## Sample Form (v15.00 - 11/05/21)

## **Randomisation Program**

Call Freefone 0800 138 5451 to contact the RECOVERY team for URGENT problems using the Randomisation Program or for medical advice. All NON-URGENT queries should be emailed to recoverytrial@ndph.ox.ac.uk

	Logged in as: F
	Section A: Baseli
Treating clinician	Date and time of randomis
A1. Name of treating clinician	
Patient details  A2. Patient surname	
Patient forename	
A3. NHS number	☐ Tick if not available
A4. What is the patient's date of birth?	·/ · ·/ ·
A5. What is the patient's sex?	~
Inclusion criteria  A6. Has consent been taken in line with the protocol?  If answer is No patient cannot be enrolled in the study	<u> </u>
A7. Does the patient have proven or suspected SARS-CoV- 2 infection? If answer is No patient cannot be enrolled in the study	_
A8. Does the patient have any medical history that might, in the opinion of the attending clinician, put the patient at	v
significant risk if they were to participate in the trial? <b>A9.</b> COVID-19 symptom onset date:	~// ~// ~
A10. Date of hospitalisation:	
A11. Date or nospitalisation:  A11. Does the patient require oxygen?	
A12. Please select one of the following to describe the	
current level of ventilation support	<b>v</b>
A12.1 Enter latest oxygen saturation measurement (%)	
A12.2 Enter latest CRP measurement since admission to hospital (mg/L) Enter 0 if below the limit of measurement	Tick if greater than limit of measurement
A12.3 Enter latest creatinine measurement since admission to hospital	μmol/L ▼ □Tick if not measured
A12.4 Enter latest D-dimer measurement since admission to hospital	ng/mL v Tick if not measured
Enter 0 if below the limit of measurement	Tick if greater than limit of measurement
A12.5 Has the patient received a COVID-19 vaccine?  Does the patient have any CURRENT comorbidities or	other medical problems or treatments?
A13.1 Diabetes	~
A13.2 Heart disease	v
A13.3 Chronic lung disease	•
A13.4 Tuberculosis	~
A13.5 HIV	~
A13.6 Severe liver disease	v
A13.7 Severe kidney impairment (eGFR<30 or on dialysis)	<b>v</b>
A13.8 Known long QT syndrome	
A13.9 Current treatment with macrolide antibiotics which are to continue  Macrolide antibiotics include clarithromycin, azithromycin and erythromycin	
A13.10 Antiplatelet therapy Includes aspirin, clopidogrel, ticagrelor, prasugrel,	<u> </u>
dipyridamole  A13.12 Has received tocilizumab therapy during this	
admission  Are the following treatments UNSUITABLE for the p	
If you answer Yes it means you think this patient sho A14B.1 Synthetic monoclonal antibodies	uld NOT receive this drug.
(REGN10933+REGN10987)	
A14D.1 Baricitinib is NOT suitable if patient (i) is pregnant; (ii) has eGFR <15 ml/min or is on dialysis/haemofiltration; (iii) has active TB; or (iv) has neutrophil count <0.5	~
Are the following treatments available?  A15B.1 Synthetic monoclonal antibodies	Yes v
(REGN10933+REGN10987)  A15D.1 Baricitinib	
Current medication	
A16.1 Is the patient currently prescribed remdesivir?	
A16.2 Is the patient currently prescribed systemic corticosteroids (dexamethasone, prednisolone, hydrocortisone, methylprednisolone)?  Please do not include topical or inhaled treatments	
A16.4 Is the patient currently on warfarin or a direct oral anticoagulant?  Includes apixaban, rivaroxaban	v
A16.5 What venous thromboembolism prophylaxis is the patient receiving?  Standard — usual for hospitalised patients (not increased due to COVID-19; higher dose — treatment dose or increased prophylaxis due to COVID-19	٧
Serum sample collection  A17.1 Have you sent a serum sample for coronavirus	<b>V</b>
antibody measurement to your transfusion laboratory?  Please sign off this form once complete	
Surname:	
Forename:	
Professional email:	
	Continue
	Cancel