





ISARIC WHO Clinical Characterisation Protocol for Severe Emerging Infections UK (CCP-UK) CASE REPORT FORMS FRONT PAGE 1 of 3

v9.3 23APR2020

DESIGN OF THE CCP-UK CASE REPORT FORM (CRF)

This CRF is divided into a "ADMISSION" form (5 pages), a "DAILY" form (1 page) for daily clinical and laboratory and data, an "OUTCOME" form (5 pages) and a "WITHDRAWAL" form (1 page).

HOW TO USE THIS CRF

The CRF is designed to complement the **Tier** of activity that a site has capacity and capability to work to. This is likely to vary over the course of an outbreak. The decision on which **Tier** to use is up to the Local Principal Investigator. All high-quality data is valuable for analysis.

Ideally, data and samples will be collected with consent using Tier 2 of the protocol schedule, as outlined below. This will be of greatest public health research value in the early stages of an outbreak.

Data can be collected as Tier Zero activity regardless of consent including retrospectively and from deceased cases.

Consent must be obtained for any biological sampling at Tier 1 and Tier 2 activity.

Tier	For sites where caseload or facilities limit research capacity to deliver Tier 1 or Tier 2 activity.
Zero	OR
	For collection of data without consent from any case; current, past and deceased.
	Please complete the ADMISSION CRF and DAILY CRF for the first day of hospital admission (day 1), the DAILY CRF for the third (d3), sixth (d6) and ninth (d9) days, then the OUTCOME CRF at discharge or death.
Tier 1	For sites where facilities limit research capacity to deliver Tier 2 activity. With consent for single timepoint biological sampling.
	Please complete the ADMISSION CRF and DAILY CRF for the first day of hospital admission (day 1), the DAILY CRF for the third (d3), sixth (d6) and ninth (d9) days, the DAILY CRF again for the first day of any ICU admission, and then the OUTCOME CRF at discharge or death.
Tier 2	For sites with available resources to deliver Tier 2 activity per the protocol schedule. With consent for multiple timepoint biological sampling.
	Please complete the ADMISSION CRF and DAILY CRF on the first day of hospital admission. Please complete the DAILY CRF on each subsequent day up to discharge or death. Please complete the OUTCOME CRF at discharge or death.

On each page above here write site code & participant number as per this example

PARTICIPANT ID I___ | I__ | I___ | I__ | I___ | I__ FRONT PAGE 2 of 3

GENERAL GUIDANCE

CASE REPORT FORMS

- 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 or deceased after admission.
- Participant Identification Numbers consist of a 5-digit CPMS / ODS site code and a 4-digit participant number. You should obtain a site code by contacting your local R&D office or CCP@liverpool.ac.uk. 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.
- CRF data should be entered to the central database at https://ncov.medsci.ox.ac.uk
- REDCap registration access is obtained by contacting CCP.REDCap@liverpool.ac.uk
- Please contact us at CCP.REDCap@liverpool.ac.uk for help with database problems.

RULES DEFINING DAYS

- 1. Day of Admission = Day of Admission regardless, e.g. even if admitted 2 months ago for a broken
- 2. For Community Acquired COVID-19 i.e. admitted with symptoms consistent with COVID-19, day 1 = first 24 hours of admission.
- 3. For those who are already admitted for any other reason and who subsequently test positive, day 1 = day the positive COVID-19 test was collected.
- 4. Rules 2 and 3 are important but we recognise that start of biological sampling for Tier 1 and 2 may be deferred or delayed for several reasons, e.g. due to a delay in the COVID-19 result being reported. If this happens, please take the d1 sample set as soon as possible and then d3 and d9 according to schedule.
- 5. For Tier Zero date of enrolment is date on which data collection started. For Tier 1 & 2 date of enrolment is enrolment = date of consent
- Complete every line of every section, except for where the instructions say to skip a section based on certain responses.
- ➤ Selections with square boxes (□) are single selection answers (choose one answer only). Selections with circles (**O**) are multiple selection answers (choose as many answers as are applicable).
- Some fields are considered URGENT AND ESSENTIAL. These are marked BOLD AND UNDERLINED IN ALL CIRCUMSTANCES PLEASE PRIORITISE THESE DATA POINTS FOR URGENT UPLOAD.
- Mark 'N/K' for any results of laboratory values that are not known or not available.
- Avoid recording data outside of the dedicated areas. Sections are available for recording additional information.
- We recommend writing clearly in black ink, using BLOCK-CAPITAL LETTERS.

