

NOVEL CORONAVIRUS (nCoV)

ACUTE RESPIRATORY INFECTION CLINICAL CHARACTERISATION DATA TOOL

DESIGN OF THIS CASE RECORD FORM (CRF)

This CRF is divided into a "CORE" form and a "DAILY" form for daily laboratory and clinical data.

Complete the CORE CRF + complete the DAILY CRF on the first day of hospital admission and on ICU admission, and daily upto 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 3 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.
 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.
- Data should be 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 for later transfer of the data onto the electronic database.
- In the case of a participant transferring between sites, 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.
- 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 (○) are multiple selection answers (choose as many answers as are applicable).
- Mark 'N/A' 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 with patient identifiable information to us via e-mail or post. All data should be transferred to the secure
 electronic database.
- Please enter data on the electronic data capture system at https://redcap.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 <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.





PARTICIPANT IDENTIFICATION #: [Ш	1[][1[II 1	[

☐ YES ☐ NO ☐ Unknown

CORE CASE RECORD FORM

CLINICAL INCLUSION CRITERIA		
Suspected or proven acute novel Coronavirus (nCoV) infection as main cause for admission:	□ YES [□ №
EPIDEMIOLOGICAL FACTORS		

*	Close	contact'	is	defined	as:

while that patient was symptomatic

- Health care associated exposure, including providing direct care for novel coronavirus patients, e.g. health care worker, working with health care workers infected with novel coronavirus, visiting patients or staying in the same close environment of a novel coronavirus patient, or direct exposure to body fluids or specimens including aerosols.

Direct contact with animals in countries where the nCoV is known to be circulating in animal populations or where

- Working together in close proximity or sharing the same classroom environment with a novel coronavirus patient.
- Traveling together with novel coronavirus patient in any kind of conveyance.

In the 14 days before onset of illness had the patient any of the following:

Presence in a healthcare facility where nCoV infections have been managed

human infections have occurred as a result of presumed zoonotic transmission

Presence in a laboratory handling suspected or confirmed nCoV samples

A history of travel to an area with documented cases of nCoV infection

Close contact* with a confirmed or probable case of nCoV infection,

- Living in the same household as a novel coronavirus patient.





PARTICIPANT IDENTIFICATION #:	11 1	П][11	П	1[

DEMOGRAPHICS
Clinical centre name:Country:
Enrolment date: [_D_][_D_]/[_M_][_M_]/[_2_][_0_][_Y_][_Y_]
Ethnic group (check all that apply): OArab OBlack OEast Asian OSouth Asian O West Asian O Latin American O White
O Aboriginal/First Nations O Other:
Employed as a Healthcare Worker? □YES □NO □N/A
Employed in a microbiology laboratory? □YES □NO □N/A
Sex at Birth: ☐ Male ☐ Female ☐ Not specified
Estimated Age [][]years OR][]months
Pregnant? ☐ YES ☐ NO ☐ Unknown ☐ N/A If YES: Gestational weeks assessment: [][] weeks
POST PARTUM? □YES □NO □ N/A (if NO or N/A skip this section - go to INFANT)
Pregnancy Outcome: ☐Live birth ☐Still birth ☐Still birth ☐Delivery date: ☐☐☐☐☐☐☐☐☐☐☐☐☐☐☐☐☐☐☐☐☐☐☐☐☐☐☐☐☐☐☐☐☐☐☐☐
Baby tested for Mother's ARI infection? □YES □NO □N/A If YES: □Positive □Negative Method: □PCR □Other:
INFANT – Less than 1 year old? □YES □NO (If NO skip this section)
Birth weight: [].[]□kg or □lbs □N/A
Gestational outcome: ☐ Term birth (≥37wk GA) ☐ Preterm birth (<37wk GA) ☐ N/A
Breastfed? ☐YES ☐NO ☐N/A If YES: ☐Currently breastfed ☐Breastfeeding discontinued at [][]weeks☐N/A
Appropriate development for age? □YES □NO □Unknown
Vaccinations appropriate for age/country? □YES □NO □Unknown □N/A





PARTICIPANT IDENTIFICATION #: [__][__]--- [__][__][__]

