		Ontology Identifier				Deprecated Label Deprecated ID	Version Tracking		
Parent Class	Field	identifier	Definition	Guidance	Examples	IMPORTANT: Only labels and/or IDs will be	Label	ID	Description/Gui
	Colour Code Legend	_				deprecated, always with replacement version	Lauci	ID.	dance
	field name in yellow = required					provided. If a term changes in its meaning, a new term will be created.			
	field name in purple = recommended field name in white = optional								
	neid name in white = optional								
	Database Identifiers	GENEPIO:0001122							
Database Identifiers	specimen collector sample ID	GENEPIO:0001123	The user-defined name for the sample.	Store the collector sample ID. If this number is considered identifiable information, provide an alternative ID. Be sure to store the key that maps between the original and alternative IDs for traceability and follow up if necessary. Every collector sample ID from a single submitter must be unique. It can have any format, but we suggest that you make it concise, unique and consistent within your lab.	prov. mpox_1234		1.0.0	1.0.0	1.0.0
Database identifiers	specimen collector sample in	GENEFIO.0001123	The user-ueimed name for the sample.	Store the primary ID of the related specimen previously	prov_mpox_1234		1.0.0	1.0.0	1.0.0
Database Identifiers	Related specimen primary ID	GENEPIO:0001128	The primary ID of a related specimen previously submitted to the repository.	submitted to the National Microbiology Laboratory so that the samples can be linked and tracked through the system.	SR20-12345		1.0.0	1.0.0	1.0.0
				Provide the case identifer. The case ID greatly facilitates linkage between laboratory and epidemiological data. The					
Database Identifiers	case ID	GENEPIO:0100281	The identifier used to specify an epidemiologically detected case of disease.	case ID may be considered identifiable information. Consulthe data steward before sharing.	ABCD1234		1.0.0	1.0.0	1.0.0
Database Identifiers	bioproject accession	GENEPIO:0001136	The INSDC accession number of the BioProject(s) to which the BioSample belongs.	Required if submission is linked to a BioProject. BioProject are an organizing tool that links together raw sequence data, assemblies, and their associated metadata. A valid BioProject accession has prefix PRIN, PRIL OF PRID, e.g., PRINA12345 and is created once at the beginning of a new sequencing project. Your laboratory can have one or many BioProjects.			1.0.0	1.0.0	1.0.0
Database Identifiers	biosample accession	GENEPIO:0001139	The identifier assigned to a BioSample in INSDC archives.	Store the accession returned from the BioSample submission. NCBI BioSamples will have the prefix SAMN, while ENA BioSamples will have the prefix SAMEA.	SAMN14180202		1.0.0	1.0.0	1.0.0
Database identifiers	INSDC sequence read accession	GENEPIO:0101203	The identifier assigned to a sequence in one of the International Nucleotide Sequence Database Collaboration (INSDC) repositories.	Store the accession assigned to the submitted sequence. European Nucleotide Archive (ENA) sequence accessions start with ERR, NCBI-SRA accessions start with SRR, DNA Data Bank of Japan (DDBJ) accessions start with DRR and Genome Sequence Archive (GSA) accessions start with CRR.	SRR123456, ERR123456, DRR123456, CRR123456		3.0.0	1.0.0	1.0.0
Database identifiers	INSDC assembly accession	GENEPIO:0101204	The versioned identifier assigned to an assembly or consensus sequence in one of the International Nucleotide Sequence Database Collaboration (INSDC) repositories.	Store the versioned accession assigned to the submitted sequence e.g. the GenBank accession version.	LZ986655.1		3.0.0	1.0.0	1.0.0
5	and the state of t	0515010 000447	T. 0'0'1'		EDI 101 400400				
Database Identifiers	GISAID accession	GENEPIO:0001147	The GISAID accession number assigned to the sequence.	Store the accession returned from the GISAID submission.	EPI_ISL_436489		1.0.0	1.0.0	1.0.0
Sample collection and processing	Sample collection and processing	GENEPIO:0001150	The name of the agency that collected the original sample.	The name of the sample collector should be written out in full, (with minor exceptions) and be consistent across multiple submissions e.g. Public Health Agency of Canada, Public Health Ontario, BC Centre for Disease Control. The sample collector specified is at the discretion of the data provider (i.e. may be hospital, provincial public health lab, or other).	BC Centre for Disease Control		1.0.0	1.0.0	1.0.0
Sample collection and processing	sample collected by		The email address of the contact responsible for follow-up	The email address can represent a specific individual or lab			1.0.0	1.0.0	1.0.0
Sample collection and processing	sample collector contact email	GENEPIO:0001156	regarding the sample.	e.g. johnnyblogs@lab.ca, or RespLab@lab.ca The mailing address should be in the format: Street numbe	RespLab@lab.ca		1.0.0	1.0.0	1.0.0
Sample collection and processing	sample collector contact address	GENEPIO:0001158	The mailing address of the agency submitting the sample.	and name, City, Province/Territory, Postal Code, Country			1.0.0	1.0.0	1.0.0
Sample collection and processing	sample collection date	GENEPIO:0001174	The date on which the sample was collected.	Sample collection date is critical for surveillance and many types of analyses. Required graunularity includes year, month and day. If this date is considered identifiable information, it is acceptable to add "litter" by adding or subtracting a calendar day (acceptable by GISAID). Alternatively, "received date" may be used as a substitute. The date should be provided in ISO 8601 standard format "YYYY-MM-DISA".	2020-03-1	6	1.0.0	1.0.0	1.0.0
				Provide the precision of granularity to the "day", "month", or "year" for the date provided in the "sample collection date" field. The "sample collection date" will be truncated to the precision specified upon export; "day" for "YYYY-MM-DD",					
Sample collection and processing Sample collection and processing	sample collection date precision sample received date	GENEPIO:0001177 GENEPIO:0001179	The precision to which the "sample collection date" was provided. The date on which the sample was received.	"month" for "YYYY-MM", or "year" for "YYYY". ISO 8601 standard "YYYY-MM-DD".	year 2020-03-2	0	1.0.0	1.0.0	1.0.0
				Provide the country name from the controlled vocabulary					
Sample collection and processing	geo_loc_name (country)	GENEPIO:0001181	The country where the sample was collected.	provided. Provide the province/territory name from the controlled	Canada		1.0.0	1.0.0	1.0.0
