite the adequate
matching, patients in the late cohort were intrinsically different from matched unexposed
patients, and indication bias might have played a stronger role, or the late administration of
CCP might not be an adequate rescue intervention. While results from multi-center, random-
ized, controlled trials will be available in the future, our study is the first adequately powered to
evaluate CCP for use in moderate-to-severe disease.

Although other studies may use different cut-offs for timing of CCP administration, our
study used hospitalization date as a reliable time point that could be easily implemented to
guide hospital treatment algorithms. It would be more challenging to establish timing of treat-
ment based on the date of symptom onset, because initial COVID-related symptoms are often
more difficult to precisely identify, and even more so in the early stages of the pandemic.
Symptom onset information was also not feasible to obtain for our entire 2,551 patient cohort
used for propensity matching. Another limitation of the present study is that each pool of
plasma was unique without individual titer levels available, given that data for this study were
collected prior to a validated serologic test. This information on donor plasma would have
enabled us to confirm a dose-response relationship, supporting the concept of the biological
effects of CCP.

Conclusion
Among patients hospitalized with moderate-to-severe COVID-19, administration of CCP
within 6 days of hospitalization in conjunction with standard of care treatment was associated
with improved in-hospital mortality compared to standard of care treatment alone. Further-
more, the early administration of CCP was associated with increased 30-day alive and ventila-
tor free days and improved WHO ordinal scale scores compared with the matched unexposed
cohort. However, these associations were not observed among patients transfused 7 days or
later into their hospital course. Further clinical trials are required to confirm the efficacy and safety of CCP, especially considering the importance of timing of administration, in order to address the limitations of this propensity-matched analysis and confirm our findings.

Supporting information
S1 File. Supplementary figures and tables.

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References

1. Zhou P, Yang XL, Wang XG, Hu B, Zhang L, Zhang W, et al. A pneumonia outbreak associated with a new coronavirus of probable bat origin. Nature. 2020; 579(7798):270–3. Epub 2020/02/06. https://doi.org/10.1038/s41586-020-2012-7 PMID: 32015507; PubMed Central PMCID: PMC7095418.

2. Wu F, Zhao S, Yu B, Chen YM, Wang W, Song ZG, et al. A new coronavirus associated with human respiratory disease in China. Nature. 2020; 579(7798):265–9. Epub 2020/02/06. https://doi.org/10.1038/s41586-020-2008-3 PMID: 32015508; PubMed Central PMCID: PMC7094943.

3. Dong E, Du H, Gardner L. An interactive web-based dashboard to track COVID-19 in real time. The Lancet Infectious Diseases. 2020; 20(5):533–4. https://doi.org/10.1016/S1473-3099(20)30120-1 PMID: 32087114.

4. Group RC, Horby P, Lim WS, Emberson JR, Mafham M, Bell JL, et al. Dexamethasone in Hospitalised Patients with Covid-19—Preliminary Report. N Engl J Med. 2020. Epub 2020/07/18. https://doi.org/10.1056/NEJMoa2021436 PMID: 32678530; PubMed Central PMCID: PMC7383959.

5. Group WHOREAfC-TW, Sterne JAC, Murthy S, Diaz JV, Slutsky AS, Villar J, et al. Association Between Administration of Systemic Corticosteroids and Mortality Among Critically Ill Patients With COVID-19: A Meta-analysis. JAMA. 2020; 324(13):1330–41. Epub 2020/09/03. https://doi.org/10.1001/jama.2020.17023 PMID: 32876694; PubMed Central PMCID: PMC7489434.

6. Beigel JH, Tomashek KM, Dodd LE, Mehta AK, Zingman BS, Kalil AC, et al. Remdesivir for the Treatment of Covid-19—Final Report. N Engl J Med. 2020. Epub 2020/05/24. https://doi.org/10.1056/NEJMoa2007764 PMID: 32445440; PubMed Central PMCID: PMC7262788.

7. Pan H, Peto R, Karim QA, Alejandria M, Henao-Restrepo AM, Garcia CH, et al. Repurposed antiviral drugs for COVID-19—interim WHO SOLIDARITY trial results. medRxiv. 2020.2020.10.15.20209817. https://doi.org/10.1009/JEM.202021364 PMID: 33264556.