CO-MORBIDITIES					
Co-morbidities and risk factors – Ch	arlson Index will be calculat	ed for each patient at analysis.			
Chronic cardiac disease, including congenital heart disease (not hypertension)	□YES □NO □N/A	Obesity (as defined by clinical staff)	□YES □NO □N/A		
Chronic pulmonary disease (not asthma)	□YES □NO □N/A	Diabetes with complications	□YES □NO □N/A		
Asthma (physician diagnosed)	□ YES □NO □N/A	Diabetes without complications	□YES □NO □N/A		
Chronic kidney disease	□YES □NO □N/A	Rheumatologic disorder	□YES □NO □N/A		
Moderate or severe liver disease	□YES □NO □N/A	Dementia	□YES □NO □N/A		
Mild liver disease	□YES □NO □N/A	Malnutrition	□YES □NO □N/A		
Chronic neurological disorder	□YES □NO □N/A	Smoking	□YES □Never smoked □Former smoker		
Malignant neoplasm	□YES □NO □N/A	Other relevant risk factor	□YES □NO □N/A		
Chronic hematologic disease	□YES □NO □N/A	If yes, specify:			
AIDS / HIV	□YES □NO □N/A				
ONSET & ADMISSION					
Onset date of first/earliest symptom	: [_D_](_M_](_M_]/[_	2_][_0_][_Y_][_Y_]			
Admission date at this facility: [_D_]	[_D_]/[_M_][_M_]/[_2_][_0][_Y_][_Y_]			
Time of admission (24-hr format):[_b	H_][_H_]/[_M_][_M_]				
Transfer from other facility? ☐YES-fa	acility is a study site	acility is not a study site □NO □N/A			
If YES: Name of transfer facility:		□N/A			
If YES: Admission date at transfer fac	cility (<i>DD/MM/YYYY</i>): [_D_][_D_]/[_M_][_M_]/[_2_][_0_][_Y_][_Y_]	□N/A		
If YES-Study Site: Participant ID # at	transfer facility: □Same as	above □Different: [][]–[_][][] □N/A		
Travel in the 14 days prior to first sy	mptom onset? □YES □	NO □Unknown			
If YES, state location(s) & date(s):	Country:	City/Geographic area:			
Return Date: [_D_][_D_]/[_M_][M_]/[_2_][_0_][_Y_][_Y_]	□ N/A (more space at the end if require	red)		
Contact with animals, raw meat or in	nsect bites in the 14 days pr	or to symptom onset?			
□YES □NO □Unknown □ N/A If YES, complete the ANIMAL EXPOSURE section					





PARTICIPANT IDENTIFICATION #: 1	11	1[][1[1[ll	1

SIGNS AND SYMPTOMS AT HOSPITAL ADMISSION (first available data at presentation/admission)	on – withii	n 24 hoi	urs)
Temperature: [][].[]□□°C or □□°F HR: [][]beats per minute RR	:[][lbreat	hs per minute
		_	•
Systolic BP: [_] [_] mmHg Diastolic BP: [_] [_] mmHg Severe dehydration: □YES	⊔NO L	JUnknov	wn
Sternal capillary refill time >2seconds □YES □NO □Unknown			
Oxygen saturation: [] []% On: □Room air □Oxygen therapy □N/A			
Admission signs and symptoms (observed/reported at admission and associated with this e	pisode of	acute i	Iness)
History of fever	□YES	□NO	□Unknown
Cough	□YES	□NO	□ Unknown
with sputum production	□YES	□NO	□ Unknown
bloody sputum/haemoptysis	□YES	□NO	□ Unknown
Sore throat	□YES	□NO	□ Unknown
Runny nose (Rhinorrhoea)	□YES	□NO	□ Unknown
Ear pain	□YES	□NO	□ Unknown
Wheezing	□YES	□NO	□ Unknown
Chest pain	□YES	□NO	□ Unknown
Muscle aches (Myalgia)	□YES	□NO	□ Unknown
Joint pain (Arthralgia)	□YES	□NO	□ Unknown
Fatigue / Malaise	□YES	□NO	□ Unknown
Shortness of breath (Dyspnea)	□YES	□NO	□ Unknown
Lower chest wall indrawing	□YES	□NO	□ Unknown
Headache	□YES	□NO	□ Unknown
Altered consciousness/confusion	□YES	□NO	□ Unknown
Seizures	□YES	□NO	□ Unknown
Abdominal pain	□YES	□NO	□ Unknown
Vomiting / Nausea	□YES	□NO	□ Unknown
Diarrhoea	□YES	□NO	□ Unknown
Conjunctivitis	□YES	□NO	□ Unknown
Skin rash	□YES	□NO	□ Unknown
Skin ulcers	□YES	□NO	□ Unknown
Lymphadenopathy	□YES	□NO	□ Unknown
Bleeding (Haemorrhage)	□YES	□NO	□ Unknown
If Bleeding: specify site(s):			





