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IMMEDIATE MESSAGE TO:

- 1. Directors of Pharmacy
- 2. Medical Directors NHS Boards

28 November 2022

Dear Healthcare Professional

CAS DRUG ALERT – No. 16 – Immediate Effect - Remdesivir For Patients Hospitalised Due To Covid-19

NB This alert updates and replaces alert CEM/CMO/2022/006, previously published in February 2022

Please see attached CMO letter about the updated UK-wide policy (with Immediate Effect) to provide access to Remdesivir for patients hospitalised due to COVID-19.

Remdesivir is an adenosine nucleotide prodrug that is metabolised intracellularly to form the pharmacologically active substrate remdesivir triphosphate. Remdesivir triphosphate inhibits SARS-CoV-2 RNA polymerase which perturbs viral replication. Evidence from the ACTT-1 trial showed that remdesivir improved time to recovery in patients hospitalised with COVID-19 by 5 days compared to placebo (Beigel et al, 2020). The World Health Organization (WHO) Solidarity trial indicated that remdesivir did not improve overall mortality, initiation of ventilation or duration of hospitalisation (Pan et al, 2020). The National Institute for Health and Care Excellence (NICE) recommends the consideration of remdesivir for up to 5 days for COVID-19 pneumonia in adults, and young people 12 years and over weighing 40 kg or more, in hospital and needing low-flow supplemental oxygen under its COVID Rapid Guideline.

Could Medical Directors and Directors of Pharmacy please forward this alert to relevant healthcare professional colleagues, including any clinicians and clinical teams involved in treatment for Patients (Adults and Children) Hospitalised with Covid-19.

Thank you for your co-operation.

Yours sincerely

Grace Jamieson Medicines Policy Team











COVID-19 Therapeutic Alert

CEM/CMO/2022/016 28 November 2022

This alert updates and replaces alert <u>CEM/CMO/2022/006</u>, previously published in February 2022.

Remdesivir for patients hospitalised due to COVID-19

Summary

Remdesivir is an adenosine nucleotide prodrug that is metabolised intracellularly to form the pharmacologically active substrate remdesivir triphosphate. Remdesivir triphosphate inhibits SARS-CoV-2 RNA polymerase which perturbs viral replication.

Evidence from the ACTT-1 trial showed that remdesivir improved time to recovery in patients hospitalised with COVID-19 by 5 days compared to placebo (Beigel et al, 2020). The World Health Organization (WHO) Solidarity trial indicated that remdesivir did not improve overall mortality, initiation of ventilation or duration of hospitalisation (Pan et al, 2020). The National Institute for Health and Care Excellence (NICE) recommends the consideration of remdesivir for up to 5 days for COVID-19 pneumonia in adults, and young people 12 years and over weighing 40 kg or more, in hospital and needing low-flow supplemental oxygen under its COVID Rapid Guideline.

Adults and paediatric patients (at least 4 weeks of age and weighing at least 3 kg) hospitalised due to symptoms of COVID-19 are eligible for treatment under the published UK clinical access policy if they fulfil the following eligibility criteria:

 SARS-CoV-2 infection is confirmed by polymerase chain reaction (PCR) test or where a multidisciplinary team (MDT) has a high level of confidence that the clinical and/or radiological features suggest that COVID-19 is the most likely diagnosis

AND

Hospitalised specifically for the management of COVID-19 symptoms
 AND

 Requiring low-flow¹ supplemental oxygen (see later section on 'Immunocompromised patients' for how this criterion applies to this group)

AND

Presented to hospital not more than 10 days since symptom onset

¹ Low-flow oxygen supplementation as defined in the <u>NICE COVID-19 Rapid Guideline: Managing COVID-19</u>.

AND

- Estimated glomerular filtration rate (eGFR) at least 30 ml/minute
 AND
- Alanine aminotransferase (ALT) below 5 times the upper limit of normal at baseline.

Exemptions to the above eligibility criteria apply to the following patient groups:

- Patients with end-stage renal disease on haemodialysis are exempt from the eGFR treatment threshold above
- Significantly immunocompromised patients (see later section on 'Immunocompromised patients' for exemptions in this cohort).

Remdesivir is also available as a treatment for COVID-19 under separately published UK clinical access policies for <u>non-hospitalised individuals at highest risk from COVID-19 infection</u> and <u>patients with hospital-onset COVID infection</u>, respectively.