CASE REPORT FORMS

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- 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.
- In the case of a participant transferring between study sites, such as to a Nightingale Hospital, or other surge facility, it is preferred to maintain the same Participant Identification Number across the sites. When this is not possible, space for recording the new number is provided.
- > Please keep all of the sheets for a single participant together e.g. with a staple or participant-unique folder.
- These three FRONT PAGES do not need to be retained.
- > DO NOT SEND CRFs to anyone by email or post.
- > See the training guide on how to send consent to CCP@liverpool.ac.uk using [SECURE] encryption
- The Dalhousie University Clinical Frailty Score is provided below for your reference.

Clinical Frailty Scale*



I Very Fit – People who are robust, active, energetic and motivated. These people commonly exercise regularly. They are among the fittest for their age.



2 Well – People who have no active disease symptoms but are less fit than category I. Often, they exercise or are very active occasionally, e.g. seasonally.



3 Managing Well – People whose medical problems are well controlled, but are not regularly active beyond routine walking.



4 Vulnerable – While not dependent on others for daily help, often symptoms limit activities. A common complaint is being "slowed up", and/or being tired during the day.



5 Mildly Frail — These people often have more evident slowing, and need help in high order IADLs (finances, transportation, heavy housework, medications). Typically, mild frailty progressively impairs shopping and walking outside alone, meal preparation and housework.



6 Moderately Frail – People need help with all outside activities and with keeping house. Inside, they often have problems with stairs and need help with bathing and might need minimal assistance (cuing, standby) with dressing.



7 Severely Frail – Completely dependent for personal care, from whatever cause (physical or cognitive). Even so, they seem stable and not at high risk of dying (within ~ 6 months).



8 Very Severely Frail – Completely dependent, approaching the end of life. Typically, they could not recover even from a minor illness.



9. Terminally III - Approaching the end of life. This category applies to people with a life expectancy <6 months, who are not otherwise evidently frail.</p>

Scoring frailty in people with dementia

The degree of frailty corresponds to the degree of dementia. Common **symptoms in mild dementia** include forgetting the details of a recent event, though still remembering the event itself, repeating the same question/story and social withdrawal.

In moderate dementia, recent memory is very impaired, even though they seemingly can remember their past life events well. They can do personal care with prompting.

In severe dementia, they cannot do personal care without help.

- * I. Canadian Study on Health & Aging, Revised 2008.
- 2. K. Rockwood et al. A global clinical measure of fitness and frailty in elderly people. CMAJ 2005;173:489-495.

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	PARTICIPANT ID I	11	1.1	1.1	1.1	l l	1.1	1.1	1.1	- 1
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Date of enrolment [_D_](_D_]/[_M_](_M_]/[_2_](_0_](_Y_](_Y_) Site Location	
For Tier Zero date of enrolment is date on which data collection started. For Tier 1 & 2 enrolment = date of consent	
CLINICAL INCLUSION CRITERIA	
Proven or high likelihood of infection with pathogen of Public Health Interest: YES NC)
OR	
Experience of the following symptoms during this illness episode: (one or more required for	inclusion)
History of self-reported feverishness or measured fever of ≥ 38°C: Cough:	☐ YES ☐ NO ☐ YES ☐ NO
Dyspnoea (shortness of breath) OR Tachypnoea*:	☐ YES ☐ NO
Clinical suspicion of Acute Respiratory Infection despite not meeting criteria above:	☐ YES ☐ NO
* respiratory rate ≥50 breaths/min for <1 year; ≥40 breaths/min for 1-4 years; ≥30 breaths/min for 5-for ≥13 years	12 years; ≥20 breaths/min
DEMOGRAPHICS	
Sex at Birth: Male Female Not specified *Date of birth [D][D]/[M][M]/[Y][<u> </u>
If date of birth is Not Known (N/K) record Age: [][]years OR [][]months	
Postcode: [][][] [][]	
England & Wales NHS, Scotland CHI, [][] [] [] [] [] [] []	
Ethnic group (check all that apply):	
OArab OBlack OEast Asian OSouth Asian OWest Asian OLatin American OWhite OAb	original/First Nations
OOther:	
Employed as a Healthcare Worker? □YES □NO □N/K	
Pregnant? ☐ YES ☐ NO ☐ N/K If YES: Gestational weeks assessment: [][] weeks	
POST PARTUM (within six weeks of delivery)? YES NO or N/K (skip this section - go to INFAI	NT)
Pregnancy Outcome: ☐Live birth ☐Still birth Delivery date: [_D_][_D_]/[_M_][_M_]/[_2	!_][_0_][_Y_][_Y_]
Has infant(s) been tested for Mother's infection? ☐YES ☐NO ☐N/K If YES: ☐Positive ☐Negative	ve
IF POSITIVE PLEASE COMPLETE A SEPARATE CASE REPORT FORM FOR THE INFANT(s)	
INFANT – Less than 1 year old? ☐YES ☐NO (skip this section) Birth weight: [].[]kg ☐N,	/K
Gestational: ☐ Term birth (≥37wk GA) ☐ Preterm birth (<37wk GA) if <37wk Estimated gestation	onweeks \square N/K
Breastfed? ☐YES ☐NO ☐N/K If YES: ☐Currently breastfed ☐Breastfeeding discontinued	□n/k



