



COVID-19 Therapeutic Alert

CEM/CMO/2021/017

17 September 2021

Casirivimab and imdevimab for patients hospitalised due to COVID-19

Summary

Neutralising monoclonal antibodies (nMABs) bind to specific sites on the spike protein of the SARS-CoV-2 virus particle, blocking its entry into cells and therefore inhibiting its replication. Ronapreve® is a combination nMAB containing equal amounts of casirivimab and imdevimab.

The RECOVERY trial has [demonstrated](#) that the casirivimab and imdevimab combination reduced the relative risk of mortality by 20%, and the absolute risk of mortality by 6%, in hospitalised patients with COVID-19 who had not mounted an antibody response of their own to the virus (i.e. were seronegative¹) at the time of treatment. Mortality was 24% in the casirivimab plus imdevimab treatment group vs 30% in those who received standard care alone. Risk of mortality in hospitalised patients has also been informed by the QCOVID® [analysis](#).

A UK-wide clinical commissioning policy [has now been published](#) recommending consideration of the intravenous use of the combination neutralising antibody casirivimab plus imdevimab at a total dose of 2.4g (1.2g of casirivimab plus 1.2g of imdevimab) in antibody seronegative patients hospitalised for the management of symptoms of COVID 19 infection who are either:

- aged 50 and above,

OR

- aged 12 to 49 AND determined to be immunocompromised by multidisciplinary team (MDT) assessment.

The casirivimab and imdevimab combination is licensed² in Great Britain for the treatment of COVID-19 in individuals aged 12 and above and weighing at least 40 kg but the published policy recommends an off-label use. A temporary regulation 174 approval is in place to cover use in Northern Ireland, pending a licensing decision by the European Medicines Agency (EMA).

Action

¹ Refers to patients who were negative for serum anti-s spike antibodies against SARS-CoV-2

² The conditional marketing authorisation covers use in Great Britain. A parallel regulation 174 approval covers use in Northern Ireland, ahead of the European Medicines Agency's determination.

NHS acute trusts / health boards are asked to take the following immediate steps to support the treatment of patients admitted to hospital for the management of symptoms of COVID-19 infection:

1. **Organisations are recommended to consider prescribing the casirivimab and imdevimab antibody combination [in line with the published policy](#) to PCR³ positive, antibody seronegative patients who have been hospitalised specifically to manage symptoms of COVID-19 infection and are either aged 50 and over, or aged between 12 and 49 and determined to be immunocompromised by multi-disciplinary team (MDT) assessment.** In the absence of a confirmed virological diagnosis, the treatment should only be used when a multidisciplinary team has a high level of confidence that the clinical and radiological features suggest that COVID-19 is the most likely diagnosis.
2. As nMAB therapies should be given to eligible patients as early as possible to maximise benefit, **organisations should ensure that anti-s spike antibody testing is undertaken for all patients hospitalised with COVID at, or as soon as possible after, the point of admission.** Supporting laboratory networks should ensure that the maximum turnaround time for IgG anti-s antibody tests should be no greater than 24 hours from the sample being taken to the result being returned. Positive and negative tests should be reported via the Second Generation Surveillance System (SGSS) to enable reimbursement of associated assay costs in England (parallel reimbursement will be available in the other administrations).
3. **Treating clinicians are asked to ensure that routine PCR testing is undertaken (to enable viral genomic sequencing) through usual testing routes:**
 - on a weekly basis as a minimum during the patient's admission
 - and
 - if and when follow up chest x-rays are undertaken post discharge or the patient attends for any other face to face follow up
4. Clinicians are also asked to ensure that any additional data collection requirements are met for the purpose of relevant surveillance, audit and research around the use of nMABs. It is expected that there will be ongoing monitoring (involving sample collection) of selected patients treated with nMABs (led by Public Health England/UK Health Security Agency, for instance around the potential generation of new variants), as well as academic research to generate new knowledge around clinical effectiveness and other relevant aspects of public health. Organisations are asked to note that intermittent blood sampling ('sparse sampling') may also be required to enable testing of serum concentration levels to complement study pharmacokinetic data. Further information will be provided in due course.
5. Discharge letters to primary care should explicitly record that the casirivimab plus imdevimab combination has been given, together with the dose and date of administration. The following **SNOMED codes should be used to support evaluation and to inform subsequent treatment decisions:**