Action

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

- Organisations are recommended to consider prescribing remdesivir to adults and paediatric patients (at least 4 weeks of age and weighing at least 3 kg) hospitalised due to COVID-19 in line with the published policy.
 - 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. All healthcare professionals are asked to ensure that any patients who receive a COVID antiviral while pregnant are reported to the UK COVID-19 antivirals in pregnancy registry on 0344 892 0909 so that they can be followed up. For more information, go to http://www.uktis.org/
- 3. Clinicians are encouraged to proactively support recruitment into trials developing further evidence in the treatment of COVID-19. Patients admitted to hospital due to COVID may be considered for entry into the RECOVERY or REMAP-CAP trials.
 - 4. Noting the critical role of surveillance, treating clinicians are strongly encouraged to actively support additional testing or data requirements as requested under country specific or UK wide surveillance programmes, in line with further guidance to be issued. Sequencing is an important part of surveillance activities to monitor for the development of new variants and drug resistance. Therefore in patients being considered for treatment with antivirals, samples pre-treatment and, where part of the clinical pathway, post-treatment, should be prioritised for sequencing. Genotype results do not form part of the eligibility criteria for treatment with remdesivir in this policy and treatment should not be delayed pending these results.

5. Discharge letters to primary care should explicitly record the treatment that 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:

Administration of Remdesivir

Procedure code: 47943005 |Administration of anti-infective agent (procedure)|

Presentation:

- 100mg powder for solution for infusion, 1 vial 38376311000001103
- 6. In England, trusts who have not yet done so should register (by site) to participate in COVID-19 specific medicine 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.
- 7. 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 may be through existing (business as usual) routes, supported by volume-based caps (reflecting estimated eligible admissions) where required.
- 8. Organisations should note that some supply of COVID medicines may be available within 'emergency supply' packaging, which differs from the Great Britain (GB) packaging / labelling aligned to the product's GB licence (or the equivalent product packaging / labelling aligned to the European Medicines Agency marketing authorisation as applicable in Northern Ireland). To preserve available supply, providers must ensure that packs with shorter use by dates are used first.
- 9. 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:
 - Remdesivir 100mg powder for concentrate for solution for infusion

Product Details

Remdesivir (Veklury) is supplied by Gilead. Delivered intravenously, it has a conditional market authorisation for use as a treatment for COVID-19 in both Great Britain (under the Medicines and Healthcare products Regulatory Authority (MHRA)) and in Northern Ireland (under the European Medicines Agency (EMA)) for 1) adults and paediatric patients (at least 4 weeks of age and weighing at least 3 kg) with pneumonia requiring supplemental oxygen (low- or high-flow oxygen or other non-invasive ventilation at start of treatment), and 2)

adults and paediatric patients (weighing at least 40 kg) who do not require supplemental oxygen and who are at increased risk of progressing to severe COVID-19.

Co-Administration

There is no interaction expected for remdesivir with other routine treatments available for COVID under published UK clinical access policies - dexamethasone or hydrocortisone, baricitinib, or tocilizumab or sarilumab.

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

Antivirals 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 <u>ISARIC-CCP study</u>.

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 an antiviral or monoclonal antibody 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 Store.

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.









An Roinn Sláinte

Männystrie O Poustie



Rapid policy statement

Interim Clinical Commissioning Policy: Remdesivir for patients hospitalised due to COVID-19

Publication date: 28 November 2022 Effective from: 28 November 2022

Commissioning position

Remdesivir is recommended to be available as a treatment option through routine commissioning for hospitalised adults and paediatric patients (at least 4 weeks of age and weighing at least 3 kg) with COVID-19 in accordance with the criteria set out in this document.

This policy covers the use of remdesivir for patients hospitalised **due to** symptoms of COVID-19 and to be treated with a 5-day course of remdesivir.

Separate policies cover the use of remdesivir in patients with hospital-onset COVID-19 and non-hospitalised patients with COVID-19.

Background and evidence

Remdesivir is an adenosine nucleotide prodrug that is metabolised intracellularly to form the pharmacologically active substrate remdesivir triphosphate. Remdesivir triphosphate inhibits SARS-CoV-2 RNA polymerase which perturbs viral replication.

