

Convalescent plasma: What you need to know

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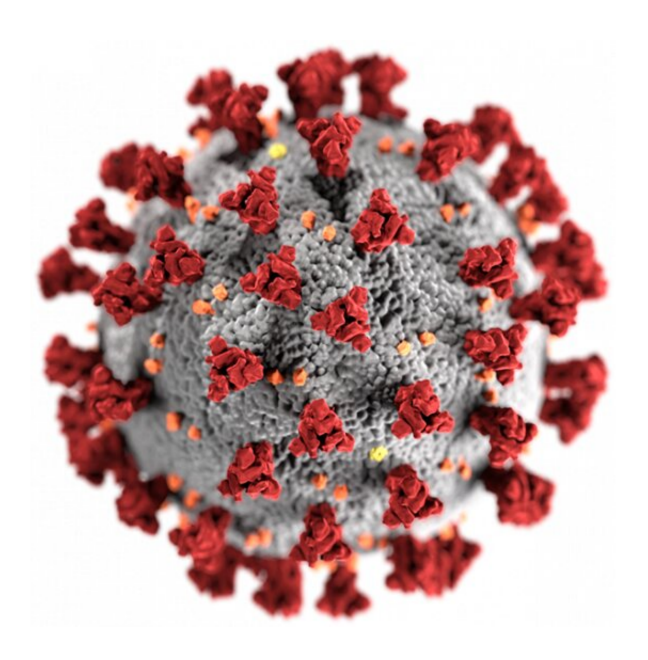


Image of the ultrastructural morphology exhibited by the 2019 Novel Coronavirus (2019-nCoV). Credit: CDC



The United States has issued an emergency authorization to use blood plasma from recovered COVID-19 patients as a treatment against the disease.

Is it safe and effective? Has President Donald Trump's administration put political considerations ahead of the science? Here's what you need to know.

What is convalescent plasma?

When a person has COVID-19, their body produces antibodies that fight the <u>coronavirus</u>. These proteins float in plasma, the liquid component of blood.

The antibodies can be harvested from patients who have recovered and injected into the blood of others to help them fight the same infection.

The idea isn't new—so-called "passive immunization"—was first tried out against diphtheria in 1892 and later against the 1918 flu pandemic.

Is it safe and effective?

The research into these questions is ongoing, but some early signs have been encouraging.

In June, the Mayo Clinic analyzed the safety of plasma following transfusion in a group of 20,000 COVID-19 patients, finding extremely low rates of side-effects such as heart failure, lung injury, allergic reaction and death.

"We concluded that the use of <u>convalescent plasma</u> was safe," Dr. Scott Wright, who led the study that was published in Mayo Clinic



Proceedings, told AFP.

Importantly, there was no sign of an effect called "antibody-dependent enhancement," when antibodies that aren't well suited to stopping a virus actually lead to more cells becoming infected.

On the question of how well it works, all experts agree on the need for more <u>clinical trials</u> to compare plasma against standard care.

Dr. Soumya Swaminathan, chief scientist of the World Health Organization, said a few smaller trials have reported their findings but "the results, in some cases, point to some benefit, but have not been conclusive."

Another study by Mayo Clinic—which was not a clinical trial and hasn't yet been peer-reviewed—suggested plasma helped reduce death rates among hospitalized patients when given early and when the antibody levels were high.

But there was no placebo, making it hard to read too much into the findings.

It enrolled 35,000 patients and showed that those who received transfusions within three days of their COVID-19 diagnosis had an 8.7 percent death rate in the next week.

Those who received plasma after four or more days had an 11.9 percent death rate.

Researchers at Johns Hopkins University are meanwhile running a trial in which they are using plasma to immunize patients before they have a chance to get sick.



Dr. David Sullivan, who is leading this trial, compared it to an "immediate vaccine."

If it works, "we can tell people that if you're high risk... you can get this early, and you don't have to worry about going to the hospital," he told AFP.

Some scientists believe that while plasma might be useful right now, in the long-run it might be more advantageous to identify the best antibodies to COVID-19 then synthesize them in labs.

These are known as "monoclonal antibodies" and they are being developed by biotech firms including Regeneron and Lilly.

The advantages are that doctors know precisely what they are getting and can dose accordingly, and the drugs can be mass-produced.

On the other hand, since the virus is continually mutating, antibodies that were produced in a lab to fight an older version might not be as effective as recently-harvested plasma.

Was the authorization politicized?

The Food and Drug Administration (FDA) has previously issued emergency authorizations for antiviral remdesivir, after it was shown to have moderate efficacy against the coronavirus, and for the antimalarial drug hydroxychloroquine, which was subsequently reversed following safety concerns.

Political commentators have questioned the timing of the FDA's latest move as Trump lags in polls before the November election.

Indeed, in making the announcement, both Trump and FDA



Commissioner Stephen Hahn misrepresented a key statistic when they said that <u>plasma</u> reduced mortality rates by 35 percent.

FDA spokeswoman Emily Miller later clarified on Twitter that the figure referred to the relative reduction in mortality risk for people who received high-levels of <u>antibodies</u> in the Mayo Clinic study compared to those who received low-levels.

"It does undermine the credibility of the entire United States government and the administration but specifically these institutions," said Dr. Matthew Heinz, a physician in Arizona who served in former president Barack Obama's health department.

But Dr. Daniel Hanley, who directs Johns Hopkins' multisite clinical trials, said that early results had met the threshold for emergency approval.

"One thing to emphasize is this information is coming in faster than it has in any other pandemic," he told AFP.

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