Commentary

Cochrane corner: convalescent plasma or hyperimmune immunoglobulin for people with COVID-19

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Cite this article: Pan Africa Medical Journal. 2020;35(2):81. DOI: 10.11604/pamj.supp.2020.35.24157
Received: 08 Jun 2020 - Accepted: 16 Jun 2020 - Published: 16 Jun 2020
Domain: Epidemiology, Haematology, Immunology
Keywords: Coronavirus, COVID-19, convalescent plasma, hyperimmune immunoglobulin, transfusion
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As the novel coronavirus continues to spread globally and across Africa, efforts are being accelerated to identify effective preventive and therapeutic measures to mitigate its burden. Convalescent plasma and hyperimmune immunoglobulin are being considered as potential therapeutic options for the coronavirus disease 2019 (COVID-19). We highlight and contextualize the findings of a recent Cochrane rapid review that evaluated the effectiveness and safety of convalescent plasma therapy in patients with COVID-19. From the eight studies it included, the review found limited and low-certainty evidence on the effectiveness and safety of convalescent plasma therapy in patients with COVID-19. The evidence was limited by the small number of participants and low-quality of included studies, as well as the inconsistency of outcome measures and reporting across studies. As African countries brace for the further spread of the virus, while exploring potential therapeutic options to mitigate its morbidity and mortality at peak, convalescent plasma therapy may offer a therapeutic ray of hope for the continent. Considering the limited evidence of the effectiveness and safety in the treatment of COVID-19, it is imperative for this therapy to be investigated within African contexts to ascertain not only its effectiveness and safety, but also its practical implications within the capacity of national blood transfusion services and health systems in the region.

Commentary

As the novel coronavirus continues to spread globally and across the Africa region, efforts are being accelerated to identify effective preventive and therapeutic measures to mitigate the impact of the coronavirus disease 2019 (COVID-19) [1]. While Africa is currently the least affected region globally, the COVID-19 pandemic continues to put enormous strain on the region’s frail health systems already strained by other disease burdens [2]. With no licensed vaccine or definitive treatment, management of COVID-19 has been limited to supportive care including fluid and electrolyte replacement, respiratory support and management of complications, in addition to several social distancing measures to prevent transmission [3]. Given the magnitude of the current pandemic, there have been growing interests in the investigation of the clinical benefits of convalescent plasma or hyperimmune immunoglobulin transfusion in the treatment of severely or critically ill COVID-19 patients. In this commentary, we highlight and contextualize the findings of a recently published Cochrane rapid review by Valk and colleagues, that evaluated...
the effectiveness and safety of convalescent plasma or hyperimmune immunoglobulin transfusion in the treatment of people with COVID-19 [4].

Convalescent plasma therapy involves the transfusion of blood plasma containing neutralising antibodies, collected from patients who recovered from COVID-19, to patients who are immunocompromised to this infection. Hyperimmune immunoglobulin therapy is similar to convalescent plasma transfusion, but is prepared from the plasma of donors with high titres of neutralizing antibodies against a specific organism or antigen [5]. Historically, both therapies have been used in the past to treat conditions for which there were no vaccines or definitive curative agents available. In this review, they were considered as potential therapeutic options for the current COVID-19 outbreak [3].

In their recently published Cochrane rapid review, Valk and colleagues considered case-series, cohort, prospectively planned, and randomised controlled trials (RCTs) eligible for inclusion, if the study evaluated convalescent plasma or hyperimmune immunoglobulin for treating people with COVID-19, irrespective of disease severity, age, gender or ethnicity. Although theirs was a rapid review, the authors followed the standard Cochrane systematic review methodology and performed all steps of study screening and selection in duplicate by two independent review authors. They searched for eligible studies in multiple electronic databases up to 23 April 2020. The searched databases included the World Health Organization (WHO) COVID-19 Global Research Database, MEDLINE, Embase, Cochrane COVID-19 Study Register and the Centers for Disease Control and Prevention (CDC) COVID-19 Research Article Database. They also searched trials registries to identify on-going trials. In their search strategy, the authors combined terms for the novel coronavirus (such as SARS-CoV-2 and nCoV-2019), the disease (such as COVID-19 and COVID-19) and convalescent plasma or hyperimmune immunoglobulin (such as plasma and immunoglobulin). The literature search yielded a total of 1267 potentially relevant references, from which eight studies (seven case-series, one prospectively planned, single-arm intervention study) with a total of 32 participants were eventually included in the review. Of the eight included studies, seven originated from China, while one was conducted in South Korea. In addition, the authors identified 48 ongoing studies in the battle against this new strain of coronavirus, including five hyperimmune immunoglobulin (one study) in COVID-19 patients, of which 22 are RCTs. The dose, volume and timing of convalescent plasma varied widely between studies. The total volume of convalescent plasma transfused was between 200mL and 2400mL, with participants receiving between one and eight doses of plasma transfusion. Most donors were male, aged between 18 and 60 years. All donors were symptom-free and had completely recovered from COVID-19 prior to plasma donation.