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(Ageusia)

ADMISSION FORM

ONSET AND ADMISSION									
Symptom onset date of first/earliest symptom: [_D_][_D_]/[_M_][_M_]/[_2_][_0_][_Y_][_Y_]									
Admission date at this facility:	[_D_][_D_]/[_M_][_M_]/[_2	:_][_0_][_Y_][_Y_]							
Is the nationt being readmitted	with Covid-192 TVES TN	io □n/k							
Is the patient being readmitted with Covid-19? □YES □NO □N/K									
Please provide reason for readmission:									
Is this a NIGHTINGALE or other S	SURGE FACILITY TYES THE	D □N/K							
Transfer from other facility? \Box Y	ES-other facility is a study sit	e □YES-other facility is not a study	site □NO □N/K						
If YES: Name of transfer facility:		□n/k							
If YES: Admission date at pre	vious facility (DD/MM/YYYY)	: [_D_][_D_]/[_M_][_M_]/[_2_][_0][_Y_][_Y_]						
If YES-Study Site: Participant I	D # at previous facility: I_I	I_II_II_II_I I_II_	II_II_I						
OR □Same as above									
VITAL SIGNS AT HOSPITAL A	DMISSION (first available o	lata at presentation/Admission – w	ithin 24 hours)						
			·						
		per minute RR: [][]breath							
Sternal capillary refill time >2se		nmHg Severe dehydration: □YES	шио шиук						
Oxygen saturation: [][][_									
	. 1/~	.,, -,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,							
Ad., 1 1 1									
acute illness)	otoms (observed/report	ed at admission and associate	ed with this episode of						
History of fever	□YES □NO □N/K	Lower cheet wall indrawing	□YES □NO □ N/K						
Cough	□YES □NO □N/K	Lower chest wall indrawing Headache	□YES □NO □ N/K						
with sputum production	□YES □NO □ N/K	Altered consciousness/confusion	□YES □NO □ N/K						
bloody sputum/haemoptysis	□YES □NO □ N/K	Seizures	□YES □NO □ N/K						
Sore throat	□YES □NO □ N/K	Abdominal pain	□YES □NO □ N/K						
Runny nose (Rhinorrhoea)	□YES □NO □ N/K	Vomiting / Nausea	□YES □NO □ N/K						
Ear pain	□YES □NO □ N/K	Diarrhoea	□YES □NO □ N/K						
Wheezing	□YES □NO □ N/K	Conjunctivitis	□YES □NO □ N/K						
Chest pain	□YES □NO □ N/K	Skin rash	□YES □NO □ N/K						
Muscle aches (Myalgia)	□YES □NO □ N/K	Skin ulcers	□YES □NO □ N/K						
Joint pain (Arthralgia)	□YES □NO □ N/K	Lymphadenopathy	□YES □NO □ N/K						
Fatigue / Malaise	□YES □NO □ N/K	Bleeding (Haemorrhage)	□YES □NO □ N/K						
Shortness of breath (Dyspnoea)	□YES □NO □ N/K	If Bleeding: specify site(s):							
Disturbance or loss of taste	LIYES LINO LIN/K	Disturbance or loss of small	□ □ □ □ N/K						

(Anosmia)



CO-MORBIDITIES (existing prior to admission)