Sample collection and processing	geo_loc_name (state/province/territory)	GENEPIO:0001185	The state/province/territory where the sample was collected.	vocabulary provided. Use "Mpox virus". This value is provided in the template.	Saskatchewan		1.0.0	1.0.0	1.0.0
Sample collection and processing	organism	GENEPIO:0001191	Taxonomic name of the organism.	Use "Mpox virus": I his value is provided in the template. Note: the Mpox virus was formerly referred to as the "Monkeypox virus" but the international nomenclature has changed (2022).	Mpox virus		1.0.0	1.0.0	1.0.0
Sample collection and processing	isolate	GENEPIO:0001195	Identifier of the specific isolate.	Provide the GISAID EpiPox virus name, which should be written in the format "hMpxV/Canada/2 digit provincial ISO code-xxxxx/year". If the province code cannot be shared fo privacy reasons, put "UN" for "Unknown".			1.0.0	1.0.0	1.0.0
Sample collection and processing	purpose of sampling	GENEPIO:0001198	The reason that the sample was collected.	As all samples are taken for diagnostic purposes, "Diagnostic Testing" should be chosen from the picklist at this time. The reason why a sample was originally collected may differ from the reason why it was selected for sequencing, which should be indicated in the "purpose of sequencing" field.	Diagnostic testing		1.0.0	1.0.0	1.0.0

				Provide an expanded description of why the sample was collected using free text. The description may include the						
			The description of why the comple was collected providing	importance of the sample for a particular public health investigation/surveillance activity/research question. If	Sumptomology and history suggested					
Sample collection and processing	purpose of sampling details	GENEPIO:0001200	The description of why the sample was collected, providing specific details.	details are not available, provide a null value.	Symptomology and history suggested Monkeypox diagnosis.		1.0	0.0	1.0.0	1.0.0
				This information is required for upload through the CNPHI						
			The type of specimen submitted to the National Microbiology	LaSER system. Select the specimen type from the pick list provided. If sequence data is being submitted rather than a						
Sample collection and processing	NML submitted specimen type	GENEPIO:0001204	Laboratory (NML) for testing.	specimen for testing, select "Not Applicable".	Nucleic Acid		1.0	0.0	1.0.0	1.0.0
				Provide the tag that describes how the previous sample is						
			The relationship of the current specimen to the specimen/sample	related to the current sample being submitted from the pick list provided, so that the samples can be linked and tracked						
Sample collection and processing	Related specimen relationship type	GENEPIO:0001209	previously submitted to the repository.	in the system.	Previously Submitted		1.0	0.0	1.0.0	1.0.0
				Provide a descriptor if an anatomical material was sampled.						
			A substance obtained from an anatomical part of an organism e.g.	Use the picklist provided in the template. If a desired term						
Sample collection and processing	anatomical material	GENEPIO:0001211	tissue, blood.		Lesion (Pustule)		1.0	0.0	1.0.0	1.0.0
				Provide a descriptor if an anatomical part was sampled.						
				Use the picklist provided in the template. If a desired term is missing from the picklist, contact emma_griffiths@sfu.ca.						
Sample collection and processing	anatomical part	GENEPIO:0001214	An anatomical part of an organism e.g. oropharynx.	If not applicable, do not leave blank. Choose a null value.	Genital area		1.0	0.0	1.0.0	1.0.0
				Provide a descriptor if a body product was sampled. Use						
			A substance excreted/secreted from an organism e.g. feces,	the picklist provided in the template. If a desired term is missing from the picklist, contact emma_griffiths@sfu.ca. If						
Sample collection and processing	body product	GENEPIO:0001216	urine, sweat.	not applicable, do not leave blank. Choose a null value.	Pus		1.0	0.0	1.0.0	1.0.0
				Provide a descriptor if a device was used for sampling. Use						
				the picklist provided in the template. If a desired term is						
Sample collection and processing	collection device	GENEPIO:0001234	The instrument or container used to collect the sample e.g. swab.	missing from the picklist, contact emma_griffiths@sfu.ca. If not applicable, do not leave blank. Choose a null value.	Swab		1.0	0.0	1.0.0	1.0.0
. ,			,	Provide a descriptor if a collection method was used for						
				sampling. Use the picklist provided in the template. If a desired term is missing from the picklist contact						
			The process used to collect the sample e.g. phlebotamy,	emma griffiths@sfu.ca. If not applicable, do not leave						
Sample collection and processing	collection method	GENEPIO:0001241	necropsy.	blank. Choose a null value.	Biopsy		1.0	0.0	1.0.0	1.0.0
				Critical for interpreting data. Select all the applicable						
				processes from the pick list. If virus was passaged, include information in "lab host", "passage number", and "passage						
			Any processing applied to the sample during or after receiving the							
Sample collection and processing	specimen processing	GENEPIO:0001253	sample.	apply, put "not applicable".	Specimens pooled		1.0	J.0	1.0.0	1.0.0
			Detailed information regarding the processing applied to a sample	Describe a few hand described of any according datable	5 swabs from different body sites were pooled					
Sample collection and processing	specimen processing details	GENEPIO:0100311	during or after receiving the sample.	applied to a sample.	and further prepared as a single sample during library prep.		1.0	0.0	1.0.0	1.0.0
				Samples can play different types of roles in experiments. A	,,,,					
				sample under study in one experiment may act as a control						
				or be a replicate of another sample in another experiment. This field is used to distinguish samples under study from						
				controls, replicates, etc. If the sample acted as an						
				experimental control or a replicate, select a role type from						
Sample collection and processing	experimental specimen role type	GENEPIO:0100921	The type of role that the sample represents in the experiment.	experimental control or a replicate, select a role type from the picklist. If the sample was not a control, leave blank or select "Not Applicable".	Positive experimental control		3.0).0	3.0.0	3.0.0
				experimental control or a replicate, select a role type from the picklist. If the sample was not a control, leave blank or select "Not Applicable".						
