

Study takes us a step closer to a universal antibody test for COVID-19

September 10 2020



Houston Methodist became the first academic medical center in the nation to infuse critically ill COVID-19 patients with convalescent plasma donated from recovered patients, which contains antibodies made by the immune system that were used to kill the virus and may transfer the power of the antibodies into a healing, possibly life-saving therapy. Credit: Houston Methodist



A new study released by Houston Methodist takes researchers a significant step closer to developing a uniform, universal COVID-19 antibody test. The multicenter collaboration tested alternative ways to measure COVID-19 antibody levels that is faster and easier and can inexpensively be used on a larger scale to accurately identify potential donors with the best chance of helping patients infected with the SARS-CoV-2 virus with convalescent plasma therapy.

The findings will also have applications beyond determining who the best plasma donors are. The consensus among the study authors is that, following donor identification, it will most likely next be used in practice to establish target levels of COVID-19 antibodies individuals will need to be considered candidates for vaccines and passive immune therapies.

Additional uses coming later that are likely to have the biggest societal impact, the researchers say, are to assess relative immunity in those previously infected by the SARS-CoV-2 virus and identifying asymptomatic individuals with high levels of neutralizing antibodies against SARS-CoV-2.

It was also found that donors who experienced shortness of breath (or dyspnea) while infected with COVID-19 and those who were hospitalized or had severe disease were more likely to have a robust immune response and, thus, had higher levels of neutralizing antibodies in all the tests. In the absence of available testing, identifying such donor characteristics may be used as a contingency plan to determine which patients have developed higher antibody levels and inform efforts to recruit plasma donors for therapeutic purposes.

In collaboration with Penn State, University of Texas at Austin and U.S. Army Medical Research Institute of Infectious Diseases, study authors James M. Musser, M.D., Ph.D., and Eric Salazar, M.D., Ph.D., physician scientists at Houston Methodist, sought to find alternatives to measuring



virus neutralization (VN) titers, which is the gold standard of COVID-19 antibody testing, as VN antibodies in the blood correlate with immunity. This kind of antibody testing, however, is not widely available, because it's technically complex, requires days to set up, run and interpret, and needs to be performed in a biosafety level 3 laboratory. This leads to most donor plasma virus antibody levels remaining unknown prior to transfusions, so an easier, more readily available method is needed to identify more suitable convalescent plasma donors.

The research team, therefore, looked to another type of test, called ELISA assays, which can be implemented and performed with relative ease in a high-throughput fashion and are widely available and extensively used in clinical labs across the world. The ELISA tests, or enzyme-linked immunosorbent assays, look at whether antibodies against the SARS-CoV-2 proteins are present and produce a quantitative measure of those antibodies. The UT Austin research team developed the ELISA antibody test for SARS-CoV-2 and provided the viral antigens for this study.

Specifically, scientists looked at the relationship of anti-spike ectodomain (ECD) and anti-receptor binding domain (RBD) IgG bloodstream antibody titers. The spike ECD and RBD proteins are physiological parts of the much-talked-about spike protein made by SARS-CoV-2 and critical to how the virus finds its way into the body, spreads and causes COVID-19 disease, so they are prime targets for antibody testing and vaccine development. The blood samples for the study were identified during an institutional surveillance program involving 2,814 Houston Methodist employees.

The goal of the study was to <u>test</u> the hypothesis that anti-ECD and anti-RBD IgG bloodstream antibody titers are correlated with VN titer, making these more accessible, easier-to-perform ELISA tests a surrogate marker to identify plasma donors with titers above the recommended



U.S. Food and Drug Administration threshold for convalescent plasma donation.

In assessing the correlation between VN antibody levels and anti-RBD and anti-ECD ELISA protein titer data, the researchers found that the ELISA tests had an 80% probability or greater of comparable antibody level to VN titers at or above the FDA-recommended levels for COVID-19 convalescent plasma. These results affirm that all three types of tests could potentially serve as a quantitative target for therapeutic and prophylactic treatments.

They also found that convalescent donors maintain high levels of immunity over the course of many weeks and that frequent plasma donations did not cause a significant decrease in antibody or virus neutralization levels.

Perhaps most surprising is that they also identified 27 individuals from the surveillance cohort with high enough antibody titers across all three tests to indicate that some asymptomatic individuals may have plasma suitable for therapeutic use and may have a degree of relative immunity against SARS-CoV-2.

Ultimately, the study successfully concluded that anti-RBD or anti-ECD IgG antibody titers can serve as a surrogate for VN titers to identify suitable <u>plasma</u> donors and that these alternate ELISA tests may provide critical information about COVID-19 immunity.

More information: Eric Salazar et al. Convalescent plasma anti-SARS-CoV-2 spike protein ectodomain and receptor binding domain IgG correlate with virus neutralization, *Journal of Clinical Investigation* (2020). DOI: 10.1172/JCI141206



Provided by Houston Methodist

Citation: Study takes us a step closer to a universal antibody test for COVID-19 (2020, September 10) retrieved 10 April 2024 from https://medicalxpress.com/news/2020-09-closer-universal-antibody-covid-.html

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