Evidence

Current evidence shows that remdesivir improves clinical outcomes in both hospitalised and non-hospitalised patients with COVID-19.

The World Health Organization (WHO) updated its 'Therapeutics and COVID-19: Living guideline' on 16 September 2022 and the recommendations have been considered in the development of this policy (WHO, September 2022).

□ Hospitalised patients: Evidence from the ACTT-1 trial showed that remdesivir improved time to recovery in patients hospitalised with COVID-19 by 5 days compared to placebo (Beigel et al, 2020). The World Health Organization (WHO) Solidarity trial indicated that remdesivir did not improve overall mortality, initiation of ventilation or duration of hospitalisation (Pan et al, 2020).

■ Non-hospitalised patients: evidence from the PINETREE trial, which studied the use of remdesivir in non-hospitalised patients within 7 days of COVID-19 symptom onset and had risk factors for disease progression indicated that a 3-day course of remdesivir resulted in a relative risk reduction of 87% in hospitalisation or death at day 28 compared with placebo (Gottlieb et al, 2020).

Marketing authorisation

Remdesivir delivered intravenously has conditional marketing authorisation for use as a treatment for COVID-19 in Great Britain (under the Medicines and Healthcare Products Regulatory Authority (MHRA)) and a full marketing authorisation in Northern Ireland (under the European Medicines Agency (EMA)) for the following indications:

treatment of COVID-19 in adults and paediatric patients (at least 4 weeks of age and weighing at least 3 kg) with pneumonia requiring supplemental oxygen (low- or high-flow oxygen or other pap invasive ventilation at start of treatment), for a treatment duration of
oxygen or other non-invasive ventilation at start of treatment), for a treatment duration of 5-10 days.

treatment of COVID-19 in adults and paediatric patients (weighing at least 40 kg) who do
not require supplemental oxygen and who are at increased risk of progressing to severe
COVID-19 within 7 days of symptom onset, for a treatment duration of 3 days.

Implementation

Eligibility criteria

Patients hospitalised due to symptoms of COVID-19

Patients are eligible for treatment under this policy if they fulfil the following eligibility criteria:

 SARS-CoV-2 infection is confirmed by polymerase chain reaction (PCR) test or where a multidisciplinary team (MDT) has a high level of confidence that the clinical and/or radiological features suggest that COVID-19 is the most likely diagnosis
 AND

Hospitalised specifically for the management of COVID-19 symptoms
 AND

• Requiring low-flow¹ supplemental oxygen (see later section on 'Immunocompromised patients' for how this criterion applies to this group)

AND

Presented to hospital not more than 10 days since symptom onset

 Estimated glomerular filtration rate (eGFR) at least 30 ml/minute AND

• Alanine aminotransferase (ALT) below 5 times the upper limit of normal at baseline.

Exemptions to the above eligibility criteria apply to the following patient groups:

- Patients with end-stage renal disease on haemodialysis are exempt from the eGFR treatment threshold above
- Significantly immunocompromised patients (see later section on 'Immunocompromised patients' for exemptions in this cohort).

¹ Low-flow oxygen supplementation as defined in the NICE COVID-19 Rapid Guideline: Managing COVID-19.

Considerations during clinical decision-making should include the following (see also the clinical pathway in Appendix 1)².

□ Risk assessment

- Clinical judgement around treatment with remdesivir can be informed by a risk score. Those with a low 4C Mortality Score³ (0 to 3) are highly likely to recover without treatment with remdesivir.
- Remdesivir should not be initiated in patients who present to hospital and are unlikely to survive (determined by clinical judgment). The 4C Mortality Score might be helpful in this assessment.

□ Initiation of treatment

- The decision to initiate treatment with remdesivir should be made by the admitting care consultant.
- Remdesivir should not be initiated in patients who present to hospital more than 10 days after symptom onset (see exemption in immunocompromised patients below).

□ Duration

- All patients treated under this policy must receive a maximum of 5 days of remdesivir in total (comprising a loading dose plus 4 further days of maintenance doses).
- Patients re-admitted for symptoms of COVID-19 (and meeting the eligibility criteria. with the exception of the requirement on the timing from symptom onset) are permitted a second course of up to 5 days upon readmission.
- Significantly immunocompromised patients (see below) are eligible for an extended course of remdesivir (up to 10 days), if agreed following multidisciplinary team assessment.

