Feasibility of a pilot program for COVID-19 convalescent plasma collection in Wuhan, China

Ling Li,1,2,† Ru Yang,3,† Jue Wang,1,2,† Qilu Lu,1,2,† Ming Ren,3 Lei Zhao,3 Hanwei Chen,3 Haixia Xu,1,2 Songli Xie,3 Jin Xie,3 Hui Lin,3 Wenjuan Li,1,2,4 Peng Fang,1,2,4 Li Gong,1,2,4 Lan Wang,3 Yanyun Wu,5 and Zhong Liu†1,2

BACKGROUND: A novel coronavirus has caused an international outbreak. Currently, there are no specific therapeutic agents for coronavirus infections. Convalescent plasma (CP) therapy is a potentially effective treatment option.

METHODS: Patients who had recovered from COVID-19 and had been discharged from the hospital for more than 2 weeks were recruited. COVID-19 convalescent plasma (CCP)-specific donor screening and selection were performed based on the following criteria: 1) aged 18-55 years; 2) eligible for blood donation; 3) diagnosed with COVID-19; 4) had two consecutive negative COVID-19 nasopharyngeal swab tests based on PCR (at least 24 hr apart) prior to hospital discharge; 5) had been discharged from the hospital for more than 2 weeks; and 6) had no COVID-19 symptoms prior to convalescent plasma donation. In addition, preference was given to CCP donors who had a fever lasting more than 3 days or a body temperature exceeding 38.5°C (101.3°F), and who donated 4 weeks after the onset of symptoms. CCP collection was performed using routine plasma collection procedures via plasmapheresis. In addition to routine donor testing, the CCP donors' plasma was also tested for SARS-CoV-2 nucleic acid and S-RBD-specific IgG antibody.

RESULTS: Of the 81 potential CCP donors, 64 (79%) plasma products were collected. There were 18 female donors and 46 male donors. There were 34 first-time blood donors and 30 repeat donors. The average time between CCP collection and initial symptom onset was 49.1 days, and the average time between CCP collection and hospital discharge was 38.7 days. The average volume of CCP collected was 327.7 mL. All Alanine transaminase (ALT) testing results met blood donation requirements. HIV Ag/Ab, anti-HCV, anti-syphilis, and HBsAg were all negative; NAT for HIV, HBV, and HCV were also negative. In addition, all of the CCP donors' plasma units were negative for SARS-CoV-2 RNA. Of the total 64 CCP donors tested, only one had an S-RBD-specific IgG titer of 1:160, all others had a titer of ≥1:320.

CONCLUSION: Based on a feasibility study of a pilot CCP program in Wuhan, China, we demonstrated the success and feasibility of CCP collection. In addition, all of the CCP units collected had a titer of ≥1:160 for S-RBD-specific IgG antibody, which met the CCP quality control requirements based on the Chinese national guidelines for CCP.

In December 2019, a new type of human coronavirus, Severe Acute Respiratory Syndrome Type 2 Coronavirus (SARS-CoV-2), was discovered. The infection caused by the virus, named COVID-19 (coronavirus disease 2019), has been spreading rapidly worldwide. The World Health Organization has declared the outbreak of COVID-19 to be a pandemic. According to the WHO data, as of April 1, 2020, COVID-19 had resulted in a total of 823,626 confirmed cases and had killed 40,598 people globally.1 Most patients with COVID-19 infection experience a series of clinical manifestations such as fever, cough, myalgia or fatigue, dyspnea, and even acute respiratory distress syndrome (ARDS) and secondary infections. Many critically ill patients have been admitted to intensive care units.2,3 Existing reports have shown that the mortality rate ranges from 1% to 4%.4 The severity and epidemic potential of COVID-19 has paralyzed the world’s health care system,

From the 1Clinical Transfusion Research Center, Institute of Blood Transfusion, Chinese Academy of Medical Sciences and Peking Union Medical College; the 2Key Laboratory of Transfusion Adverse Reactions, CAMS, Chengdu, Sichuan; the 3Wuhan Blood Center, Wuhan, Hubei; the 4Anhui Medical University, Hefei, China; and the 5University of Miami, Miami, Florida.

Address reprint requests to: Lan Wang, Baofeng 1st Road, Qiaoakou District, Wuhan, Hubei Province, China; e-mail: wlan66@126.com; Yanyun Wu, University of Miami, Department of Pathology, Miami, FL 33136; e-mail: yxw1366@med.miami.edu; Zhong Liu, 26 Huacai Rd, Longtian Industry Zone, Chenghua District, Chengdu, Sichuan, China; e-mail: liuz@ibt.pumc.edu.cn.

†Equal contributors.

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claimed many lives, and threatened economic stability. Unfortunately, to date, apart from symptomatic treatment and supportive care, no specific antiviral treatment or vaccine has been proven effective.\(^5\)

Convalescent plasma (CP) containing SARS-CoV-2-specific antibodies from recovered patients is now being entertained as a potential treatment option.\(^6\) CP has been used to treat several other viral infections, including severe acute respiratory syndrome coronavirus (SARS-CoV),\(^7\) Ebola virus,\(^8\) Middle East respiratory syndrome coronavirus (MERS-CoV),\(^9\) and avian influenza A(H5N1) virus.\(^10\) A recent report by Shen et al. showed that the transfusion of COVID-19 convalescent plasma (CCP) had resulted in significant clinical improvement in five critically ill patients with COVID-19 and ARDS.\(^11\) CCP has been included as a treatment option in the Chinese COVID-19 treatment guidelines and a viral titer of 1:160 has been recommended as a product quality control indicator.\(^12\) Recently, the US Food and Drug Administration created pathways for using CCP either under an emergency IND or expanded access.\(^13\)

For the past few months, CCP has been collected and used in China empirically. In this study, we highlight key elements of a pilot program for collecting CCP in Wuhan, China.

**METHODS**

**Operation protocol for CCP collection**

Figure 1 illustrates the workflow of the CCP collection program in Wuhan, China. Potential donors were recruited through mass media and social media such as WeChat. Please see Figure S2 for sample recruitment material in Chinese, with English translation in supplemental material.

The primary recruitment targets were patients who had recovered from COVID-19 and had been discharged from the hospital for more than 2 weeks. Hospital discharge criteria include: 1) body temperature returns to normal for more than 7 days; 2) respiratory symptoms improve significantly; 3) lung imaging shows significant absorption of inflammation; and 4) two consecutive negative COVID-19 nasopharyngeal swab tests based on PCR (at least 24 hr apart). Consent was obtained via telephone by qualified and trained staff to review a COVID-19-specific health history questionnaire and to qualify them based on the following criteria: 1) aged 18-55 years; 2) eligible for blood donation; 3) diagnosed with COVID-19; 4) had two consecutive negative COVID-19 nasopharyngeal swab tests based on PCR (at least 24 hr apart) prior to hospital discharge; 5) had been discharged from the hospital for more than 2 weeks; and 6) had no COVID-19 symptoms prior to CCP donation. In addition, preference was given to CCP donors who had a fever lasting more than 3 days or a body temperature exceeding 38.5°C (101.3°F), and 4 weeks after symptom onset.\(^14\) If they were deemed qualified at the pre-screening, they were asked to bring their medical records to an in-person appointment at the Wuhan Blood Center. A healthcare worker from the blood center examined the donors, reviewed their medical records, and made sure that the donors had two negative COVID-19 nasopharyngeal swab tests based on PCR (at least 24 hr apart) and no COVID-19 symptoms. If they were determined to be eligible for CCP donation, they then underwent pre-donation testing that included alanine transaminase (ALT), HBsAg, and hemoglobin. If they met all blood donation requirements, they would proceed with a CCP donation via plasmapheresis that was performed according to routine procedures and requirements for plasma donation. After plasma donation, they were monitored for adverse events and awarded with a CCP-specific blood donation certificate.

**Study recruitment, screening, and sample collection**

Donor recruitment, consent, screening, and blood specimen collection were conducted at the Wuhan Blood Center. Protocols for donor selection, plasma collection, specimen
collection and testing, and clinical trials were approved by the Ethics Review Committee of the Institute of Blood Transfusion, Chinese Academy of Medical Sciences. All CCP donors gave written, informed consent. The CCP products collected from March 3, 2020 to March 18, 2020 were examined for the CCP program and product quality control.

**Plasma collection**
CCP collection was performed according to routine procedures and requirements for plasma donation. Plasma collection was performed via plasmapheresis (DigiPla80, Sichuan Nangeer Biotechnology Co., Ltd; XCF 3000, Sichuan Nangeer Biotechnology Co., Ltd). The volume of plasma to be collected was determined based on the donor’s weight. Female donors and male donors weighing less than 70 kg were given 0.9% sodium chloride (NS) during collection. The volume of the replacement fluid was half of the plasma collected. All plasma products were frozen within 6 hours.

**Laboratory testing**

**Sample collection and preparation**
All donor samples were collected in a sodium citrate tube for the ELISA assay and an Ethylene Diamine Tetraacetic Acid (EDTA) tube with gel separator for the NAT assay, and were centrifuged and separated within 4 hours.

**Routine blood product testing**
All blood products were tested for HCV, HBV, HIV by ELISA and NAT, and for syphilis by ELISA.

