Covid-19: 40% of patients with weakened immune system mount lower response to vaccines

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Four in 10 people who are clinically vulnerable generate lower levels of antibodies than healthy recipients after two shots of vaccine against SARS-CoV-2, a study has found.\(^1\)

The Octave (Observational Cohort Trial T cells Antibodies and Vaccine Efficacy in SARS-CoV-2) trial is one of the largest in the world to have looked at the response to covid-19 vaccination in patients who are immunocompromised. It compared 600 patients, who had a weakened immune system because of their disease process or treatment, with the antibody response of healthy people from the Pitch (Protective Immunity from T Cells in Healthcare workers) study.

The trial included patients with solid organ and haematological cancers, end stage kidney and liver disease, organ transplants, and immune mediated inflammatory disease such as inflammatory bowel disease, vasculitis, or rheumatoid arthritis—patients who were not included in original vaccine trial data.

The findings, published as a preprint on the Lancet site,\(^1\) showed that 89% of immunocompromised patients seroconverted within four weeks of the second vaccine dose, as compared with 100% of healthy participants in the Pitch trial.

Overall, 60% of immunocompromised patients had an antibody response equivalent to that of healthy vaccine recipients, but 11% of those with a weakened immune system failed to generate any antibodies.

Failure to seroconvert was particularly high in certain groups: 72.4% of patients with ANCA associated vasculitis and 98% of patients with inflammatory arthritis were in this category. Notably, all of the patients with ANCA associated vasculitis had received rituximab, a targeted B cell depletion therapy, and the researchers suspect a possible link between low seroconversion and rituximab because of the importance of B cells in the immune response to covid-19.

**Immunological analysis**

The findings have been shared with the UK’s Joint Committee on Vaccination and Immunisation, which is poised to decide on provision of booster vaccines in early September. Some countries that are already administering booster doses have prioritised people with chronic diseases or who are taking immunosuppressive therapies.\(^2\)

Iain McInnes, chief investigator of Octave, clarified that this was an immunological analysis, looking at antibody and cellular (T cell) response, not a clinical effectiveness trial examining responses to specific vaccines or mixed doses.

While there was concern about the lack of seroconversion in some people, the researchers noted that these patients had a cellular (T cell) response, the body’s “back-up” system. This suggests that the vaccine is immunologically active in all patients but that the type and quality of that response varies.

The trial, funded by the Medical Research Council, is a multicentre, UK-wide trial, led by the University of Glasgow and coordinated by the University of Birmingham’s Cancer Research UK Clinical Trials Unit.

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