Sample collection and processing	experimental specimen role type experimental control details	GENEPIO:0100921 GENEPIO:0100922	The type of role that the sample represents in the experiment. The details regarding the experimental control contained in the sample represents in the experimental control contained in the sample represents in the experimental control contained in the sample represents in the experimental control contained in the sample represents in the experimental control contained in the sample represents in the experiment.	experimental control or a replicate, select a role type from the picklist. If the sample was not a control, leave blank or select "Not Applicable".		sample as process control	3.0		3.0.0	3.0.0
				experimental control or a replicate, select a role type from the picklist. If the sample was not a control, leave blank or select "Not Applicable".		sample as process control		0.0		
	experimental control details Lineage and Variant information	GENEPIO:0100922 GENEPIO:0001498		experimental control or a replicate, select a role type from the picklist. If the sample was not a control, leave blank or select "Not Applicable".		sample as process control	3.0	0.0	3.0.0	3.0.0
Sample collection and processing	experimental control details Lineage and Variant information Host Information	GENEPIO:0100922 GENEPIO:0001498 GENEPIO:0001268	The details regarding the experimental control contained in the sar	experimental control or a replicate, select a role type from the picklist. If the sample was not a control, leave blank or select "Not Applicable". **Provide details regarding the nature of the reference strain to the ref	i Human coronavirus 229E (HCoV-229E) spiked in	sample as process control	3.0 1.0 1.0	0.0	3.0.0 1.0.0 1.0.0	3.0.0 1.0.0 1.0.0
Sample collection and processing Host Information	experimental control details Lineage and Variant information Host Information host (common name)	GENEPIO:0010922 GENEPIO:0001498 GENEPIO:0001268 GENEPIO:0001386	The details regarding the experimental control contained in the sar The commonly used name of the host.	experimental control or a replicate, select a role type from the picklist. If the sample was not a control, leave blank or select "Not Applicable". Provide details regarding the nature of the reference strain the control of the reference strain that the control of the reference str	i Human coronavirus 229E (HCoV-229E) spiked in	sample as process control	3.0 1.0 1.0	0.0 0.0 0.0 0.0	3.0.0 1.0.0 1.0.0 1.0.0	3.0.0 1.0.0 1.0.0
Sample collection and processing Host Information Host Information	experimental control details Lineage and Variant information Host Information host (common name) host (scientific name)	GENEPIO:0100922 GENEPIO:0001498 GENEPIO:0001268 GENEPIO:0001386 GENEPIO:0001387	The details regarding the experimental control contained in the sar The commonly used name of the host. The taxonomic, or scientific name of the host.	experimental control or a replicate, select a role type from the picklist. If the sample was not a control, leave blank or select "Not Applicable". Provide details regarding the nature of the reference strain to Common name or scientific name are required if there was Common name or scientific name are required if there was	u Human coronavirus 229E (HCoV-229E) spilked in 3 Human 1 Human 1 Homo sapiens	sample as process control	3.0 1.0 1.0 1.0	0.0 0.0 0.0 0.0 0.0	3.0.0 1.0.0 1.0.0 1.0.0	3.0.0 1.0.0 1.0.0 1.0.0
Sample collection and processing Host Information	experimental control details Lineage and Variant information Host Information host (common name)	GENEPIO:0010922 GENEPIO:0001498 GENEPIO:0001268 GENEPIO:0001386	The details regarding the experimental control contained in the sar The commonly used name of the host. The taxonomic, or scientific name of the host. Health status of the host at the time of sample collection.	experimental control or a replicate, select a role type from the picklist. If the sample was not a control, leave blank or select "Not Applicable". Provide details regarding the nature of the reference strain of the refer	i Human coronavirus 229E (HCoV-229E) spiked in	sample as process control	3.0 1.0 1.0	0.0 0.0 0.0 0.0 0.0	3.0.0 1.0.0 1.0.0 1.0.0	3.0.0 1.0.0 1.0.0
Sample collection and processing Host Information Host Information	experimental control details Lineage and Variant information Host Information host (common name) host (scientific name)	GENEPIO:0100922 GENEPIO:0001498 GENEPIO:0001268 GENEPIO:0001386 GENEPIO:0001387	The details regarding the experimental control contained in the sar The commonly used name of the host. The taxonomic, or scientific name of the host.	experimental control or a replicate, select a role type from the picklist. If the sample was not a control, leave blank or select "Not Applicable". Provide details regarding the nature of the reference strain to Common name or scientific name are required if there was Common name or scientific name are required if there was	u Human coronavirus 229E (HCoV-229E) spilked in 3 Human 1 Human 1 Homo sapiens	sample as process control	3.0 1.0 1.0 1.0	0.0 0.0 0.0 0.0 0.0 0.0 0.0	3.0.0 1.0.0 1.0.0 1.0.0	3.0.0 1.0.0 1.0.0 1.0.0
Sample collection and processing Host Information Host Information Host Information	experimental control details Lineage and Variant information Host Information host (common name) host (scientific name) host health state	GENEPIO:0100922 GENEPIO:0001498 GENEPIO:0001268 GENEPIO:0001386 GENEPIO:0001387 GENEPIO:0001388	The details regarding the experimental control contained in the sar The commonly used name of the host. The taxonomic, or scientific name of the host. Health status of the host at the time of sample collection. Further details pertaining to the health or disease status of the	experimental control or a replicate, select a role type from the picklist. If the sample was not a control, leave blank or select "Not Applicable". Provide details regarding the nature of the reference strain to the reference strain is common name or scientific name are required if there was form name or scientific name are required if there was if known, select a value from the pick list. If known, select a descriptor from the pick list provided in	Human coronavirus 229E (HCoV-229E) spilked in il Human il Homo sapiens Asymptomatic	sample as process control	3.0 1.0 1.0 1.0 1.0	0.0 0.0 0.0 0.0 0.0 0.0 0.0	3.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0	3.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0