The authors assessed therapeutic effectiveness by improvement of clinical symptoms (assessed by respiratory support), time to discharge from hospital, admission on the intensive care unit (ICU) and length of stay in the ICU and all-cause mortality at hospital discharge. All included studies reported some improvement in clinical symptoms in at least some participants. Time to discharge ranged from 4 to 35 days after convalescent plasma therapy. The majority of patients who were admitted in the intensive care unit (ICU) were no longer on the ICU or no longer required mechanical ventilation at the end of the reporting period. Length of stay on the ICU after receiving plasma (11 days) was reported for only one participant. In terms of safety, the review found that the majority of the included studies reported that no serious adverse events occurred following plasma transfusion; however one study reported a serious adverse event (severe anaphylactic shock). Reported non-serious adverse events included moderate fever (38.9°C). All participants were alive at the end of the reporting period, but not all participants had been discharged from hospital by the end of the study. Overall, these findings represent very limited and low-certainty evidence on the effectiveness and safety of convalescent plasma therapy for people with COVID-19, due to the low-quality of included studies; small number of participants; inconsistency of outcome measures; variation in outcome reporting across studies; and differences in the type of participants with COVID-19. The authors highlighted that more high-quality RCTs are needed to determine the effectiveness and safety of convalescent plasma or hyperimmune immunoglobulin in large, randomized clinical trials or well-designed observational studies to obtain better-certainty evidence on the clinical benefits of these potential therapies for COVID-19.

While Africa currently bears the least burden of the COVID-19 pandemic, the enormous threats posed on the region’s frail health systems by the possibility of further spread of the virus are dire [2]. Therefore, African countries must brace for these threats by exploring potential therapeutic options to mitigate the morbidity and fatality of the pandemic at peak. In spite of its limited effectiveness and safety evidence in the context of COVID-19, convalescent plasma transfusion may offer a therapeutic ray of hope for the continent. Also, the previous successes in the treatment of similar respiratory diseases like the avian influenza A (H5N1), the 2009 pandemic influenza A (H1N1) and severe acute respiratory syndrome (SARS) coronavirus, convalescent plasma may offer some optimism in mitigating the potential benefits of the therapeutic plasma treatment. Of the eight included studies, seven originated from China, while one was conducted in South Korea. In addition, the authors identified 48 ongoing studies in the battle against this new strain of coronavirus, including five hyperimmune immunoglobulin (one study) in COVID-19 patients, of which 22 are RCTs. The dose, volume and timing of convalescent plasma varied widely between studies. The total volume of convalescent plasma transfused was between 200mL and 2400mL, with participants receiving between one and eight doses of plasma transfusion. Most donors were male, aged between 18 and 60 years. All donors were symptom-free and had completely recovered from COVID-19 prior to plasma donation.

Conclusion

From the highlighted Cochrane review, there is limited and low-certainty evidence on the effectiveness and safety of convalescent plasma or hyperimmune immunoglobulin therapy for people with COVID-19, due to small sample sizes and study design limitations of the included studies.
African countries may consider further investigation to ascertain not only the effectiveness and safety but the practicality of these therapeutic options within the capacity of their national blood transfusion services.

**Competing interests**
The authors declare no competing interests.

**Authors’ contributions**
CSW conceived the manuscript. CAN wrote the first draft. CSW contributed important intellectual input to subsequent versions of the manuscript and approved the final version.

**Acknowledgements**
The views expressed in this article are those of the authors and do not necessarily reflect the views or policies of Cochrane or the South African Medical Research Council.

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