³ polymerase chain reaction (PCR) test

Presentations:

- Casirivimab 300 mg per 2.5 mL (120 mg/mL) with Imdevimab 300 mg per 2.5 mL (120 mg/mL) 2 vial pack - 40025711000001108
- Casirivimab 1332 mg per 11.1 mL (120 mg/mL) with Imdevimab 1332 mg per 11.1 mL (120 mg/mL) 2 vial pack - 39654011000001101

6. Any organisation treating patients with the casirivimab and imdevimab antibody combination as an off-label product will be required to assure itself that the necessary internal governance arrangements have been completed before the medicine is prescribed. These arrangements may be through the health board / trust drugs and therapeutics committee, or equivalent.
7. Organisations should adhere to the procedures outlined in the [institutional readiness document](#) which has been developed by the Specialist Pharmacy Service to support product storage, preparation and administration.
8. In England, trusts who have not yet done so should register (by site) to participate in COVID-19 specific casirivimab and imdevimab supply arrangements, via Blueteq™. Blueteq should also then be used to confirm pre-authorisation for individual patients. HSC Trusts in Northern Ireland should liaise with the Regional Pharmaceutical Procurement Service to register interest. In Scotland, Health Board Directors of Pharmacy should notify NHS National Procurement if they wish to participate. Health Boards in Wales should notify the All Wales Specialist Procurement Pharmacist of their intention to participate.
9. Organisations should note that following initial nationally determined allocations to participating hospitals, ongoing supplies to each hospital will be replenished on the basis of relative use / need. Ongoing ordering will be through existing (business as usual) routes, supported by volume-based caps (reflecting estimated eligible admissions) where required.
10. Organisations should note that initial supply will be available within 'global pandemic' packaging, which differs from the planned Great Britain (GB) packaging / labelling aligned to the product's GB licence. For example, pandemic packs display a use by date 24 months from manufacture, but current GB regulatory requirements require the product is used within 12 months of manufacture. A 'Dear Healthcare Professional' letter will be available to explain any differences in packaging and supplied packs will be approved for use in both Great Britain and Northern Ireland. **To preserve available supply, hospitals must ensure that packs with shorter use by dates are used first.**
11. Regular stock updates should be provided to trust / hospital and regional pharmacy procurement lead / chief pharmacists. Hospitals should enter the product onto stock control and prescribing systems as described below:

Casirivimab 300 mg per 2.5 mL (120 mg/mL) with Imdevimab 300 mg per 2.5 mL (120 mg/mL) with the dose description as: 2 vial pack

AND/OR

Casirivimab 1332 mg per 11.1 mL (120 mg/mL) with Imdevimab 1332 mg per 11.1 mL (120 mg/mL) with the dose description as: 2 vial pack

Product Details

Ronapreve® is supplied to the UK by Roche. It is a combination neutralising monoclonal antibody (casirivimab plus imdevimab) used to inhibit viral replication in individuals who have not yet mounted an adequate antibody response to the SARS-COV-2 virus following either exposure or vaccination.

The casirivimab plus imdevimab combination for intravenous and subcutaneous use is authorised for use in the treatment and prophylaxis of COVID positive individuals aged 12 and above and weighing at least 40 kg. Supply of the casirivimab and imdevimab combination is subject to the same requirements in both Great Britain and Northern Ireland, and the product information in the Summary of Product Characteristics should be considered applicable across the UK.

Prescribing

The casirivimab plus imdevimab combination product is authorised as a treatment for COVID-19 but the published policy recommends an off-label use. As such, clinicians prescribing this treatment should follow trust / hospital governance procedures in relation to the prescribing of off-label medicines.