Sample collection and processing Host Information Host Information Host Information	experimental control details Lineage and Variant information Host Information host (common name) host (scentific name) host health state host health status details	GENEPIO:0100922 GENEPIO:0001498 GENEPIO:0001268 GENEPIO:0001386 GENEPIO:0001387 GENEPIO:0001388 GENEPIO:0001389	The details regarding the experimental control contained in the sar The commonly used name of the host. The taxonomic, or scientific name of the host. Health status of the host at the time of sample collection. Further details pertaining to the health or disease status of the host at time of collection.	experimental control or a replicate, select a role type from the picklist. If the sample was not a control, leave blank or select "Not Applicable". Provide details regarding the nature of the reference strain is common name or scientific name are required if there was. Common name or scientific name are required if there was if known, select a value from the pick list. If known, select a value from the pick list provided in the template. If known, select a value from the pick list provided in the template.	I Human coronavirus 229E (HCoV-229E) spilked in I Human I Human I Homo sapiens Asymptomatic Hospitalized	sample as process control	3.0 1.0 1.0 1.0 1.0	0.0 0.0 0.0 0.0 0.0 0.0 0.0	3.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0	3.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0
Sample collection and processing Host Information Host Information Host Information	experimental control details Lineage and Variant information Host Information host (common name) host (scentific name) host health state host health status details	GENEPIO:0100922 GENEPIO:0001498 GENEPIO:0001268 GENEPIO:0001386 GENEPIO:0001387 GENEPIO:0001388 GENEPIO:0001389	The details regarding the experimental control contained in the sar The commonly used name of the host. The taxonomic, or scientific name of the host. Health status of the host at the time of sample collection. Further details pertaining to the health or disease status of the host at time of collection.	experimental control or a replicate, select a role type from the picklist. If the sample was not a control, leave blank or select "Not Applicable". Provide details regarding the nature of the reference strain of the refer	I Human coronavirus 229E (HCoV-229E) spilked in I Human I Human I Homo sapiens Asymptomatic Hospitalized	sample as process control	3.0 1.0 1.0 1.0 1.0	0.0 0.0 0.0 0.0 0.0 0.0 0.0	3.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0	3.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0
Sample collection and processing Host Information Host Information Host Information	experimental control details Lineage and Variant information Host Information host (common name) host (scentific name) host health state host health status details	GENEPIO:0100922 GENEPIO:0001498 GENEPIO:0001268 GENEPIO:0001386 GENEPIO:0001387 GENEPIO:0001388 GENEPIO:0001389	The details regarding the experimental control contained in the sar The commonly used name of the host. The taxonomic, or scientific name of the host. Health status of the host at the time of sample collection. Further details pertaining to the health or disease status of the host at time of collection.	experimental control or a replicate, select a role type from the picklist. If the sample was not a control, leave blank or select "Not Applicable". Provide details regarding the nature of the reference strain is common name or scientific name are required if there was. Common name or scientific name are required if there was if known, select a value from the pick list. If known, select a value from the pick list provided in the template. If known, select a value from the pick list provided in the template.	I Human coronavirus 229E (HCoV-229E) spilked in I Human I Human I Homo sapiens Asymptomatic Hospitalized	sample as process control	3.0 1.0 1.0 1.0 1.0	0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0	3.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0	3.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0
Sample collection and processing Host Information	experimental control details Lineage and Variant information Host Information host (common name) host (scientific name) host health state host health status details host health outcome	GENEPIO:0010922 GENEPIO:0001498 GENEPIO:0001268 GENEPIO:0001386 GENEPIO:0001387 GENEPIO:0001389 GENEPIO:0001389 GENEPIO:0001389 GENEPIO:0001391	The details regarding the experimental control contained in the sar The commonly used name of the host. The taxonomic, or scientific name of the host. Health status of the host at the time of sample collection. Further details pertaining to the health or disease status of the host at time of collection. Disease outcome in the host. The name of the disease experienced by the host. Age of host at the time of sampling.	experimental control or a replicate, select a role type from the picklist. If the sample was not a control, leave blank or select "Not Applicable". Provide details regarding the nature of the reference strain in the picklist. If the provide details regarding the nature of the reference strain in the picklist. Common name or scientific name are required if there was: If known, select a value from the pick list. If known, select a descriptor from the pick list provided in the template. If known select a value from the pick list provided in the template. Note: the Mpox disease was formerly referred to as "Monkeypox' but the international nomenclature has changed (2022). If known, provide age. Age-binning is also acceptable.	Human coronavirus 229E (HCoV-229E) spilked in a Human la Homo sapiens Asymptomatic Hospitalized Recovered	sample as process control	3.0 1.0 1.0 1.0 1.0 1.0 1.0	0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0	3.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0	3.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0
Sample collection and processing Host Information Host Information Host Information Host Information Host Information Host Information	experimental control details Lineage and Variant information Host Information host (common name) host (scientific name) host health state host health status details host health outcome	GENEPIC.0100922 GENEPIC.0001498 GENEPIC.0001268 GENEPIC.0001386 GENEPIC.0001387 GENEPIC.0001389 GENEPIC.0001389 GENEPIC.0001389	The details regarding the experimental control contained in the sar The commonly used name of the host. The taxonomic, or scientific name of the host. Health status of the host at the time of sample collection. Further details pertaining to the health or disease status of the host at time of collection. Disease outcome in the host. The name of the disease experienced by the host.	experimental control or a replicate, select a role type from the picklist. If the sample was not a control, leave blank or select "Not Applicable". Provide details regarding the nature of the reference strain is common name or scientific name are required if there was. Common name or scientific name are required if there was if known, select a value from the pick list. If known, select a value from the pick list provided in the template. If known, select a value from the pick list. If known, select a value from the pick list. Select "Mook" from the pick list provided in the template. Note: the Mpox disease was formerly referred to as "Monkeypor" but the international nomenclature has changed (2022). If known, provide age. Age-binning is also acceptable. If known, provide the age units used to measure the host's it known, provide the age units used to measure the host's it known, provide the age units used to measure the host's it known, provide the age units used to measure the host's it known, provide the age units used to measure the host's it known, provide the age units used to measure the host's it known, provide the age units used to measure the host's it has the provide the great and the pick list.	Human coronavirus 229E (HCoV-229E) spilked in a Human la Homo sapiens Asymptomatic Hospitalized Recovered	sample as process control	3.0 1.0 1.0 1.0 1.0 1.0 1.0	0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0	3.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0	3.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0