□ Reassessment and review

The use of remdesivir should be reassessed daily. Consider stopping remdesivir if:

- The patient clinically improves and no longer requires supplemental oxygen 72 hours after commencement of treatment: or
- The patient continues to deteriorate despite 48 hours of sustained mechanical ventilation.

☐ Immunocompromised patients

For significantly immunocompromised patients⁴ hospitalised for COVID-19 symptoms:

- A course of remdesivir can be extended to a maximum of 10 days
- The criterion on time between symptom onset and treatment initiation does not apply
- The criterion on the need for supplemental oxygen requirement does not apply.

² Clinical judgement in the initiation, review, escalation and de-escalation of patients receiving remdesivir treatment should be supported where possible by multidisciplinary team assessment.

³ The 4C Mortality Score (available at https://isaric4c.net/risk/) is a validated risk stratification score, which can help inform clinical decision making for patients admitted to hospital with COVID-19 (Knight et al, 2020). Other clinical risk scores are available.

⁴ Refers to patients with a significant impairment of humoral immune response (antibody production) and/or cellular immune competence.

Patients with hospital-onset COVID-19

Please refer to the UK Clinical Commissioning Policy for antivirals or nMABs for <u>hospitalised</u> <u>patients with COVID-19</u>.

Non-hospitalised patients with COVID-19

Please refer to the UK Clinical Commissioning Policy for antivirals or nMABs for non-hospitalised patients with COVID-19.

Where possible, all patients being considered for treatment with remdesivir should have samples taken for serology (anti-S (spike) antibody) prior to treatment. However, SARS-CoV-2 antibody tests or results are **not** a requirement for treatment with remdesivir under the criteria specified in this policy.

Exclusion criteria and cautions

Patients who meet any of the following exclusion criteria are NOT eligible for treatment with remdesivir under this policy:

- Children aged less than 4 weeks of age and/or weighing less than 3kg.
- Known hypersensitivity reaction to the active substances or to any of the excipients of remdesivir as listed in the Summary of Product Characteristics for <u>Great Britain</u> and <u>Northern Ireland</u>.
- Estimated glomerular filtration rate (eGFR) <30 mL/min (except in patients with endstage renal disease on haemodialysis).
- Alanine transaminase (ALT) ≥ 5 times the upper limit of normal.

Remdesivir should be discontinued in patients who develop **any** of the following:

- ALT ≥ 5 times the upper limit of normal during treatment with remdesivir (remdesivir may be restarted when ALT is < 5 times the upper limit of normal)
- ALT elevation accompanied by signs or symptoms of liver inflammation or increasing conjugated bilirubin, alkaline phosphatase, or international normalised ratio (INR)

An individual clinical decision should be made as to whether pre-treatment urea and electrolytes and liver function tests are required, based upon whether recent bloods are available or the patient is considered at risk of undiagnosed liver or kidney disease.

Hypersensitivity reactions including infusion-related and anaphylactic reactions have been observed during and following administration of remdesivir. Signs and symptoms may include hypotension, hypertension, tachycardia, bradycardia, hypoxia, fever, dyspnoea, wheezing, angioedema, rash, nausea, vomiting, diaphoresis, and shivering. Slower infusion rates, with a maximum infusion time of up to 120 minutes, can be considered to potentially prevent these signs and symptoms. Patients should be monitored for hypersensitivity reactions during and following administration of remdesivir as clinically appropriate. If signs and symptoms of a clinically significant hypersensitivity reaction occur, administration of remdesivir should be discontinued immediately and appropriate treatment initiated.

Dose

The recommended dosage for adults and paediatric patients (weighing at least 40kg) under this policy is a single loading dose of remdesivir 200 mg intravenously on day 1, followed by a once daily maintenance dose of remdesivir 100 mg for the remainder of the treatment course, which should not exceed five days (see exemption in immunocompromised patients above).

The recommended dosage for paediatric patients at least 4 weeks old (weighing at least 3kg but less than 40kg) under this policy is a single loading dose of remdesivir 5mg/kg intravenously on day 1, followed by a once daily maintenance dose of remdesivir 2.5mg/kg for the remainder of the treatment course, which should not exceed five days (see exemption in immunocompromised patients above).