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Chronic cardiac disease, including congenital heart disease. (not hypertension)	□YES	□no	□n/k	Obesity (as defined by clinical staff)	□YES	□no	□n/ĸ
Hypertension (physician diagnosed)	□YES	□no	□n/k	Diabetes and Type	□no	1 1	2 □n/k
Chronic pulmonary disease (not asthma)	□YES	□по	□n/k	Diabetes (any) with complications	□YES	□по	□n/ĸ
Asthma (physician diagnosed)	□YES	□по	□n/k	Diabetes (any) without complications	□YES	□no	□n/ĸ
Chronic kidney disease	□YES	□по	□n/k	Rheumatologic disorder	□YES	□по	□N/K
Moderate / severe liver disease	□YES	□по	□n/k	Dementia	□YES	□по	□N/K
Mild liver disease	□YES	□по	□n/k	Malnutrition	□YES	□по	□n/k
Chronic neurological disorder	□YES	□по	□n/ĸ	Smoking □YES □Never smoked □	Former s	moker	□п/к
Malignant neoplasm	□YES	□по	□n/k	Other relevant risk factor			
Chronic hematologic disease	□YES	□по	□n/k	□YES □NO □N/K			
AIDS / HIV	□YES	□по	□n/k	If yes, specify			
Is the patient thought to be	a meml	ber of	a CLINIC	ALLY EXTREMELY VULNERABLE GR	OUP		
Solid organ transplant recipients:	□YES	□no	□n/k				
People with specific cancers: \square Y	FS □N	о Пи	/ĸ				
people with cancer who are			-	therapy			
 people with lung cancer wh 	_	_					
people with cancers of the treatment	blood or	bone m	narrow suc	h as leukaemia, lymphoma or myeloma w	ho are at	: any sta	ge of
 people having immunother 	apy or ot	her cor	ntinuing an	tibody treatments for cancer			
 people having other targets or PARP inhibitors 	ed cance	rtreatm	nents whic	h can affect the immune system, such as I	orotein ki	nase inh	nibitors
 people who have had bone marrow or stem cell transplants in the last 6 months, or who are still taking immunosuppression drugs 							
People with <u>severe</u> respiratory conditions including all cystic fibrosis, severe asthma requiring daily oral steroid or injectable maintenance therapy and severe chronic obstructive pulmonary requiring oxygen (COPD):							
People with rare diseases and inborn errors of metabolism that significantly increase the risk of infections (such as Severe combined immunodeficiency (SCID), homozygous sickle cell):							
People on immunosuppression the	rapies su	ıfficient	to signific	antly increase risk of infection: ☐YES	□no [∃n/κ	
Women who are pregnant with sig	nificant l	neart di	sease, con	genital or acquired: □YES □NO □N	I/K		



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PRE-ADMISSION MEDICATION Were any of the following taken within 14 days of admission?

CLINICAL FRAILTY SC With reference to th Clinical Frailty Score CURRENT MEDICATION	ness episode prior N/K If yes, specif ORE e Dalhousie Un	TYES NO N/K r to admission? y: niversity Clinical Fra	Angiotensin converting enzyme inhibitors (ACEI)? YES NO N/K Angiotensin II receptor blockers (ARBs)? YES NO N/K Non-steroidal anti-inflammatory (NSAID)? YES NO N/K ailty Score (see guidance page 3) value 1 to 9 or N/K				
Medication name (generic name preferred)	Dose & unit Dose & unit microgram milligram gram gram int. unit	Dose Frequency □q.d - once a day □b □t.i.d - three times a □q.h.s - before bed □	day □q.i.d - four times a day □5X a day - five times a day ours □ q.6h - every six hours day □prn - as needed ecify	Route of administration ON Oral Oinhaled Oother ON/K Specify Other:			
	(specify)	\square q.h.s - before bed [day □q.i.d - four times a day □5X a day - five times a day ours □ q.6h - every six hours day □prn - as needed ecify	□IV □oral □inhaled □other □N/K Specify Other:			
	□microgram □milligram □gram □int. unit □other (specify)	\square q.h.s - before bed [day □q.i.d - four times a day □5X a day - five times a day ours □ q.6h - every six hours day □prn - as needed ecify	□IV □oral □inhaled □other □N/K Specify Other:			