Sample collection and processing Host Information	experimental control details Lineage and Variant information Host Information host (common name) host (scientific name) host health state host health status details host health outcome	GENEPIO:0010922 GENEPIO:0001498 GENEPIO:0001268 GENEPIO:0001386 GENEPIO:0001387 GENEPIO:0001389 GENEPIO:0001389 GENEPIO:0001389 GENEPIO:0001391	The details regarding the experimental control contained in the sar The commonly used name of the host. The taxonomic, or scientific name of the host. Health status of the host at the time of sample collection. Further details pertaining to the health or disease status of the host at time of collection. Disease outcome in the host. The name of the disease experienced by the host. Age of host at the time of sampling.	experimental control or a replicate, select a role type from the picklist. If the sample was not a control, leave blank or select "Not Applicable". Provide details regarding the nature of the reference strain in the picklist. If the provide details regarding the nature of the reference strain in the picklist. Common name or scientific name are required if there was. Common name or scientific name are required if there was. If known, select a value from the pick list. If known, select a descriptor from the pick list provided in the template. If known, select a descriptor from the pick list. Select "Mpox" from the pick list provided in the template. Note: the Mpox disease was formerly referred to as "Monkeypox" but the international nonencitative has changed (2022). If known, provide age. Age-binning is also acceptable. If known, provide the age units used to measure the host's Age bins in 10 year intervals have been provided. If a	Human coronavirus 229E (HCoV-229E) spilked in a Human le Homo sapiens Asymptomatic Hospitalized Recovered Mpox 79 year	sample as process control	3.0 1.0 1.0 1.0 1.0 1.0 1.0	0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0	3.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0	3.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0
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Sample collection and processing Host Information	experimental control details Lineage and Variant information Host Information host (common name) host (scientific name) host health state host health status details host health outcome host disease host age host age host age unit host gender host gender host residence geo_loc name (country) host residence geo_loc name (state/province/territory) symptom onset date signs and symptoms	GENEPIC.0010922 GENEPIC.001498 GENEPIC.0001498 GENEPIC.0001386 GENEPIC.0001387 GENEPIC.0001389 GENEPIC.0001389 GENEPIC.0001389 GENEPIC.0001393 GENEPIC.0001393 GENEPIC.0001396 GENEPIC.0001396 GENEPIC.0001396 GENEPIC.0001396 GENEPIC.0001399 GENEPIC.0001399	The details regarding the experimental control contained in the said the commonly used name of the host. The taxonomic, or scientific name of the host. Health status of the host at the time of sample collection. Further details pertaining to the health or disease status of the host at time of collection to the collection. Disease outcome in the host. The name of the disease experienced by the host. Age of host at the time of sampling. The units used to measure the host's age. The age category of the host at the time of sample collection. The country of residence of the host. The state/province/territory of residence of the host. The state/province/territory of residence of the host. The date on which the symptoms began or were first noted. A perceived change in function or sensation, (loss, disturbance or appearance) indicative of a disease, reported by a patient.	experimental control or a replicate, select a role type from the picklist. If the sample was not a control, leave blank or select "Not Applicable". Provide details regarding the nature of the reference strain is common name or scientific name are required if there was. Common name or scientific name are required if there was. If known, select a value from the pick list provided in the template. If known, select a value from the pick list provided in the template. When the pick list provided in the template. Note: the Mpox disease was formerly referred to as "Monkeypox" but the international nomenciature has changed (2022). If known, provide age, Age-binning is also acceptable. If known, provide the age units used to measure the host's Age bins in 10 year intervals have been provided. If a host's age cannot be specified due to provacy concerns, an age bin can be used as an alternative. If known, provide the most is the provided in the template. If known, provide the age units used to measure the host's Age bins in 10 year intervals have been provided. If a host's age cannot be specified due to provacy concerns, an age bin can be used as an alternative. If known, provide the growner from pick list provided in the temple Select the province/territory name from pick list provided in If known, provide the symptom onset date in ISO 8601 standard format "YYYY-MM-DD". Select all of the symptoms experienced by the host from the pick list.	Human coronavirus 229E (HCoV-229E) spiked in a Human coronavirus 229E (HCoV-229E) spiked in a Human and them sapiens Asymptomatic Hospitalized Recovered Mpox 79 year 79 year 202-05-25 Lesion (Pustule), Swollen Lymph Nodes, Myalgia (muscle pain)		3.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1	0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	1.0.0 1.0.0
Sample collection and processing Host Information	experimental control details Lineage and Variant information Host Information host (common name) host (scientific name) host health state host health status details host health outcome host age host age host age unit host gender host residence geo_loc name (country) host residence geo_loc name (state/province/ferritory) symptom onset date	GENEPIO:0010922 GENEPIO:001493 GENEPIO:0001288 GENEPIO:0001386 GENEPIO:0001389 GENEPIO:0001389 GENEPIO:0001389 GENEPIO:0001393 GENEPIO:0001393 GENEPIO:0001393 GENEPIO:0001393 GENEPIO:0001393 GENEPIO:0001393 GENEPIO:0001393 GENEPIO:0001393	The details regarding the experimental control contained in the sar The commonly used name of the host. The taxonomic, or scientific name of the host. Health status of the host at the time of sample collection. Further details pertaining to the health or disease status of the host at time of collection. Disease outcome in the host. The name of the disease experienced by the host. Age of host at the time of sampling. The units used to measure the host's age. The age category of the host at the time of sampling. The gender