Administration

For adults and paediatric patients (weighing at least 40kg), 200mg of remdesivir (day 1 loading dose) and 100mg of remdesivir (days 2-5 maintenance doses) should be diluted in either a 250ml or 100ml pre-filled bag of 0.9% sodium chloride solution and infused over a minimum of 30 minutes. Treatment should be initiated as soon as possible after admission to hospital and within 10 days of symptom onset.

For paediatric patients at least 4 weeks old (weighing at least 3kg but less than 40kg), the single loading dose of remdesivir (5mg/kg intravenously on day 1) and the once daily maintenance dose (2.5mg/kg on days 2-5) should be diluted in either a 25ml, 50ml or 100ml pre-filled bag of 0.9% sodium chloride solution and infused over a minimum of 30 minutes. Treatment should be initiated as soon as possible after admission to hospital and within 10 days of symptom onset.

Renal and liver function should be monitored carefully during treatment with remdesivir as clinically appropriate.

Genotyping and sequencing of samples

Sequencing is an important part of surveillance activities to monitor for the development of new variants and drug resistance. Therefore, in patients being considered for treatment with remdesivir, samples pre-treatment and where part of the clinical pathway, post-treatment, should be prioritised for sequencing. Genotype results do not form part of the eligibility criteria for treatment with remdesivir and treatment should not be delayed pending these results.

Pregnancy

There are no or limited amount of data from the use of remdesivir in pregnant women. Remdesivir should be **avoided** in pregnancy unless clinicians believe the benefits of treatment outweigh the risks to the individual (please see the relevant SmPC for further information). Women of child-bearing potential have to use effective contraception during treatment.

Co-administration

There is no interaction expected between remdesivir and other commissioned treatments for COVID-19. For further information please visit the University of Liverpool COVID-19 Drug Interactions website (https://www.covid19-druginteractions.org/checker).

Please refer to other published UK clinical commissioning policies setting out available COVID-19 treatments here.

Safety reporting

Any suspected adverse drug reactions (ADRs) for patients receiving remdesivir should be reported directly to the MHRA via the new dedicated COVID-19 Yellow Card reporting site at: https://coronavirus-yellowcard.mhra.gov.uk/

Governance

Data collection requirement

Provider organisations in England should register all patients using prior approval software

(alternative arrangements in Scotland, Wales and Northern Ireland will be communicated) and ensure monitoring arrangements are in place to demonstrate compliance against the criteria as outlined.

Clinical outcome reporting

Hospitals managing COVID-19 patients are encouraged to submit data through the ISARIC 4C Clinical Characterisation Protocol (CCP) case report forms (CRFs), as coordinated by the COVID-19 Clinical Information Network (CO-CIN) (https://isaric4c.net/protocols/).

Effective from

This policy will be in effect from the date of publication.

Policy review date

This is an interim rapid clinical policy statement, which means that the full process of policy production has been abridged: public consultation has not been undertaken. This policy may need amendment and updating if, for instance, new trial data emerges, supply of the drug changes, or a new evidence review is required. A NICE Technology Appraisal or Scottish Medicines Consortium (SMC) Health Technology Assessment or All Wales Medicines Strategy Group (AWMSG) appraisal of remdesivir for COVID-19 would supersede this policy when completed.

Surveillance and service evaluation

There is an urgent need to generate more evidence and greater understanding around use of treatments in patients with COVID-19. Both surveillance and service evaluation are necessary to gain knowledge around the following: factors of relevance in determining neutralising monoclonal antibodies (nMABs) and antiviral treatment; the impact of nMAB and antiviral treatment in the community and hospital settings on the immune/virologic response and clinical recovery; and the public health sequelae of nMAB and antiviral use, such as generation of new mutations and/or variants.

Treating clinicians are asked to ensure that all PCR tests undertaken as an inpatient and/or in the community where any patient who is receiving ongoing PCR testing as part of secondary care (for example, through an outpatient clinic) should do this through the hospital laboratory where these samples should be retained for sequencing. Further serial sampling for specific patient groups may be requested as part of UKHSA genomic surveillance purposes, or country specific programmes.

