

Global COVID-19 Clinical Platform

NOVEL CORONAVIRUS (COVID-19) - RAPID VERSION

DESIGN OF THIS CASE RECORD FORM (CRF)

This CRF has 3 modules:

Module 1 to be completed on the first day of admission to the health centre.

Module 2 to be completed on first day of admission to ICU or high dependency unit. Module 2 should also be completed daily for as many days as resources allow. Continue to follow-up patients who transfer between wards.

Module 3 to be completed at discharge or death.

GENERAL GUIDANCE

- The CRF is designed to collect data obtained through examination, interview and review of hospital notes. Data may be collected retrospectively if the patient is enrolled after the admission date.
- Participant Identification Numbers consist of a site code and a participant number. You can obtain a site code and register on the data management system by contacting <u>ncov@isaric.org</u>.
 Participant numbers should be assigned sequentially for each site beginning with 00001. In the case of a single site recruiting participants on different wards, or where it is otherwise difficult to assign sequential numbers, you can assign numbers in blocks or incorporate alpha characters. E.g. Ward X will assign numbers from 00001 or A0001 onwards and Ward Y will assign numbers from 50001 or B0001 onwards. Enter the Participant Identification Number at the top of every page.
- Data are entered to the central electronic REDCap database at https://ncov.medsci.ox.ac.uk or to your site/network's independent database. Printed paper CRFs may be used and the data can be typed into the electronic database afterwards.
- Complete every section. Questions marked "If yes,..." should be left blank when they do not apply (i.e. when the answer is not yes).
- Selections with square boxes (
) are single selection answers (choose one answer only).
- Selections with circular boxes (**O**) are multiple selection answers (choose all that apply).
- Mark 'Unknown' for any data that are not available or unknown.
- Avoid recording data outside of the dedicated areas.
- If using paper CRFs, we recommend writing clearly in ink, using BLOCK-CAPITAL LETTERS.
- Place an (X) in the boxes to mark the answer. To make corrections, strike through (------) the data you wish to delete and write the correct data above it. Please initial and date all corrections.
- Please keep all of the sheets for a single participant together e.g. with a staple or participant-unique folder.
- Please transfer all paper CRF data to the electronic database. All paper CRFs can be stored by the institution responsible for them. All data should be transferred to the secure electronic database.
- Please enter data on the electronic data capture system at https://ncov.medsci.ox.ac.uk. If your site would like to collect data independently, we can support the establishment of locally hosted databases.
- Please contact us at <u>ncov@isaric.org</u>. If we can help with databases, if you have comments and to let us know that you are using the forms.



MODULE 3: complete at discharge/death

DIAGNOSTIC/PATHOGEN TESTING Chest X-Ray /CT performed? Yes No Unknown If Yes: infiltrates present? Yes No Unknown Was pathogen testing done during this illness episode? DYes DNo DUnknown If yes, complete all below: Influenza virus: Positive Negative Not done If positive, type Coronavirus: Desitive Desitive Negative Not done If positive: MERS-CoV DSARS-CoV-2 Other **Other respiratory pathogen:** Positive Degative Dot done **If positive,** specify Viral haemorrhagic fever: Positive Negative Not done If positive, specify virus Other pathogen of public health interest detected: If yes, specify: _____ Falciparum malaria: Positive Negative Not done Non-falciparum malaria: Positive Negative Not done **HIV:** Positive Negative Not done COMPLICATIONS: At any time during hospitalisation did the patient experience: □Yes □No □Unknown Bacteraemia □Yes □No □Unknown Shock □Yes □No □Unknown □Yes □No □Unknown Bleeding Seizure Meningitis/Encephalitis □Yes □No □Unknown Endocarditis □Yes □No □Unknown Anaemia □Yes □No □Unknown Myocarditis/Pericarditis □Yes □No □Unknown □Yes □No Cardiac arrhythmia □Yes □No □Unknown Acute renal injury □Unknown □Yes □No □Unknown □Yes □No □Unknown Pancreatitis Cardiac arrest □No □Unknown □Yes □No □Unknown □Yes Liver dysfunction Pneumonia □No □Unknown □Unknown **Bronchiolitis** □Yes Cardiomyopathy □Yes □No Acute Respiratory Distress □Yes □No □Unknown Other □Yes □No □Unknown If Yes, specify Syndrome MEDICATION: While hospitalised or at discharge, were any of the following administered? Oral/orogastric fluids? UYes No Unknown Intravenous fluids? UYes No Unknown Antiviral? DYes Do DUnknown If yes: ORibavirin OLopinavir/Ritonavir ONeuraminidase inhibitor OInterferon alpha OInterferon beta OOther, specify: Antibiotic? DYes DNo DUnknown If yes, specify: Corticosteroid? UYes No Unknown If yes, route: OOral OIntravenous OInhaled If yes, specify agent and maximum daily dose: _ Antifungal agent? Yes No Unknown If yes, specify: _ Antimalarial agent? UYes No Unknown If yes, specify: _ Experimental agent? DYes DNo DUnknown If yes, specify: _ SUPPORTIVE CARE: At ANY time during hospitalisation, did the patient receive/undergo: ICU or High Dependency Unit admission? UYes No Unknown If yes, total duration: davs Date of ICU admission:[_D_][_D_]/[_M_][_M_]/[_2_][_0_][_Y_][_Y_] □N/A **Oxygen therapy?** TYes No Unknown If yes, complete all: Total duration: days O₂ flow volume: O1-5 L/min O6-10 L/min O11-15 L/min O>15 L/min Source of oxygen: OPiped OCylinder OConcentrator Interface: ONasal prongs OHF nasal cannula OMask OMask with reservoir OCPAP/NIV mask Non-invasive ventilation? (e.g. BIPAP, CPAP) □Yes □No □ Unknown If yes, total duration: days Invasive ventilation (Any)? □Yes □No □Unknown If yes, total duration: ____ days Extracorporeal (ECMO) support? Yes No Unknown If yes, total duration: davs Prone position? UYes No Unknown If yes, total duration: days Renal replacement therapy (RRT) or dialysis? UYes UNo Unknown Inotropes/vasopressors? Yes No Unknown If yes, total duration: _____ days OUTCOME Outcome: Discharged alive Hospitalized Transfer to other facility Death Palliative discharge Unknown **Outcome date:** [_D_][_D_]/[_M_][_M_]/[_2_][_0_][_Y_][_Y_] □Unknown If Discharged alive: Ability to self-care at discharge versus before illness: Same as before illness: Worse □Better □Unknown