PARTICIPANT IDENTIFICATION #: [Ш	1[][Ш	Ш	Ш	- 1
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PATHOGEN TESTING:						
Was pathogen testing do	ne during this illness episode? ☐YES (comp	lete section) NO [□N/A			
Influenza: ☐ YES- Confirmed ☐ YES- Probable ☐ NO If YES: ☐ A/H3N2 ☐ A/H1N1pdm09 ☐ A/H7N9						
□ A/H5N1 □]A, not typed □ B □ Other:					
Coronavirus: ☐ YES- Con	firmed ☐ YES- Probable ☐ NO If YES: ☐	☐ Novel CoV ☐ MERS (CoV			
☐ Other CoV:	:					
RSV: ☐ YES- Co	onfirmed □ YES- Probable □ NO					
Adenovirus: □ YES- Co	onfirmed 🗆 YES- Probable 🗆 NO					
Bacteria: : ☐ Yes — co	onfirmed : □ No					
Other Infectious Respirat	tory diagnosis: YES- Confirmed YES- P	robable 🗆 NO				
If ves Other Infectious Re	espiratory diagnosis, specify:					
				-		
-	ES		infective: U YI			
Collection Date (DD/MM/YYYY)	Biospecimen Type	Laboratory test Method	Result	Pathogen Tested/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, <i>Specify:</i>	□Positive □Negative □N/A			
//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/A			
//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/A			
//20	□Nasal/NP swab □Throat swab □Combined nasal/NP+throat swab □Sputum □BAL □ETA □Urine □Faeces/rectal swab □Blood □Other, Specify:	□PCR □Culture □Other, Specify:	□Positive □Negative □N/A			
//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/A			





PARTICIPANT IDENTIFICATION #: 1	11	Ш][1[11 11	[

DAILY CASE RECORD FORM (complete one form on admission, one form on admission to ICU, and daily up to 14 days or until discharge or death if earlier)

DAILY ASSESSMENT FORM (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'):
Current admission to ICU/ITU/IMC/HDU? □YES □NO □N/A
Record the worst value (within the previous 24 hours (if Not Available write 'N/A')):
Done □YES □NO FiO ₂ (0.21-1.0) [].[] or []L/min
Done □YES □NO SaO₂ [][]%
Done ☐YES ☐NO PaO ₂ at time of FiO ₂ above [][] ☐kPa or ☐mmHg
Done □YES □NO PaO₂ sample type: □ Arterial □ Venous □ Capillary □N/A
Done □YES □NO From same blood gas record as PaO ₂ PCO ₂ □kPa <i>or</i> □mmHg
Done □YES □NO pH
Done MYES NO HCO3mEq/L
Done TYES NO Base excess mmol/L
AVPU Alert [] Verbal[] Pain [] Unresponsive[]
Glasgow Coma Score (GCS / 15) [][]
Done ☐YES ☐NO Richmond Agitation-Sedation Scale (RASS) []
Done □YES □NO Riker Sedation-Agitation Scale (SAS) []
Done YES NO Systolic Blood Pressure [][]mmHg
Done □YES □NO Diastolic Blood Pressure [][]mmHg
Done □YES □NO Mean Arterial Blood Pressure [][]mmHg
Done ☐YES ☐NO Urine flow rate [][][]mL/24 hours ☐ Check if estimated
Is the patient currently receiving, or has received (between 00:00 to 24:00 on day of assessment) (apply to all questions in this
section): Non-invasive ventilation (e.g. BIPAP, CPAP)? ☐ YES ☐ NO ☐ N/A Invasive ventilation? ☐ YES ☐ NO ☐N/A
Extra corporeal life support (ECLS)? YES NO N/A High-flow nasal canula oxygen therapy YES NO N/A
Dialysis/Hemofiltration? □YES □NO □N/A
Any vasopressor/inotropic support? \square YES \square NO (if NO, answer the next 3 questions NO) \square N/A
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: ΥΕS ΠΟ
Dopamine >15μg/k/min OR Epinephrine/Norepinephrine > 0.1μg/kg/min:
Neuromuscular blocking agents? □ YES □ NO □ N/A Inhaled Nitric Oxide? □ YES □ NO □ N/A Tracheostomy inserted? □ YES □ NO □ N/A Prone positioning? □ YES □ NO □ N/A
Other intervention or procedure: YES NO N/A If YES, Specify:





