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## Provisional Authorisation for COVID-19 Tests

<i>Applicant</i>	Restalyst Pte Ltd
<i>Name of test</i>	COVID19-REAAD™ Anti-SARS-CoV-2 IgG ELISA
<i>Intended purpose (As per manufacturer's information for use)</i>	<p>COVID19-REAAD™ Anti-SARS-CoV-2 IgG ELISA (Coronavirus Disease 2019– REcombinant Antigen-Antibody Detection) is an enzyme–linked immunosorbent assay intended for qualitative and semi-quantitative detection of SARS-CoV-2 IgG antibodies in human serum and plasma. COVID19-REAAD™ is intended for use as an aid in identifying individuals with an adaptive immune response to SARS-CoV-2, indicating recent or prior infection. COVID19-REAAD™ should not be used as the sole basis for diagnosis.</p> <p>COVID19-REAAD™ is intended for professional use in detecting COVID-19 patients by detecting patient's anti-SARS-CoV-2 antibodies. IgG antibodies to SARS-CoV-2 are generally detectable in blood several days after initial infection, although the post-infection duration of antibodies presence is not well characterised. Individuals may have detectable virus present for several weeks following seroconversion.</p> <p>The sensitivity of COVID19-REAAD™ early after infection is unknown. Negative results do not preclude SARS-CoV-2 infection and should not be used as the sole basis for patient management decisions. If acute infection is suspected, direct testing for SARS-CoV-2 is necessary. False positive results for IgG antibodies may occur due to cross-reactivity from pre-existing antibodies or other possible causes.</p>
<i>Date of Provisional Authorisation</i>	29 June 2021

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