

To:	Carter BloodCare Customers
From:	Hospital Relations
Date:	December 30, 2021
Re:	FDA revisions affecting the use of COVID-19 Convalescent Plasma
	Usage

## Background

On December 28, 2021, the FDA revised the emergency use authorization (EUA) associated with COVID-19 Convalescent Plasma (CCP) restricting the use of CCP with high titers of anti-SARS-COV-2 antibodies for the treatment of COVID-19 in patients with immunosuppressive disease or receiving immunosuppressive treatment in the inpatient or outpatient setting; revised the acceptable tests and increased qualifying antibody test result cutoffs to be used for manufacturing CCP, and made corresponding changes to the authorized Fact Sheets.

Effective immediately, Carter BloodCare will discontinue distribution of CCP until we can confirm that any previously collected units meet the new high titer requirements. We will notify you if and when CCP distribution is resumed. Please review the revisions to the EUA <u>here</u>.

## **Client Action**

- Notify appropriate personnel that CCP distribution is discontinued effective immediately.