PARTICIPANT IDENTIFICATION #:	11 1	11][11	П	11	

DAILY CASE RECORD FORM (complete one form on admission, one form on admission to ICU, and daily up to 14 days or till discharge or death if earlier)

DAILY 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'):				
Done □YES □NO Haemoglobin □g/L or □g/dL				
Done □YES □NO WBC count □x10 ⁹ /L or □x10 ³ /μL				
Done Output Done Output Do				
Done One Ore Ore Ore Ore Ore Ore Ore Ore Ore Or				
Done YES NO Haematocrit [][]%				
Done □YES □NO Platelets □x10 ⁹ /L or □x10 ³ /μL				
Done TYES TO APTT/APTR				
Done TYES TNO PTseconds				
Done YES NO INR				
Done □YES □NO ALT/SGPT U/L				
Done □YES □NO Total Bilirubin□μmol/L <i>or</i> □mg/dL				
Done □YES □NO AST/SGOT U/L				
Done □YES □NO Glucose□mmol/L or □mg/dL				
Done □YES □NO Blood Urea Nitrogen (urea) □mmol/L or □mg/dL				
Done □YES □NO Lactate□mmol/L or □mg/dL				
Done Output Done Output Do				
Done TYES NO Sodium [][] mEq/L				
Done TYES NO Potassium [][] mEq/L				
Done □YES □NO Procalcitonin [][].[]ng/mL				
Done □YES □NO CRP_[][].[]_mg/L				
Chest X-Ray performed? □YES □NO □N/A				





PARTICIPANT IDENTIFICATION #: [Ш	Ш][Ш	11 11	
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COMPLICATIONS: At any time during hospitalisation did the patient experience:							
Viral pneumonitis	□ YES	□ NO	□N/A	Cardiac arrest	☐ YES	□ №	□N/A
Bacterial pneumonia	☐ YES	□ NO	□N/A	Bacteremia	☐ YES	□NO	□N/A
Acute Respiratory Distress Syndrome	□ YES	□ NO	□N/A	Coagulation disorder / Disseminated Intravascular Coagulation	☐ YES	□ NO	□N/A
IF yes, specify: ☐ Mild ☐ Unkr	☐ Mode nown	erate [□ Severe	Anemia	☐ YES	□ NO	□N/A
Pneumothorax	☐ YES	□ №	□N/A	Rhabdomyolysis / Myositis	☐ YES	□ №	□N/A
Pleural effusion	□ YES	□ №	□N/A	Acute renal injury/ Acute renal failure	☐ YES	□ №	□N/A
Cryptogenic organizing pneumonia (COP)	☐ YES	□ NO	□N/A	Gastrointestinal haemorrhage	☐ YES	□ №	□N/A
Bronchiolitis	□ YES	□NO	□N/A	Pancreatitis	☐ YES	□ №	□N/A
Meningitis / Encephalitis	□ YES	□ №	□N/A	Liver dysfunction	☐ YES	□ №	□N/A
Seizure	□ YES	□ №	□N/A	Hyperglycemia	☐ YES	□ №	□N/A
Stroke / Cerebrovascular accident	□ YES	□ №	□N/A	Hypoglycemia	☐ YES	□ №	□N/A
Congestive heart failure	□ YES	□NO	□N/A	Other	☐ YES	□ №	□N/A
Endocarditis / Myocarditis / Pericarditis	□ YES	□ NO	□N/A	If yes specify:	<u>'</u>		
Cardiac arrhythmia	□ YES	□ NO	□N/A				
Cardiac ischaemia	□ YES	□ №	□N/A				





PARTICIPANT IDENTIFICATION#: [__|[__]--- [__][__]|__]