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ADMISSION FORM

CURRENT MEDICATION to		SION CONTINUED Irrently taking or has taken within the past 14 day	/ S
Medication name (generic name preferred)	Dose & unit	Dose Frequency	Route of administration
	□microgram □milligram □gram □int. unit □other (specify)	□q.d - once a day □b.i.d - twice a day □t.i.d - three times a day □q.i.d - four times a day □q.h.s - before bed □5X a day - five times a day □ q.4h - every four hours □ q.6h - every six hours □q.o.d - every other day □prn - as needed □Other frequency Specify Other:	□IV □oral □inhaled □other □N/K Specify Other:
	□microgram □milligram □gram □int. unit □other (specify)	□q.d - once a day □b.i.d - twice a day □t.i.d - three times a day □q.i.d - four times a day □q.h.s - before bed □5X a day - five times a day □ q.4h - every four hours □ q.6h - every six hours □q.o.d - every other day □prn - as needed □Other frequency Specify Other:	□IV □oral □inhaled □other □N/K Specify Other:
	□microgram □milligram □gram □int. unit □other (specify)	□q.d - once a day □b.i.d - twice a day □t.i.d - three times a day □q.i.d - four times a day □q.h.s - before bed □5X a day - five times a day □ q.4h - every four hours □ q.6h - every six hours □q.o.d - every other day □prn - as needed □Other frequency Specify Other:	□IV □oral □inhaled □other □N/K Specify Other:
	□microgram □milligram □gram □int. unit □other (specify)	□q.d - once a day □b.i.d - twice a day □t.i.d - three times a day □q.i.d - four times a day □q.h.s - before bed □5X a day - five times a day □ q.4h - every four hours □ q.6h - every six hours □q.o.d - every other day □prn - as needed □Other frequency Specify Other:	□IV □oral □inhaled □other □N/K Specify Other:



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ISARIC WHO Clinical Characterisation Protocol for Severe Emerging Infections UK DAILY FORM complete per Tier of activity AND if research samples are collected Page 1 of 1

DAILY TREATMENT (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/K'):
Is the patient in a high level care area i.e. admitted to ICU/ITU/IMC/HDU ☐YES ☐NO ☐N/K
Highest Temperature: [][_]•[_] □°C
Any Supplemental Oxygen See NO N/K FiO2 (0.21-1.0) [].[] or [][]% or [][] L/min (highest)
Oxygen saturation TYES TNO N/K SpO ₂ [][][]% (lowest)
AVPU Alert[] Verbal[] Pain [] Unresponsive[] or $\square N/K$ Glasgow Coma Score (GCS / 15) [][] or $\square N/K$
Is the patient currently receiving, or has received (from 00:00 to 24:00) on day of assessment: Non-invasive respiratory support (e.g. NIV, BIPAP, CPAP)? YES NO N/K Invasive ventilation? YES NO N/K High-flow nasal canula oxygen therapy (>2L/min)? YES NO N/K
DAILY LABORATORY RESULTS
Record the values of laboratory results taken between 00:00 to 24:00 on day of assessment (if Not Available write 'N/K, if multiple record the values for the blood draw taken closest to midday'):
Done □YES □NO □N/K Haemoglobin □g/L <i>or</i> □g/dL
Done One
Done Output Done Of Section O
Done Output Done Of Section O
Done One OYES ONO ON/K Platelets Ox109/L or Ox103/μL Done OYES ONO ON/K APTT/APTR
Done □YES □NO □N/K PT seconds <i>or</i> Done □YES □NO □N/K INR
Done □YES □NO □N/K ALT/SGPT U/L
Done □YES □NO □N/K Total Bilirubin□μmol/L <i>or</i> □mg/dL
Done □YES □NO □N/K AST/SGOT U/L
Done □YES □NO □N/K Glucose □mmol/L <i>or</i> □mg/dL
Done □YES □NO □N/K Blood Urea Nitrogen (urea) □mmol/L <i>or</i> □mg/dL
Done □YES □NO □N/K Lactate□mmol/L <i>or</i> □mg/dL
Done □YES □NO □N/K LDH [][].[]_U/L
Done TYES NO N/K Creatinine Kinase (CPK) [][].[]_U/L
Done \square YES \square NO \square N/K Creatinine $_____$ \square μ mol/L or \square mg/dL
Done □YES □NO □N/K Sodium □mmol/L <i>or</i> □mEq/L
Done □YES □NO □N/K Potassium □mmol/L <i>or</i> □mEq/L
Done ☐YES ☐NO ☐N/K Procalcitonin [][].[]ng/mL
Done ☐YES ☐NO ☐N/K CRP [][].[] mg/L
Most recent HbA1c N/K
Chest X-Ray /CT performed? ☐YES ☐NO ☐N/K IF Yes: Were infiltrates present? ☐YES ☐NO ☐N/K
DESEADOU SAMDI ES
Where biological samples have been taken for research KIT NUMBER [C] [C] [P] [] [] [] []
venicre viological samples have been taken for research

please record the KIT number here.