Clinicians must ensure that any additional data collection requirements are met for the purpose of relevant surveillance, audit and evaluation around the use of nMABs and antivirals. It is expected that there will be ongoing monitoring (involving sample collection) of selected patients treated with nMABs and antivirals (led by UKHSA, 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.

Equality statement

Promoting equality and addressing health inequalities are at the heart of the four nations' values. Throughout the development of the policies and processes cited in this document, we have:

- Given due regard to the need to eliminate discrimination, harassment and victimisation, to advance equality of opportunity, and to foster good relations between people who share a relevant protected characteristic (as cited under the Equality Act 2010 or equivalent equality legislation) and those who do not share it; and
- Given regard to the need to reduce inequalities between patients in access to and

outcomes from healthcare services and to ensure services are provided in an integrated way where this might reduce health inequalities.

Definitions

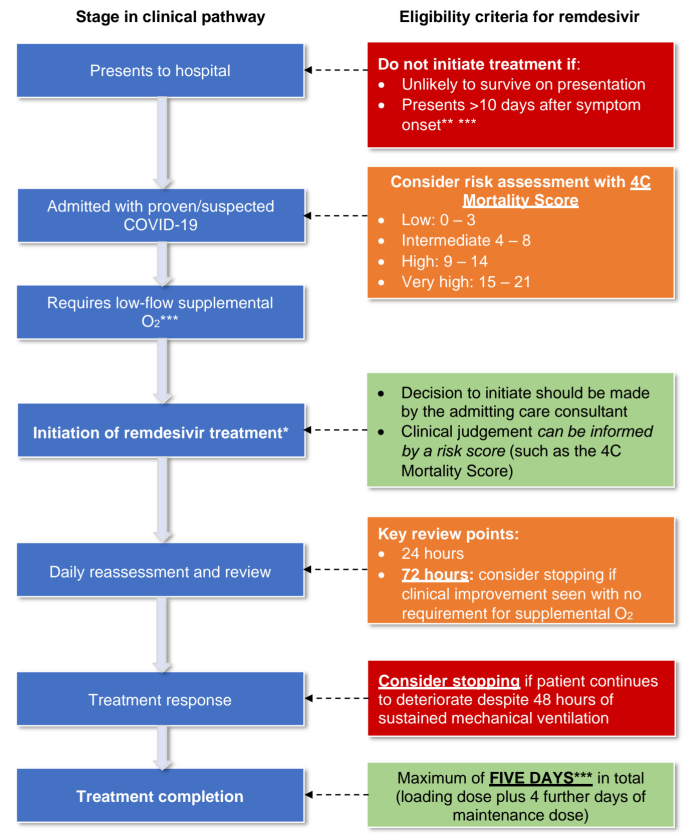
COVID-19	Refers to the disease caused by the severe acute respiratory syndrome coronavirus-2 (SARS-CoV-2) virus.
Mechanical ventilation	A life support treatment which helps people breathe when they are not able to breathe enough on their own.
Extra Corporeal Membrane Oxygenation	A life support machine for people who have a severe and life-threatening illness that stops their heart or lungs from working properly.

References

- 1. Beigel JH, Tomashek KM, Dodd LE, et al. Remdesivir for the Treatment of Covid-19 -Final Report. N Engl J Med. 2020;383(19):1813-1826. doi:10.1056/NEJMoa2007764
- 2. Gottlieb RL, Vaca CE, Paredes R, et al. Early Remdesivir to Prevent Progression to Severe Covid-19 in Outpatients. N Engl J Med. 2022;386(4):305-315. doi:10.1056/NEJMoa2116846
- 3. WHO Solidarity Trial Consortium, Pan H, Peto R, et al. Repurposed Antiviral Drugs for Covid-19 - Interim WHO Solidarity Trial Results. N Engl J Med. 2021;384(6):497-511. doi:10.1056/NEJMoa2023184

Appendix 1

Clinical pathway and criteria for the use of remdesivir in patients hospitalised with COVID-19



^{*}There should be careful consideration before initiating remdesivir treatment

^{**}Unless patient is readmitted with COVID-19 (see main policy document)