CORE CASE RECORD FORM

If YES, please provide type and dose: __ Antifungal agent? □YES □NO □N/A

TREATMENT: At ANY time du	ring hospitalisation, did the patient receive/undergo:			
ICU or High Dependency Unit a	dmission? □YES □NO □N/A If YES, total duration:days			
If yes, date of I				
, ,				
	CU discharge: [_D_][_D_]/[_M_][_M_]/[_2_][_0_][_Y_][_Y_]			
Oxygen therapy? □YES □NO [LIN/A			
Non-invasive ventilation? (e.g. E	BIPAP, CPAP) □YES □NO □N/A			
Invasive ventilation (Any)?	□YES □NO □N/A If YES, total duration:days			
Prone Ventilation?	□YES □NO □N/A			
Inhaled Nitric Oxide?	□YES □NO □N/A			
Tracheostomy inserted	□YES □NO □N/A,			
Extracorporeal support?	□YES □NO □N/A			
Renal replacement therapy (RRT) or dialysis? □YES □NO □N/A				
Inotropes/vasopressors?				
If YES: First/Start date: [_D_][_D_]/[_M_][_M_]/[_2_][_0_][_Y_][_Y_]				
Last/End date: $[_D][_D]/[_M][_M]/[_2][_0][_Y][_Y]$ \square N/A				
OTHER intervention or procedure (please specify):				
MEDICATION: While hospitalised or at discharge, were any of the following administered?				
Antiviral agent? ☐YES ☐NO ☐N/A If YES: ☐Ribavirin ☐Lopinavir/Ritonavir ☐Interferon alpha ☐Interferon beta				
□ Neuraminidase inhibitor □Other				
Antibiotic? □YES □NO □N/A				
Corticosteroid? □YES □NO □N	N/A If YES, Route: □Oral □Intravenous □Inhaled			





PARTICIPANT IDENTIFICATION #: [][][_][_][]	[][
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OUTCOME				
Outcome: ☐ Discharged alive ☐ Hospitalization ☐ Transfer to other facility ☐ Death				
☐ Palliative discharge ☐ Unknown				
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/A				
If Discharged alive: Post-discharge treatment: Oxygen therapy? ☐ YES ☐ NO ☐ N/A Dialysis/renal treatment? ☐ YES ☐ NO ☐ N/A Other intervention or procedure? ☐ YES ☐ NO ☐ N/A				
If YES: Specify (multiple permitted):				
If Transferred: Facility name: \ \ \ \ \ \ \ \ \ \ \ \ \ \ \				
If Transferred: Is the transfer facility a study site? ☐ YES ☐ NO ☐ N/A				
If a Study Site: Participant ID# at new facility: ☐ Same as above ☐ Different: [][] — [][] ☐ N/A				





PARTICIPANT IDENTIFICATION#: [__][__]---[__][__][__]

CORE CASE RECORD FORM

TRAVEL. Did the nations travel in the 14 days prior to first symptom enset? If > 1 location 9 data lists						
TRAVEL: Did the patient travel in the 14 days prior to first symptom onset? If > 1 location & date list:						
Country: City/Geogra	phic area:	Return Date (<i>DD/MM/20YY</i>): / /20				
Country: City/Geogra	phic area:	Return Date (<i>DD/MM/20YY</i>)://20				
Country: City/Geogra	phic area:	Return Date (<i>DD/MM/20YY</i>)://20				
ANIMAL EXPOSURES: Did the patient prior to first symptom onset? □YES If YES, specify the animal/insect, type	□NO □N/A If yes, Complete 6					
Bird/Aves (e.g. chickens, turkeys, ducks)	□YES □NO □N/A					
Bat	□YES □NO □N/A					
Livestock (e.g. goats, cattle, camels)	□YES □NO □N/A					
Horse	□YES □NO □N/A					
Hare/ Rabbit	□YES □NO □N/A					
Pigs	□YES □NO □N/A					
Non-human primates	□YES □NO □N/A					
Rodent (e.g. rats, mice, squirrels)	□YES □NO □N/A					
Insect or tick bite (e.g. tick, flea, mosquito)	□YES □NO □N/A					
Reptile / Amphibian	□YES □NO □N/A					
Domestic animals living in his/her home (e.g. cats, dogs, other)	□YES □NO □N/A					
Animal feces or nests	□YES □NO □N/A					
Sick animal or dead animal	□YES □NO □N/A					
Raw animal meat / animal blood	□YES □NO □N/A					
Skinned, dressed or eaten wild game	□YES □NO □N/A					
Visit to live animal market, farm or zoo	□YES □NO □N/A					
Participated in animal surgery or necropsy	□YES □NO □N/A					
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Other animal contacts:

□YES □NO □N/A