8. Rojas M, Rodríguez Y, Monsalve DM, Acosta-Ampudia Y, Camacho B, Gallo JE, et al. Convalescent Plasma in Covid-19: Possible mechanisms of action. Autoimmunity Reviews. 2020; 19(7):102554. https://doi.org/10.1016/j.autrev.2020.102554 PMID: 32380316.

9. Cheng Y, Wong R, Soo YO, Wong WS, Lee CK, Ng MH, et al. Use of convalescent plasma therapy in SARS patients in Hong Kong. Eur J Clin Microbiol Infect Dis. 2005; 24(1):44–6. Epub 2004/12/24. https://doi.org/10.1007/s10096-004-1271-9 PMID: 15916939; PubMed Central PMCID: PMC7088355.

10. Hung IFN, To KK, Lee CK, Lee KL, Yan WW, Chan K, et al. Hyperimmune IV immunoglobulin treatment: a multicenter double-blind randomized controlled trial for patients with severe 2009 influenza A (H1N1) infection. Chest. 2013; 144(2):464–73. Epub 2013/03/02. https://doi.org/10.1378/chest.12-2907 PMID: 23450336.

11. Duan K, Liu B, Li C, Zhang H, Yu T, Qu J, et al. Effectiveness of convalescent plasma therapy in severe COVID-19 patients. Proc Natl Acad Sci U S A. 2020; 117(17):9490–6. Epub 2020/04/08. https://doi.org/10.1073/pnas.2004168117 PMID: 32253318; PubMed Central PMCID: PMC7196337.

12. Salazar E, Perez KK, Ashraf M, Chen J, Castillo B, Christensen PA, et al. Treatment of Coronavirus Disease 2019 (COVID-19) Patients with Convalescent Plasma. Am J Pathol. 2020; 190(6):1680–90. Epub
13. Perotti C, Baldanti F, Bruno R, Del Fante C, Seminari E, Casari S, et al. Mortality reduction in 46 severe Covid-19 patients treated with hyperimmune plasma. A proof of concept single arm multicenter trial. Haematologica. 2020. Epub 2020/07/25. https://doi.org/10.3324/haematol.2020.261784 PMID: 32356382.

14. Joyner MJ, Senefeld JW, Klassen SA, Mills JR, Johnson PW, Theel ES, et al. Effect of Convalescent Plasma on Mortality among Hospitalized Patients with COVID-19: Initial Three-Month Experience. medRxiv. 2020:2020.08.12.20169359. https://doi.org/10.1101/2020.08.12.20169359 PMID: 32817978.

15. Liu STH, Lin HM, Baine I, Wajnb erg A, Gumprecht JP, Rahman F, et al. Convalescent plasma treatment of severe COVID-19: a propensity score-matched control study. Nat Med. 2020. Epub 2020/09/17. https://doi.org/10.1038/s41591-020-1088-9 PMID: 32934372.

16. Li L, Zhang W, Hu Y, Tong X, Zheng S, Yang J, 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(5):460–70. Epub 2020/06/04. https://doi.org/10.1001/jama.2020.10044 PMID: 32942084; PubMed Central PMC: PMC7270883.

17. Arvind Gharbar an CCEJ, Corine Geurtsvan Kessel, den Hollande r Jan G., Karim Faiz, Mollema Femke P.N., Stalenhoef Janneke E., Dofferhoff Anton, Ludwig Inge, Koster Ad, Hassing Robert-Jan, Bos Jeannet C., van Pottelberge Geert R., Vlaaijer Immo N., Ammerlaan Heidi S.M., Segarcanu Elena, Miedema Jelle, van der Eerden Menno, Papageorgiou Grigoris, te Broekhorst Peter, Swan eveld Francis H., Katsikis Peter D., Mueller Yvonne, Ofda Nisreen M.A., Koopmans Marion P.G., Haag mans Bart L., Rokx Casper, Rijnders Bart. Convalescent Plasma for COVID-19. A randomized clinical trial. medRxiv. 2020;Preprint. https://doi.org/10.1101/2020.07.01.20139857.

18. Aganwal A, Mukherjee A, Kumar G, Chatterjee P, Bhatnagar T, Malhotra P. 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; 371:m3939. https://doi.org/10.1136/bmj.m3939 PMID: 33093056.