of the host at the time of sample collection. The country of residence of the host. The state/province/territory of residence of the host. The date on which the symptoms began or were first noted. A perceived change in function or sensation, (loss, disturbance or	experimental control or a replicate, select a role type from the picklist. If the sample was not a control, leave blank or select "Not Applicable". Trovide details regarding the nature of the reference strain is provided details regarding the nature of the reference strain is common name or scientific name are required if there was. Common name or scientific name are required if there was. If known, select a value from the pick list provided in the template. If known, select a descriptor from the pick list provided in the template. If known, select a descriptor from the pick list provided in the template. Note: the Mpox disease was formerly referred to as "Monkeypor's but the international nomenclature has changed (2022). If known, provide age. Age-binning is also acceptable. If known, provide the age units used to measure the host's a Age bins in 10 year intervals have been provided. If a Age bins in 10 year intervals have been provided. If a Robins in 20 year intervals have been provided. If a Robins in 20 year intervals have been provided. If a Robins in 20 year intervals have been provided. If a Robins in 20 year intervals have been provided in the template in the control of the country name from pick list provided in the template in the country name from pick list provided in the template and the province-ferritory name from pick list provided in the template and the province-ferritory name from pick list provided in the template and and form the pick list. Select all of the pre-existing conditions and risk factors expe	Human coronavirus 229E (HCoV-229E) spiked in a Human coronavirus 229E (HCoV-229E) spiked in a Human and them sapiens Asymptomatic Hospitalized Recovered Mpox 79 year 79 year 202-05-25 Lesion (Pustule), Swollen Lymph Nodes, Myalgia (muscle pain)		3.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1	0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0	3.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0	3.0.0 1.0.0
Sample collection and processing Host Information	experimental control details Lineage and Variant information Host Information host (common name) host (scientific name) host scientific name) host health state host health status details host health outcome host disease host age host age host age unit host sesidence geo_loc name (country) host sesidence geo_loc name (state/province/territory) symptom onset date signs and symptoms pre-existing conditions and risk factors	GENEPIO.0010922 GENEPIO.0001498 GENEPIO.0001268 GENEPIO.0001386 GENEPIO.0001387 GENEPIO.0001389 GENEPIO.0001389 GENEPIO.0001393 GENEPIO.0001393 GENEPIO.0001393 GENEPIO.0001395 GENEPIO.0001396 GENEPIO.0001397 GENEPIO.0001396 GENEPIO.0001397 GENEPIO.0001397 GENEPIO.0001397 GENEPIO.0001397	The details regarding the experimental control contained in the sair The taxonomic, or scientific name of the host. The taxonomic, or scientific name of the host. Health status of the host at the time of sample collection. Further details pertaining to the health or disease status of the host at time of collection. Disease outcome in the host. The name of the disease experienced by the host. Age of host at the time of sampling. The units used to measure the host's age. The age category of the host at the time of sampling. The gender of the host at the time of sample collection. The country of residence of the host. The date on which the symptoms began or were first noted. A perceived change in function or sensation, (loss, disturbance or appearance) indicative of a disease, reported by a patient. Patient pre-existing conditions and risk factors. Si-Pre-existing co	experimental control or a replicate, select a role type from the picklist. If the sample was not a control, leave blank or select "Not Applicable". Provide details regarding the nature of the reference strain is provided details regarding the nature of the reference strain is common name or scientific name are required if there was. Common name or scientific name are required if there was. If known, select a value from the pick list provided in the template. If known, select a descriptor from the pick list provided in the template. If known, select a value from the pick list provided in the template. Note: the Mpox disease was formerly referred to as "Monkeypor's but the international nomenclature has changed (2022). If known, provide age. Age-binning is also acceptable. If known, provide the age units used to measure the host's af Age bins in 10 year intervals have been provided. If a flag bins in 10 year intervals have been provided. If a flag bins in 10 year intervals have been provided. If a flown, provide and the specified due to provary concerns, an age bin can be used as an afternative. If known, provide the symptom onset date in ISO 8601 istandard format "YYYY-MM-DD". Select all of the symptom onset date in ISO 8601 standard format from pick list provided in the templity of the providence of the symptom onset date in ISO 8601 istandard format from pick list the desired term is missing, contact the	a Human coronavirus 229E (HCoV-229E) spilked in a Human la Homo sapiens Asymptomatic Hospitalized Recovered Mpox 79 a year 50 - 59 Male Canada I Quebec 2022-05-25 Lesion (Pustule), Swollen Lymph Nodes, Myaljai (muscle pain) rienced by the host from the pick list. If the desired		3.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1	0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	1.0.0 1.0.0
Sample collection and processing Host Information	experimental control details Lineage and Variant information Host Information host (common name) host (scientific name) host health state host health status details host health outcome host disease host age host age host age unit host gender host gender host residence geo_loc name (country) host residence geo_loc name (state/province/territory) symptom onset date signs and symptoms	GENEPIC.0010922 GENEPIC.001498 GENEPIC.0001498 GENEPIC.0001386 GENEPIC.0001387 GENEPIC.0001389 GENEPIC.0001389 GENEPIC.0001389 GENEPIC.0001393 GENEPIC.0001393 GENEPIC.0001396 GENEPIC.0001396 GENEPIC.0001396 GENEPIC.0001396 GENEPIC.0001399 GENEPIC.0001399	The details regarding the experimental control contained in the said. The taxonomic, or scientific name of the host. The taxonomic, or scientific name of the host. Health status of the host at the time of sample collection. 