^{***}See additional comments on immunocompromised patients

Clinical pathway: Therapies for patients hospitalised due to COVID-19

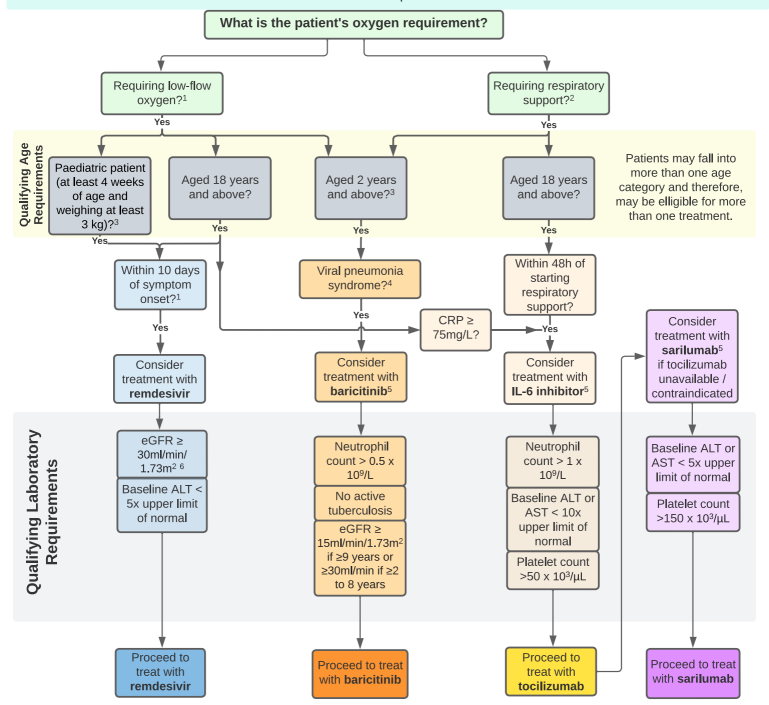
- This guide aims to support treatment decisions for commissioned COVID-19 therapies and outlines their position in the treatment pathway
 for patients hospitalised due to COVID-19. The relevant clinical commissioning policies should be consulted for further details
- Patients must be hospitalised specifically for management of COVID-19 and must be receiving supplemental oxygen or receiving respiratory support
- · Consult the relevant Summary of Product Characteristics for advice on contraception and use in pregnancy
- Please refer to the NICE COVID-19 Rapid Guideline (NG 191) for other treatments

CORTICOSTEROIDS

Consider dexamethasone (or hydrocortisone or prednisolone if treatment with dexamethasone is unavailable/not possible) in patients who require supplemental oxygen to maintain prescribed oxygen saturation levels

TRIALS

All **hospitalised** patients can consider joining the RECOVERY trial or the pandemic aspects of the REMAP-CAP trial. To enter RECOVERY, they should have: a **viral pneumonia syndrome**; confirmed **SARS-CoV-2 infection**; and no **medical history** that might put the patient at risk from entering a trial. To enter REMAP-CAP, they should be in critical care with an **acute illness due to suspected pandemic illness**. Patients can be referred for entry into clinical trials at any stage in this clinical pathway and will continue to receive treatment under this pathway in addition to any trial medication prescribed.



Deterioration - Consider other therapeutic agent(s) from group above in accordance with respective clinical policies

¹ For treatment with remdesivir, the criteria relating to supplemental oxygen and the treatment window from symptom onset do not apply to significantly immunocompromised patients.

² Defined as: high-flow nasal oxygen, continuous positive airway pressure (CPAP) or non-invasive ventilation, or invasive mechanical ventilation.
³ Clinicians should seek paediatric MDT advice for paediatric patients to determine clinical capacity to benefit from treatment.

⁴ In general, viral pneumonia should be suspected when a patient presents with: a) typical symptoms (e.g. influenza-like illness with fever and muscle pain, or respiratory illness with cough and shortness of breath); AND b) compatible chest X-ray findings (consolidation or ground-glass shadowing); AND c) alternative causes have been considered unlikely or excluded (e.g. heart failure, bacterial pneumonia).

⁵ Baricitinib may be administered in combination with IL-6 receptor blockers (as well as corticosteroids, unless contraindicated), according to clinical judgement, in patients with severe or critical COVID-19. If an IL-6 inhibitor is not deemed suitable, or eligibility criteria (for an IL-6 inhibitor) are unmet, baricitinib treatment may still be considered.

⁶ Patients with end-stage renal disease on haemodialysis are exempt from the specified eGFR threshold.