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PATHOGEN TESTING									
Was pathogen testing done during this illness episode? \ YES \ NO \ N/K \ Influenza : \ YES \ NO \ If YES: \ A/H3N2 \ A/H1N1pdm09 \ A/H7N9 \ A/H5N1 \ \ \ A not typed, other A \ \ B not typed \ Other type (specify): \ Coronavirus: \ YES \ NO \ If YES: \ COVID-19/SARS-COV-2 2019 \ \ \ Other CoV (specify): \ RSV: \ YES \ NO \ Adenovirus: \ YES \ NO \ Bacteria: \ YES : specify : \ list all below \ No \ Other : \ YES \ NO \ If yes Other, specify \									
Collection Date (DD/MM/YYYY)	Bio specimen Type	Laboratory Test Method	Result	Pathogen Detected					
//20	□ Nasal/NP swab □ Throat swab □ Combined nasal/NP+throat swab □ Sputum □ BAL □ ETA □ Urine □ Feces/rectal swab □ Blood □ Other, Specify:	□PCR □Culture □Other, Specify:	□Positive □Negative □N/K						
//20	□ Nasal/NP swab □ Throat swab □ Combined nasal/NP+throat swab □ Sputum □ BAL □ ETA □ Urine □ Feces/rectal swab □ Blood □ Other, Specify:	□PCR □Culture □Other, Specify:	□Positive □Negative □N/K						
//20	□ Nasal/NP swab □ Throat swab □ Combined nasal/NP+throat swab □ Sputum □ BAL □ ETA □ Urine □ Feces/rectal swab □ Blood □ Other, Specify: □ □ Nasal □	□PCR □Culture □Other, Specify:	□Positive □Negative □N/K						



PARTICIPANT ID I	1.1	1.1	1.1	1.1	I I	1.1	1.1	1.1	- 1
IANTICHANTIDI									

OUTCOME FORM Page 2 of 4

MEDICATION: While hospitalised or at discharge, were any of the following administered?						
Antiviral agent? □YES □NO □ N/K If YES	S, tick all the apply: O	Ribavirin OLopinavir/Ritonavir OInterferon alpha				
OInterferon beta OChloroquine / Hydro	xychloroquine O Rer	ndesivir				
ONeuraminidase inhibitor if YES: Which OOther antiviral						
Antibiotic?						
Corticosteroid? □YES □NO □N/K If YES,						
<u> </u>	nouse. Oral initiate					
If YES, please provide type and maximum da	aily dose:	-				
Antifungal agent? □YES □NO □N/K If YE	S: which					
Off-label / Compassionate Use medications?						
TREATMENT: At ANY time during hospitalisation, did the patient receive/undergo:						
ICU or High Dependency Unit admission? ☐YES ☐NO ☐N/K If YES, total duration:days						
Date of ICU/HDU admission: D] D] D] M] M] D 2] D] N M] D 1 D N M D N						
ICU/HDU discharge date: [_D_][_D_]/[_M_][_M_]/[_2_][_0_][_Y_][_Y_] □N/K						
Any Oxygen therapy? ☐YES ☐NO ☐N/K High Flow Oxygen therapy? (>2I/min) ☐YES ☐NO ☐N/K						
Non-invasive ventilation? (e.g. BIPAP, CPAP) □YES □NO □N/K						
Invasive ventilation (Any intubation)?	□YES □NO □N/K	If YES, total duration:days				
Prone Ventilation?	□YES □NO □N/K					
Inhaled Nitric Oxide?	□YES □NO □N/K					
Tracheostomy inserted?	□YES □NO □N/K					
Extracorporeal (ECMO) support?	□YES □NO □N/K	If YES, total duration:days				
Renal replacement therapy (RRT) or dialysis?	□YES □NO □N/K					
Inotropes/vasopressors?	□YES □NO □N/K	If YES, total duration:days				
Blood Group (please check past as well as cu	rrent medical record)	oA oB oAB oO oN/K				