19. Libster R, Pérez Marc G, Wappner D, Coviello S, Bianchi A, Braem V, et al. Early High-Titer Plasma Therapy to Prevent Severe Covid-19 in Older Adults. N Engl J Med. 2021. Epub 2021/01/07. https://doi.org/10.1056/NEJMoa2033700 PMID: 33406353; PubMed Central PMC: PMC7793608.

20. Austin PC. Propensity-score matching in the cardiovascular surgery literature from 2004 to 2006: a systematic review and suggestions for improvement. J Thorac Cardiovasc Surg. 2007; 134(5):1128–35. Epub 2007/11/03. https://doi.org/10.1016/j.jtcvs.2007.07.021 PMID: 17976439.

21. Wölfel R, Corman VM, Guggemos W, Seilmaier M, Zange S, Müller MA, et al. Virological assessment of hospitalized patients with COVID-2019. Nature. 2020; 581(7809):465–9. https://doi.org/10.1038/s41586-020-2196-x PMID: 32235945.

22. Cevik M, Kuppalli K, Kindrachuk J, Peiris M. Virology, transmission, and pathogenesis of SARS-CoV-2. BMJ. 2020; 371:m3862. https://doi.org/10.1136/bmj.m3862 PMID: 33097561.

23. Long Q-X, Liu B-Z, Deng H-J, Wu G-C, Deng K, Chen Y-K, et al. Antibody responses to SARS-CoV-2 in patients with COVID-19. Nature Medicine. 2020; 26(6):845–8. https://doi.org/10.1038/s41591-020-0897-1 PMID: 32350462.

24. Group Fl-C-. Prevention of severe COVID-19 in the elderly by early high-titer plasma. medRxiv. 2020. https://doi.org/10.1101/2020.11.20.2034013.

25. Group RC. Convalescent plasma in patients admitted to hospital with COVID-19 (RECOVERY): a randomised controlled, open-label, platform trial. Lancet. 2021. Epub 2021/05/18. https://doi.org/10.1016/S0140-6736(21)00897-7 PMID: 34002557.

26. Zeng QL, Yu ZZ, Gou JJ, Li GM, Ma SH, Zhang GF, et al. Effect of Convalescent Plasma Therapy on Viral Shedding and Survival in Patients With Coronavirus Disease 2019. J Infect Dis. 2020; 222(1):38–43. Epub 2020/04/30. https://doi.org/10.1093/infdis/jiaa228 PMID: 32348485; PubMed Central PMC: PMC7197534.

27. Simonovich VA, Burgos Pratx LD, Scibona P, Beruto MV, Vallone MG, Vazquez C, et al. A Randomized Trial of Convalescent Plasma in Covid-19 Severe Pneumonia. N Engl J Med. 2020. Epub 2020/11/25. https://doi.org/10.1056/NEJMoa2031304 PMID: 33232588.

28. Shen C, Wang Z, Zhao F, Yang Y, Li J, Yuan J, et al. Treatment of 5 Critically Ill Patients With COVID-19 With Convalescent Plasma. JAMA. 2020; 323(16):1582–9. Epub 2020/03/29. https://doi.org/10.1001/jama.2020.4783; PubMed Central PMC: PMC7101507.

29. Zhang B, Liu S, Tan T, Huang W, Dong Y, Chen L, et al. Treatment With Convalescent Plasma for Critically Ill Patients With Severe Acute Respiratory Syndrome Coronavirus 2 Infection. Chest. 2020; 158 (1):e9–e13. Epub 2020/04/04. https://doi.org/10.1016/j.chest.2020.03.039 PMID: 32243945; PubMed Central PMC: PMC7195335.
30. Nahid P, Mase SR, Migliori GB, Sotgiu G, Bothamley GH, Brozek JL, et al. Treatment of Drug-Resistant Tuberculosis. An Official ATS/CDC/ERS/IDSA Clinical Practice Guideline. Am J Respir Crit Care Med. 2019; 200(10):e93–e142. Epub 2019/11/16. https://doi.org/10.1164/rcrm.201909-1874ST PMID: 31729908; PubMed Central PMCID: PMC6857485.