**Special testing for CCP**

**Testing for SARS-CoV-2 nucleic acid.** All collected donor plasma samples were screened for SARS-CoV-2 RNA by individual donor testing, using the PerkinElmer New Coronavirus Nucleic Acid Detection Kit (PerkinElmer Healthcare Diagnostics Co., Ltd.). The test kits used were approved and licensed by the FDA, and the assays were performed according to the manufacturers’ instructions.

**Titer of S-RBD-specific IgG antibody.** For product quality control, S-RBD-specific IgG antibody was measured. The S-RBD-specific IgG antibody ELISA kit was made in-house. In brief, 96 well plates (Thermo Scientific) were coated with 100 ng of recombinant Receptor binding domain (RBD) polypeptides per well (Sino Biological). The plates were decanted of the coating solution after overnight coating and were blocked with B3T at 37°C (98.6°F) for 1 hour. Plasma samples were diluted 160-, 320-, 640-, and 1280-fold with 0.5% triton X-100 phosphate buffered saline (TPBS) and 5% fetal calf serum (FBS) (Gibco). After washing, the plates had mouse to human secondary antibody added and were observed for horseradish peroxidase (HRP) reaction. The changes in the absorbance at 450 nm and 630 nm were measured using an automatic microplate reader (Sunrise Tecan GmbH), and the OD values were calculated. The results were reported as the S/CO value, which was calculated as the ratio of OD value to the cutoff value. If the ELISA assay was positive, the highest dilutions were reported as the titers, which ranged between 1:160 and 1:1280 (ELISA endpoint dilution titers).

**Statistical analysis**
The descriptive data were reported. The minimum, maximum, median, average, and SD were calculated by Excel.

**RESULTS**

**CCP donor and product summary**
From 81 potential donors for CCP, 64 (79% of donors) plasma products were collected; 17 of the potential donors could not donate plasma because they did not meet the blood donation requirements (2 had hypertension and 15 had abnormal ALT). There were 18 female donors and 46 male donors. Of the 64 donors, 34 were first-time blood donors and 30 were repeat donors. The average time between CCP collection and symptom onset was 49.1 days, and the average time between CCP collection and hospital discharge was 38.7 days. The average volume of CCP collected was 327.7 mL.

**Infectious disease testing**
All ALT testing met the blood donation criteria. The anti-HIV1/2, anti-HCV, syphilis testing, and HBsAg were all negative; the NAT for HIV, HBV, and HCV were also negative.

In addition, all CCP donors’ plasma tested negative for SARS-CoV-2 RNA.

**Titer of S-RBD-specific IgG antibody**
Among the total 64 analyzed CCP donors, only one had an S-RBD-specific IgG titer of 1:160, all others had a titer ≥1:320.

**DISCUSSION**
The COVID-19 pandemic has become a major public health challenge, not only for China but also for countries around the world. The development of an effective and safe CCP program may provide an effective treatment option to help control the pandemic situation.

Here we report a summary of a pilot CCP collection program (Table 1). The program was able to attract recovered patients and successfully collected CCP products. The high rate of program exclusion is not surprising, because this was not a typical blood donation, and some patients may not have been fully recovered from COVID-19 or were not qualified as blood donors at baseline despite a high level of enthusiasm. It is also very encouraging to note that more than half of the CCP donors are committed repeat
blood donors who were willing to come back after being very ill.

From blood safety perspective for CCP, it is also notable that we did not observe any positivity for routine TTD testing, or NAT for SARS-CoV-2 nucleic acid. This is rather reassuring. From product quality control perspective, it is notable that all CCP donors in this pilot program had an S-RBD-specific IgG titer of $\geq 1:160$, which met the Chinese national CCP standard.

While the data are very limited, we did demonstrate the success and feasibility of a pilot CCP program. In addition, clinical validation based on well-designed clinical trials such as RCT will be needed for the safety and therapeutic efficacy of CCP. Clinical correlation with the donor selection criteria and product quality control indicators will help further refine the CCP program requirements.

**CONCLUSION**

Based on a feasibility study of a pilot CCP program in Wuhan, China, we demonstrated the success and feasibility of CCP collection. All donors who donated CCP during the study period had negative routine TTD, including negative NAT for HIV, HBV, and HCV, and negative NAT for SARS-CoV-2. In addition, all donors had a titer $\geq 1:160$ for S-RBD-specific IgG antibody, which met the CCP quality control requirements based on the Chinese national guidelines for CCP.

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**CONFLICT OF INTEREST**

The authors have disclosed no conflicts of interest.

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SUPPORTING INFORMATION

Additional Supporting Information may be found in the online version of this article.

Supplemental material

Figure S2: Recruitment material in Chinese, with English translation in supplemental material.