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Sample collection and processing Host Information	experimental control details Lineage and Variant information Host Information host (common name) host (scientific name) host scientific name) host health state host health status details host health outcome host disease host age host age host age unit host sesidence geo_loc name (country) host sesidence geo_loc name (state/province/territory) symptom onset date signs and symptoms pre-existing conditions and risk factors	GENEPIO.0010922 GENEPIO.0001498 GENEPIO.0001268 GENEPIO.0001386 GENEPIO.0001387 GENEPIO.0001389 GENEPIO.0001389 GENEPIO.0001393 GENEPIO.0001393 GENEPIO.0001393 GENEPIO.0001395 GENEPIO.0001396 GENEPIO.0001397 GENEPIO.0001396 GENEPIO.0001397 GENEPIO.0001397 GENEPIO.0001397 GENEPIO.0001397	The details regarding the experimental control contained in the sail. The taxonomic, or scientific name of the host. Health status of the host at the time of sample collection. Further details pertaining to the health or disease status of the host at time of collection. Disease outcome in the host. The name of the disease experienced by the host. Age of host at the time of sampling. The units used to measure the host's age. The gender of the host at the time of sampling. The gender of the host at the time of sample collection. The country of residence of the host. The date on which the symptoms began or were first noted. A perceived change in function or sensation, floss, disturbance or appearance) indicative of a disease, reported by a patient. Patient pre-existing conditions and risk factors. <i>Pre-existing conditions are risk factors. <i>Pre-existing conditions and risk factors. <i>Pre-existing conditions are risk factors. <i-pre-existing are="" conditions="" factors.<="" risk="" td=""><td>experimental control or a replicate, select a role type from the picklist. If the sample was not a control, leave blank or select "Not Applicable". Provide details regarding the nature of the reference strain is provided to the select of the select and the select of the select and select of the select and select and</td><td>a Human coronavirus 229E (HCoV-229E) spiked in a Human la Homo sapiens Asymptomatic Hospitalized Recovered Mpox 79 year 79 Male 1 Canada 1 Quebec 2022-05-25 Lesion (Pustule), Swollen Lymph Nodes, Mylagia (muscle pain) reneed by the host from the pick list. If the desired Delayed wound healing (lesion healing) Tecovirimat used to treat current Monkeypox</td><td></td><td>3.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1</td><td>0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0</td><td>1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00</td><td>1.0.0 1.0.0</td></i-pre-existing></i></i></i></i></i></i></i></i></i></i></i></i></i></i></i>	experimental control or a replicate, select a role type from the picklist. If the sample was not a control, leave blank or select "Not Applicable". Provide details regarding the nature of the reference strain is provided to the select of the select and the select of the select and select of the select and	a Human coronavirus 229E (HCoV-229E) spiked in a Human la Homo sapiens Asymptomatic Hospitalized Recovered Mpox 79 year 79 Male 1 Canada 1 Quebec 2022-05-25 Lesion (Pustule), Swollen Lymph Nodes, Mylagia (muscle pain) reneed by the host from the pick list. If the desired Delayed wound healing (lesion healing) Tecovirimat used to treat current Monkeypox		3.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1	0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	1.0.0 1.0.0
Sample collection and processing Host Information	experimental control details Lineage and Variant information Host Information host (common name) host (scientific name) host scientific name) host health state host health status details host health outcome host disease host age host age host age unit host sesidence geo_loc name (country) host sesidence geo_loc name (state/province/territory) symptom onset date signs and symptoms pre-existing conditions and risk factors	GENEPIO.0010922 GENEPIO.0001498 GENEPIO.0001268 GENEPIO.0001386 GENEPIO.0001387 GENEPIO.0001389 GENEPIO.0001389 GENEPIO.0001393 GENEPIO.0001393 GENEPIO.0001393 GENEPIO.0001395 GENEPIO.0001396 GENEPIO.0001397 GENEPIO.0001396 GENEPIO.0001397 GENEPIO.0001397 GENEPIO.0001397 GENEPIO.0001397	The details regarding the experimental control contained in the sair The taxonomic, or scientific name of the host. The taxonomic, or scientific name of the host. Health status of the host at the time of sample collection. Further details pertaining to the health or disease status of the host at time of collection. Disease outcome in the host. The name of the disease experienced by the host. Age of host at the time of sampling. The units used to measure the host's age. The age category of the host at the time of sampling. The gender of the host at the time of sample collection. The country of residence of the host. The date on which the symptoms began or were first noted. A perceived change in function or sensation, (loss, disturbance or appearance) indicative of a disease, reported by a patient. Patient pre-existing conditions and risk factors. Si-Pre-existing co	experimental control or a replicate, select a role type from the picklist. If the sample was not a control, leave blank or select "Not Applicable". Provide details regarding the nature of the reference strain is considered to the select of the select and the select and select and select a select and select	a Human coronavirus 229E (HCoV-229E) spiked in a Human and a Human sapiens Asymptomatic Hospitalized Recovered Mpox 79 a year 79 a Year 2022-05-25 Lesion (Pustule), Swollen Lymph Nodes, Myalgia (muscle pain) reinced by the host from the pick list. If the desired Delayed wound healing (lesion healing)		3.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1	0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	1.0.0 1.0.0
Sample collection and processing Host Information	experimental control details Lineage and Variant information Host Information host (common name) host (scientific name) host health state host health status details host health outcome host disease host age host age host age host age unit host gender host residence geo_loc name (country) host residence geo_loc name (state/province/territory) symptom onset date signs and symptoms pre-existing conditions and risk factors complications antiviral therapy	GENEPIC.0010922 GENEPIC.0001498 GENEPIC.0001386 GENEPIC.0001387 GENEPIC.0001387 GENEPIC.0001389 GENEPIC.0001389 GENEPIC.0001389 GENEPIC.0001399 GENEPIC.0001399 GENEPIC.0001399 GENEPIC.0001399 GENEPIC.0001399 GENEPIC.0001399 GENEPIC.0001400 GENEPIC.0001401	The details regarding the experimental control contained in the sail. The taxonomic, or scientific name of the host. The taxonomic, or scientific name of the host. Health status of the host at the time of sample collection. Further details pertaining to the health or disease status of the host at time of collection. Disease outcome in the host. The name of the disease experienced by the host. Age of host at the time of sampling. The units used to measure the host's age. The age category of the host at the time of sampling. The gender of the host at the time of sample collection. The country of residence of the host. The state/province/territory of residence of the host. The date on which the symptoms began or were first noted. A perceived change in function or sensation, (loss, disturbance or appearance) indicative of a disease, reported by a patient. Patient pre-existing conditions and risk factors. <in>Pre-existing co</in>	experimental control or a replicate, select a role type from the picklist. If the sample was not a control, leave blank or select "Not Applicable". Provide details regarding the nature of the reference strain is considered to the select of the select and the select and select and select a select and select	a Human coronavirus 229E (HCoV-229E) spiked in a Human coronavirus 229E (HCoV-229E) spiked in a Human and a Human aspiens Asymptomatic Hospitalized Recovered Mpox 79 New 79		3.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1	0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	10.0 10.0 10.0 10.0 10.0 10.0 10.0 10.0