OUTCOME FORM

Myocarditis/Pericarditis

Cardiomyopathy

	PARTICIPANT ID I	1.1	1.1	1.1	1.1	l l	1.1	1.1	1.1	- 1
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□YES □NO □N/K

 COMPLICATIONS: At any time during hospitalisation did the patient experience:

 Viral pneumonia
 YES NO N/K
 Cardiac arrhythmia
 YES NO N/K

 Bacterial pneumonia
 YES NO N/K
 Cardiac ischemia
 YES NO N/K

 Acute Respiratory Distress Syndrome
 YES NO N/K
 Cardiac arrest
 YES NO N/K

 Cryptogenic organizing pneumonia (COP)
 YES NO N/K
 Bacteraemia

'			
Cryptogenic organizing pneumonia (COP)	□YES □NO □N/K	Bacteraemia	□YES □NO □N/K
Pneumothorax	□YES □NO □N/K	Coagulation disorder / Disseminated Intravascular Coagulation	□YES □NO □N/K
Pleural effusion	□YES □NO □N/K	Anaemia	□YES □NO □N/K
Bronchiolitis	□YES □NO □N/K	Rhabdomyolysis / Myositis	□YES □NO □N/K
Meningitis / Encephalitis	□YES □NO □N/K	Acute renal injury/acute renal failure	□YES □NO □N/K
Seizure	□YES □NO □N/K	Gastrointestinal haemorrhage	□yes □no □n/k
Stroke / Cerebrovascular accident	□YES □NO □N/K	Pancreatitis	□YES □NO □N/K
Other neurological complication	□YES □NO □N/K	Liver dysfunction	□YES □NO □N/K
Congestive heart failure	□YES □NO □N/K	Hyperglycaemia	□YES □NO □N/K
Endocarditis	□YES □NO □N/K	Hypoglycaemia	□YES □NO □N/K

□YES □NO □N/K

□YES □NO □N/K

Other, if yes specify below

Other

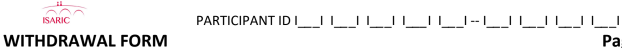
STUDY PARTICIPATION
Is / Has the participant being/ been recruited to a trial or multi-centre study during the period of their current illness (including
initiation in the community and hospital)? \square YES \square NO
IF YES , specify
Name of study
Study Participant ID
Add another study? ☐ YES ☐ NO
IF YES , specify
Name of study
Study Participant ID
Add another study? ☐ YES ☐ NO
IF YES , specify
Name of study
Study Participant ID



	PARTICIPANT ID I	1.1	1.1	1.1	1.1	l l	1.1	1.1	1.1	ı
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OUTCOME FORM Page 4 of 4

OUTCOME									
Outcome: Discharged alive	Outcome: Discharged alive expected to survive								
☐ Hospitalisation	= Remains i	n Hospital ≥ Day 28 after sym _l	otom onset						
- if so	☐ Ongoi	ng health care needs relating	to this admission for C	COVID-19					
	OR								
☐ Medically fit for discharge (COVID-19 resolved) but remains in hospital for other reason (e.g. awaiting suitable care in community, resident in long term health care or mental health facility)									
☐ Transfer to oth	er facility	☐ Palliative discharge	☐ Death	□ n/ K					
Outcome date: [_D_][_D_]/[_	_M_][_M_]/[_2_][_0_][_Y_][_Y_]							
If Discharged alive:									
Ability to self-care at discharge versus before illness: ☐ Same as before illness ☐ Worse ☐ Better ☐ N/K									
If Discharged alive: Post-disc Oxygen t	_	ent: ES □ NO □ N/K							
If Transferred: Facility name: \ _ N/K									
If Transferred: Is the transfe	er facility a stu	dy site? ☐ YES ☐ NO ☐ N/K							
If a Study Site: Participant II	D # at new fac	ility: 🗆 Same as above							
☐ Different: [][][_][][]-	[][][] □n/k							



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WITHDRAWAL
Date of withdrawal: $\[\] \] \[\] \[\$
Type of withdrawal: ☐ Withdrawal from samples only ☐ Other Please specify:
Reason for withdrawal: