

## COVID-19 CORE CASE REPORT FORM

### ACUTE RESPIRATORY INFECTION CLINICAL CHARACTERISATION DATA TOOL

#### DESIGN OF THIS CASE REPORT FORM (CRF)

This CRF is set up in modules to be used for recording data on the ISARIC\_nCov Core Database or for independent studies.

**Module 1 and Module 2** complete on the first day of presentation/admission or on first day of COVID-19 assessment.

**Module 2** also complete on first day of admission to ICU or high dependency unit. In addition, complete daily for as many days as resources allow up to a maximum of 14 days. Continue to follow-up patients who transfer between wards.

**Module 3 (Outcome)** complete 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 prospectively or retrospectively if the patient is enrolled after the admission date.
- Participant Identification Numbers consist of a 5 digit site code and a 4 digit participant number. You can obtain a site code and registering on the data management system by contacting [ncov@isaric.org](mailto:ncov@isaric.org). Participant numbers should be assigned sequentially for each site beginning with 0001. In the case of a single site recruiting participants on different wards, or where it is otherwise difficult to assign sequential numbers, it is acceptable to assign numbers in blocks or incorporating alpha characters. E.g. Ward X will assign numbers from 0001 or A001 onwards and Ward Y will assign numbers from 5001 or B001 onwards. Enter the Participant Identification Number at the top of every page.
- Printed paper CRFs may be used for later transfer of the data onto the electronic database.
- For participants who return for re-admission to the same site, **start a new form with the same Participant Identification Number**. Please check “YES-admitted previously” in the ONSET & ADMISSION section. Enter as 2 separate entries in the electronic database.
- For participants who transfer between two sites that are both collecting data on this form, it is preferred to have the data entered by a single site as a single admission, under the same Participant Identification Number. When this is not possible, the first site should record “Transfer to other facility” as an OUTCOME, and the second site should start a new form with a new patient number and indicate “YES-transferred” in ONSET & ADMISSION.
- Complete every line of every section, except where the instructions say to skip a section based on a response.
- Selections with circles (●) are single selection answers (choose one answer only). Selections with square boxes (□) are multiple selection answers (choose as many answers as are applicable).
- Mark ‘Not done’ for any results of laboratory values that are not available, not applicable or unknown.
- Avoid recording data outside of the dedicated areas. Sections are available for recording additional information.
- If using paper CRFs, we recommend writing clearly in ink, using BLOCK-CAPITAL LETTERS.
- Place an (X) when you choose the corresponding 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 needs to be stored locally, do not send any forms to us. Data are accepted only via 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 are happy to support the establishment of locally hosted databases.
- Please contact us at [ncov@isaric.org](mailto:ncov@isaric.org) if you need help with databases, if you have comments and to let us know that you are using the forms.



**MODULE 1: PRESENTATION/ADMISSION CASE REPORT FORM**

<b>SIGNS AND SYMPTOMS ON ADMISSION</b> ( <i>Unk = Unknown</i> )			
History of fever	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk	Fatigue / Malaise	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
Cough	<input type="radio"/> YES-non-productive <input type="radio"/> YES-productive	Anorexia	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
	<input type="radio"/> YES-with haemoptysis <input type="radio"/> NO <input type="radio"/> Unk	Altered consciousness/confusion	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
Sore throat	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk	Muscle aches (myalgia)	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
Runny nose (rhinorrhoea)	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk	Joint pain (arthralgia)	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
Wheezing	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk	Inability to walk	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
Shortness of breath	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk	Abdominal pain	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
Lower chest wall indrawing	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk	Diarrhoea	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
Chest pain	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk	Vomiting / Nausea	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
Conjunctivitis	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk	Skin rash	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
Lymphadenopathy	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk	Bleeding (Haemorrhage) If YES, specify site(s):	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
Headache	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk		
Loss of smell (Anosmia)	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk	Other symptom(s) If YES, specify:	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
Loss of taste (Ageusia)	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk		
Seizures	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk		

<b>PRE-ADMISSION MEDICATION</b> ( <i>taken within 14 days of admission/presentation at healthcare facility</i> )	
Angiotensin converting enzyme inhibitors (ACE inhibitors)	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
Angiotensin II receptor blockers (ARBs)	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
Non-steroidal anti-inflammatory (NSAIDs)	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
Oral steroids	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk If YES, agent(s):
Other immunosuppressant agents (not oral steroids)	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk If YES, agent(s):
Antivirals	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk If YES, agent(s):
Antibiotics	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk If YES, agent(s):
Other targeted COVID-19 Medications	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk If YES, agent(s):

<b>CO-MORBIDITIES AND RISK FACTORS</b> ( <i>existing prior to admission and ongoing</i> )			
Chronic cardiac disease ( <i>not hypertension</i> )	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk	Chronic hematologic disease	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
Hypertension	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk	AIDS / HIV	<input type="radio"/> YES-on ART <input type="radio"/> YES-not on ART <input type="radio"/> NO <input type="radio"/> Unk
Chronic pulmonary disease ( <i>not asthma</i> )	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk	Diabetes Mellitus	<input type="radio"/> YES-Type 1 <input type="radio"/> YES -Type 2 <input type="radio"/> NO <input type="radio"/> Unk
Asthma ( <i>physician diagnosed</i> )	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk	Rheumatologic disorder	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
Chronic kidney disease	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk	Dementia	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
Obesity ( <i>as defined by clinical staff</i> )	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk	Tuberculosis	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
Moderate or severe liver disease	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk	Malnutrition	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
Mild liver disease	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk	Smoking	<input type="radio"/> YES <input type="radio"/> Never smoked <input type="radio"/> Former smoker <input type="radio"/> Unk
Asplenia	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk	Other relevant risk factor(s) If YES, specify:	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
Chronic neurological disorder	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk		
Malignant neoplasm	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk		

## MODULE 2: DAILY CASE REPORT FORM

Complete on the day of admission or first COVID-19 investigation, and on the first day of ICU admission (if different from day of admission). In addition, depending on available resources, complete every day for a maximum of 14 days, or for days when biochemical results are available.

**SIGNS AND SYMPTOMS (Record the worst value between 00:00 to 24:00 on day of assessment)(worst=furthest from normal range)**
**DATE OF ASSESSMENT (DD/MM/YYYY):** [ \_ ][ \_ ]/[ \_ M ][ \_ M ]/[ \_ 2 ][ \_ 0 ][ \_ Y ][ \_ Y ]

**Temperature:** [ ][ ][ ]·[ ][ ] °C or [ ][ ][ ] °F      **HR:** [ ][ ][ ] beats per minute      **RR:** [ ][ ][ ] breaths per minute

**Systolic BP:** [ ][ ][ ] mmHg      **Diastolic BP:** [ ][ ][ ] mmHg      **Oxygen saturation SaO<sub>2</sub>** [ ][ ][ ] %

**Any supplemental oxygen: FiO<sub>2</sub> (0.21-1.0)** [ ][ ]·[ ][ ][ ] or [ ][ ][ ] % or [ ][ ][ ] L/min

**Sternal capillary refill time >2seconds**    YES    NO    Unknown

**AVPU:** Alert [ ][ ] Verbal [ ][ ] Pain [ ][ ] Unresponsive [ ][ ]      **Glasgow Coma Score (GCS / 15)** [ ][ ][ ]

**Is the patient currently receiving, or has received (between 00:00 to 24:00 on day of assessment)**
**High-flow nasal cannula oxygen therapy?**       YES    NO    Unknown

**Non-invasive ventilation (Any)?**  YES    NO    Unknown    **If YES:**  BIPAP    CPAP    Other    Unknown

**Invasive ventilation?**       YES    NO    Unknown

**Prone positioning?**       YES    NO    Unknown

**Inhaled Nitric Oxide?**       YES    NO    Unknown

**Tracheostomy inserted?**       YES    NO    Unknown

**Extra corporeal life support (ECLS/ ECMO)?**       YES    NO    Unknown    **If YES:**  VV    AV    Central    Unknown

**Renal replacement therapy (RRT) or dialysis?**    YES    NO    Unknown

**Any vasopressor/inotropic support?**  YES    NO    Unknown (if NO, select NO for the next 3 questions)

**Dopamine <5µg/kg/min OR Dobutamine OR milrinone OR levosimendan:**       YES    NO

**Dopamine 5-15µg/kg/min OR Epinephrine/Norepinephrine < 0.1µg/kg/min OR vasopressin OR phenylephrine:**  YES    NO

**Dopamine >15µg/kg/min OR Epinephrine/Norepinephrine > 0.1µg/kg/min:**       YES    NO

**Neuromuscular blocking agents?**  YES    NO    Unknown

**Other intervention(s) or procedure(s)?**  YES    NO    Unknown    **If YES, Specify:** \_\_\_\_\_

**Current admission to ICU/ITU/IMC/HDU?**  YES    NO    Unknown      (Record the worst value on day of assessment)

**PaO<sub>2</sub> (at time nearest to the FiO<sub>2</sub> recorded at top of page)** [ ][ ][ ][ ]  kPa or  mmHg     Not done

**PaO<sub>2</sub> sample type:**  Arterial    Capillary    Unknown

**From same blood gas record as PaO<sub>2</sub>:**
**PCO<sub>2</sub>** \_\_\_\_\_  kPa or  mmHg    |    **pH** \_\_\_\_\_    |    **HCO<sub>3</sub><sup>-</sup>** \_\_\_\_\_ mEq/L    |    **Base excess** \_\_\_\_\_ mmol/L

**Richmond Agitation-Sedation Scale (RASS)** [ ][ ]    or    **Riker Sedation-Agitation Scale (SAS)** [ ][ ]     Unknown

**Mean Arterial Blood Pressure** [ ][ ][ ][ ] mmHg     Unknown

**Urine flow rate** [ ][ ][ ][ ][ ][ ][ ][ ] mL/24 hours     Check if estimated     Unknown

## MODULE 2: DAILY CASE REPORT FORM

Complete on the day of admission or first COVID-19 investigation, and on the first day of ICU admission (if different from day of admission). In addition, depending on available resources, complete every day for a maximum of 14 days, or for days when biochemical results are available.

**LABORATORY RESULTS (on admission, on any admission to ICU, then daily) – complete every line**
**DATE OF ASSESSMENT (DD/MM/YYYY):** [ \_D\_ ][ \_D\_ ]/[ \_M\_ ][ \_M\_ ]/[ \_2\_ ][ \_0\_ ][ \_Y\_ ][ \_Y\_ ]

Record the worst value between 00:00 to 24:00 on day of assessment (if Not Available write 'N/A'):

**LABORATORY RESULTS** (\*record units if different from those listed)

Parameter	Value*	Not done	Parameter	Value*	Not done
Haemoglobin (g/L)		<input type="radio"/>	Urea (BUN) (mmol/L)		<input type="radio"/>
WBC count (x10 <sup>9</sup> /L)		<input type="radio"/>	Lactate (mmol/L)		<input type="radio"/>
Lymphocyte count (10 <sup>9</sup> /L)		<input type="radio"/>	Creatinine (µmol/L)		<input type="radio"/>
Neutrophil count (10 <sup>9</sup> /L)		<input type="radio"/>	Sodium (mmol/L)		<input type="radio"/>
Haematocrit (%)		<input type="radio"/>	Potassium (mmol/L)		<input type="radio"/>
Platelets (x10 <sup>9</sup> /L)		<input type="radio"/>	Procalcitonin (ng/mL)		<input type="radio"/>
APTT (seconds)		<input type="radio"/>	CRP (mg/L)		<input type="radio"/>
APTR		<input type="radio"/>	LDH (U/L)		<input type="radio"/>
PT (seconds)		<input type="radio"/>	Creatine kinase (U/L)		<input type="radio"/>
INR		<input type="radio"/>	Troponin I (ng/mL)		<input type="radio"/>
ALT/SGPT (U/L)		<input type="radio"/>	D-dimer (mg/L)		<input type="radio"/>
Total bilirubin (µmol/L)		<input type="radio"/>	Ferritin (ng/mL)		<input type="radio"/>
AST/SGOT (U/L)		<input type="radio"/>	IL-6 (pg/mL)		<input type="radio"/>
Glucose (mmol/L)		<input type="radio"/>			

### MODULE 3: OUTCOME CASE REPORT FORM

**TREATMENT: At ANY time during hospitalisation, did the patient receive/undergo:**
**Any Oxygen therapy?**  YES  NO  Unknown      **If YES, total duration:** \_\_\_\_\_ days  Unknown

**Maximum O<sub>2</sub> flow volume:**  <2 L/min  2-5 L/min  6-10 L/min  11-15 L/min  >15 L/min

**Non-invasive ventilation? (Any)**  YES  NO  Unknown      **If YES, total duration:** \_\_\_\_\_ days  Unknown

**Invasive ventilation? (Any)**  YES  NO  Unknown      **If YES, total duration:** \_\_\_\_\_ days  Unknown

**Prone Positioning?**  YES  NO  Unknown      **If YES, total duration:** \_\_\_\_\_ days  Unknown

**Inhaled Nitric Oxide?**  YES  NO  Unknown

**Tracheostomy inserted?**  YES  NO  Unknown

**Extracorporeal support (ECMO)?**  YES  NO  Unknown      **If YES, total duration:** \_\_\_\_\_ days  Unknown

**Renal replacement therapy (RRT) or dialysis?**  YES  NO  Unknown

**Inotropes/vasopressors?**  YES  NO  Unknown      **If YES, total duration:** \_\_\_\_\_ days  Unknown

**ICU or High Dependency Unit admission?**  YES  NO  Unknown      **If YES, total duration:** \_\_\_\_\_ days  Unknown

**If YES, date of ICU admission:** [ \_ ] [ \_ ] / [ \_ ] [ \_ ] / [ \_ ] [ \_ ] [ \_ ] [ \_ ]  Unknown

**date of ICU discharge:** [ \_ ] [ \_ ] / [ \_ ] [ \_ ] / [ \_ ] [ \_ ] [ \_ ] [ \_ ]  Unknown

**COMPLICATIONS: At any time during hospitalisation did the patient experience: (Unk = Unknown)**

Viral pneumonia/pneumonitis	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk	Stroke / Cerebrovascular accident	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
Bacterial pneumonia	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk	Meningitis / Encephalitis	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
Acute Respiratory Distress Syndrome	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk	Bacteremia	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
If YES, specify: <input type="radio"/> Mild <input type="radio"/> Moderate <input type="radio"/> Severe <input type="radio"/> Unk		Coagulation disorder / DIC	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
Pneumothorax	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk	Pulmonary embolism	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
Pleural effusion	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk	Anemia	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
Cryptogenic organizing pneumonia (COP)	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk	Rhabdomyolysis / Myositis	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
Bronchiolitis	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk	Acute renal injury/ Acute renal failure	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
Cardiac arrest	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk	Gastrointestinal haemorrhage	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
Myocardial infarction	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk	Pancreatitis	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
Cardiac ischaemia	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk	Liver dysfunction	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
Cardiac arrhythmia	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk	Hyperglycemia	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
Myocarditis / Pericarditis	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk	Hypoglycemia	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk
Endocarditis	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk	Other  If YES specify:	
Cardiomyopathy	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk		
Congestive heart failure	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk		
Seizure	<input type="radio"/> YES <input type="radio"/> NO <input type="radio"/> Unk		

### MODULE 3: OUTCOME CASE REPORT FORM

**DIAGNOSTICS**

**Was patient clinically diagnosed with COVID-19?**  YES  NO  Unknown

**Was pathogen testing done during this illness episode?**  YES (complete section)  NO  Unknown

**Coronavirus:**  Positive  Negative  Not done **If Positive:**  COVID-2019/ SARS-CoV2  MERS CoV  
 Other CoV: \_\_\_\_\_  Unknown

**Influenza :**  Positive  Negative  Not done **If Positive:**  A/H3N2  A/H1N1pdm09  A/H7N9  A/H5N1  A-not typed  B  
 Other: \_\_\_\_\_  Unknown

**RSV:**  Positive  Negative  Not done

**Adenovirus:**  Positive  Negative  Not done

**Bacteria:**  Positive  Negative  Not done **If Positive, specify:** \_\_\_\_\_  Unknown

**Other pathogen/s detected:**  YES  NO  Unknown **If YES, specify all:** \_\_\_\_\_  Unknown

\*\*\*\*\*

**Clinical pneumonia diagnosed?**  YES  NO  Unknown

**Chest X-Ray performed?**  YES  NO  Unknown **If Yes: Were infiltrates present?**  YES  NO  Unknown

**CT performed?**  YES  NO  Unknown **If Yes: Were infiltrates present?**  YES  NO  Unknown

Collection Date (DD/MM/YYYY)	Biospecimen Type	Laboratory test Method	Result	Pathogen Tested/Detected
_D_ / _M_ / 20_Y_Y	<input type="radio"/> Nasal/NP swab <input type="radio"/> Throat swab <input type="radio"/> Combined nasal/NP+throat swab <input type="radio"/> Sputum <input type="radio"/> BAL <input type="radio"/> ETA <input type="radio"/> Urine <input type="radio"/> Feces/rectal swab <input type="radio"/> Blood <input type="radio"/> Other, Specify: _____	<input type="radio"/> PCR <input type="radio"/> Culture <input type="radio"/> Other, Specify: _____	<input type="radio"/> Positive <input type="radio"/> Negative <input type="radio"/> Unknown	_____
_D_ / _M_ / 20_Y_Y	<input type="radio"/> Nasal/NP swab <input type="radio"/> Throat swab <input type="radio"/> Combined nasal/NP+throat swab <input type="radio"/> Sputum <input type="radio"/> BAL <input type="radio"/> ETA <input type="radio"/> Urine <input type="radio"/> Feces/rectal swab <input type="radio"/> Blood <input type="radio"/> Other, Specify: _____	<input type="radio"/> PCR <input type="radio"/> Culture <input type="radio"/> Other, Specify: _____	<input type="radio"/> Positive <input type="radio"/> Negative <input type="radio"/> Unknown	_____
_D_ / _M_ / 20_Y_Y	<input type="radio"/> Nasal/NP swab <input type="radio"/> Throat swab <input type="radio"/> Combined nasal/NP+throat swab <input type="radio"/> Sputum <input type="radio"/> BAL <input type="radio"/> ETA <input type="radio"/> Urine <input type="radio"/> Feces/rectal swab <input type="radio"/> Blood <input type="radio"/> Other, Specify: _____	<input type="radio"/> PCR <input type="radio"/> Culture <input type="radio"/> Other, Specify: _____	<input type="radio"/> Positive <input type="radio"/> Negative <input type="radio"/> Unknown	_____
_D_ / _M_ / 20_Y_Y	<input type="radio"/> Nasal/NP swab <input type="radio"/> Throat swab <input type="radio"/> Combined nasal/NP+throat swab <input type="radio"/> Sputum <input type="radio"/> BAL <input type="radio"/> ETA <input type="radio"/> Urine <input type="radio"/> Faeces/rectal swab <input type="radio"/> Blood <input type="radio"/> Other, Specify: _____	<input type="radio"/> PCR <input type="radio"/> Culture <input type="radio"/> Other, Specify: _____	<input type="radio"/> Positive <input type="radio"/> Negative <input type="radio"/> Unknown	_____
_D_ / _M_ / 20_Y_Y	<input type="radio"/> Nasal/NP swab <input type="radio"/> Throat swab <input type="radio"/> Combined nasal/NP+throat swab <input type="radio"/> Sputum <input type="radio"/> BAL <input type="radio"/> ETA <input type="checkbox"/> Urine <input type="radio"/> Feces/rectal swab <input type="radio"/> Blood <input type="radio"/> Other, Specify: _____	<input type="radio"/> PCR <input type="radio"/> Culture <input type="radio"/> Other, Specify: _____	<input type="radio"/> Positive <input type="radio"/> Negative <input type="radio"/> Unknown	_____