Further guidance on the prescribing of off-label medicines can be found below:

- <https://www.gov.uk/drug-safety-update/off-label-or-unlicensed-use-of-medicines-prescribers-responsibilities>
- <https://www.gmc-uk.org/ethical-guidance/ethical-guidance-for-doctors/good-practice-in-prescribing-and-managing-medicines-and-devices/prescribing-unlicensed-medicines>
- <https://www.rpharms.com/Portals/0/RPS%20document%20library/Open%20access/Professional%20standards/Prescribing%20competency%20framework/prescribing-competency-framework.pdf>

Administration

The casirivimab plus imdevimab combination should be administered as an intravenous infusion at a dose of 2.4g (1.2g casirivimab plus 1.2g imdevimab).

A single dose is to be administered. A second dose should not be considered, given the uncertainty over evidence of additional benefit as well as the need to maximise available supply.

Co-Administration

The RECOVERY study demonstrated a mortality benefit for the combination of casirivimab plus imdevimab over and above standard of care in hospitalised COVID positive antibody seronegative patients. Its use should therefore be considered alongside other relevant COVID therapies (including corticosteroids, remdesivir, and tocilizumab or sarilumab) where appropriate and unless contraindicated. There is no interaction of the casirivimab

plus imdevimab combination expected for either dexamethasone or hydrocortisone, remdesivir, or tocilizumab or sarilumab.

For further information please visit the University of Liverpool COVID-19 Drug Interactions website (<https://www.covid19-druginteractions.org/checker>).

The casirivimab plus imdevimab combination should not be infused concomitantly in the same IV line with other medications.

Monitoring, tracking and follow-up

Monitoring of longer-term progress is strongly recommended via recruitment of patients receiving COVID therapies to the [ISARIC-CCP study](#).

All handovers of clinical care (including between hospitals if patients are transferred, between levels of care and clinical teams within hospitals, and between hospitals and primary care) should explicitly record that the casirivimab plus imdevimab combination has been given together with the dose and date of administration. SNOMED codes (see action section, above) should be used in discharge letters to primary care.

Healthcare professionals are asked to report any suspected adverse reactions via the United Kingdom Yellow Card Scheme www.mhra.gov.uk/yellowcard or search for MHRA Yellow Card in the Google Play or Apple App Store8.

Distribution

NHS Trusts (NHS boards in Scotland and Wales)
National / Regional Medical Directors
National / Regional Chief Pharmacists
Lead/Senior Pharmacists and Regional Procurement Pharmacy Leads
Trust/Hospital Pathology Directors (to circulate to pathology networks and laboratory staff)
Trust / Hospital Medical Directors (to circulate to medical and nursing staff managing admitted patients infected with COVID-19)

Enquiries

England

Enquiries from NHS trusts in England should in the first instance be directed to your trust pharmacy team who will escalate issues to the Regional Chief Pharmacist and national teams if required. Further information can be requested from the dedicated email address: england.spoc-c19therapeutics@nhs.net.

Northern Ireland

Enquiries from hospitals in Northern Ireland should in the first instance be directed to your hospital pharmacy team who will escalate issues to the Regional Pharmaceutical Procurement Service or Pharmaceutical Directorate at the Department of Health if required. Further information can be obtained by contacting RPHPS.Admin@northerntrust.hscni.net

Scotland

Enquiries from hospitals in Scotland should in the first instance be directed to your hospital pharmacy team who will escalate issues to either NHS National Procurement or the Scottish Government's Medicines Policy Team if required. Contact should be made using the following emails: nss.nhssmedicineshortages@nhs.scot or medicines.policy@gov.scot

Wales

Enquiries from hospitals in Wales should in the first instance be directed to the health board's Chief Pharmacist who will escalate issues to the Pharmacy and Prescribing Team at Welsh Government if required. Enquiries to the Welsh Government should be directed to: COVID-19.Pharmacy.Prescribing@gov.wales.