Sample collection and processing Host Information	experimental control details Lineage and Variant information Host Information host (common name) host (scientific name) host health state host health status details host health outcome host disease host age host age host age host age unit host spender host residence geo_loc name (country) host residence geo_loc name (state/province/territory) symptom onset date signs and symptoms pre-existing conditions and risk factors complications	GENEPIO.0010922 GENEPIO.0001498 GENEPIO.0001386 GENEPIO.0001387 GENEPIO.0001387 GENEPIO.0001389 GENEPIO.0001389 GENEPIO.0001389 GENEPIO.0001389 GENEPIO.0001393 GENEPIO.0001393 GENEPIO.0001393 GENEPIO.0001393 GENEPIO.0001393 GENEPIO.0001394 GENEPIO.0001396 GENEPIO.0001397 GENEPIO.0001397 GENEPIO.0001399 GENEPIO.0001400 GENEPIO.0001400	The details regarding the experimental control contained in the sair The taxonomic, or scientific name of the host. The taxonomic, or scientific name of the host. Health status of the host at the time of sample collection. Further details pertaining to the health or disease status of the host at time of collection. Disease outcome in the host. The name of the disease experienced by the host. Age of host at the time of sampling. The units used to measure the host's age. The age category of the host at the time of sampling. The gender of the host at the time of sample collection. The country of residence of the host. The state/province/territory of residence of the host. The date on which the symptoms began or were first noted. A perceived change in function or sensation, (loss, disturbance or appearance) indicative of a disease, reported by a patient. Patient pre-existing conditions and risk factors. <i>Pre-existing conditions and risk factors. <i>Pre-existing conditions are result of host disease. Treatment of viral infections with agents that prevent viral replication in infected cells without impairing the host cell function.</i></i>	experimental control or a replicate, select a role type from the picklist. If the sample was not a control, leave blank or select "Not Applicable". Provide details regarding the nature of the reference strain is considered to the select of the select and the select and select and select a select and select	a Human coronavirus 229E (HCoV-229E) spiked in a Human coronavirus 229E (HCoV-229E) spiked in a Human and a Human aspiens Asymptomatic Hospitalized Recovered Mpox 79 New 79		3.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1	0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	10.0 10.0 10.0 10.0 10.0 10.0 10.0 10.0
Sample collection and processing Host Information	experimental control details Lineage and Variant information Host Information host (common name) host (scientific name) host health state host health status details host health outcome host disease host age host age host age host age unit host gender host residence geo_loc name (country) host residence geo_loc name (state/province/territory) symptom onset date signs and symptoms pre-existing conditions and risk factors complications antiviral therapy	GENEPIC.0010922 GENEPIC.0001498 GENEPIC.0001386 GENEPIC.0001387 GENEPIC.0001387 GENEPIC.0001389 GENEPIC.0001389 GENEPIC.0001389 GENEPIC.0001399 GENEPIC.0001399 GENEPIC.0001399 GENEPIC.0001399 GENEPIC.0001399 GENEPIC.0001399 GENEPIC.0001400 GENEPIC.0001401	The details regarding the experimental control contained in the sail. The taxonomic, or scientific name of the host. The taxonomic, or scientific name of the host. Health status of the host at the time of sample collection. Further details pertaining to the health or disease status of the host at time of collection. Disease outcome in the host. The name of the disease experienced by the host. Age of host at the time of sampling. The units used to measure the host's age. The age category of the host at the time of sampling. The gender of the host at the time of sample collection. The country of residence of the host. The state/province/territory of residence of the host. The date on which the symptoms began or were first noted. A perceived change in function or sensation, (loss, disturbance or appearance) indicative of a disease, reported by a patient. Patient pre-existing conditions and risk factors. <in>Pre-existing co</in>	experimental control or a replicate, select a role type from the picklist. If the sample was not a control, leave blank or select "Not Applicable". Provide details regarding the nature of the reference strain is considered to the select of the select and the select and select and select a select and select	i Human coronavirus 229E (HCoV-229E) spiked in		3.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1	0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	10.0 10.0 10.0 10.0 10.0 10.0 10.0 10.0

Host vaccination information	host vaccination status	GENEPIO:0001404	The vaccination status of the host (fully vaccinated, partially vaccinated, or not vaccinated).	Select the vaccination status of the host from the pick list.	Not Vaccinated [GENEPIO:0100102]	1.0.0	1.0.0	1.0.0
		GENEPIO:0001406	The same of days of the same in a short but he had	Record how many doses of the vaccine the host has		100	100	100
ost vaccination information	number of vaccine doses received		The number of doses of the vaccine recived by the host. The name of the vaccine administered as the first dose of a	received. Provide the name and the corresponding manufacturer of	1	1.0.0	1.0.0	1.0.0
est vaccination information	vaccination dose 1 vaccine name	GENEPIO:0100313	vaccine regimen.	the Smallpox vaccine administered as the first dose.	IMVAMUNE (Bavarian Nordic)	1.0.0	1.0.0	1.0.0
ost vaccination information	vaccination dose 1 vaccination date	GENEPIO:0100314	The date the first dose of a vaccine was administered.	Provide the date the first dose of Smallpox vaccine was administered. The date should be provided in ISO 8601 standard format "YYYY-MM-DD".	2022-06-01	1.0.0	1.0.0	1.0.0
lost vaccination information	vaccination history	GENEPIO:0100321	A description of the vaccines received and the administration dates of a series of vaccinations against a specific disease or a set of diseases.	Free text description of the dates and vaccines administered against a particular disease/set of diseases. It is also acceptable to concatenate the individual dose information (vaccine name, vaccination date) separated by semicolons.	IMVAMUNE (Bavarian Nordic); 2022-06-01	1.0.0	1.0.0	1.0.0
	Host exposure information	GENEPIO:0001409	The country where the host was likely exposed to the causative	Select the country name from the pick list provided in the		1.0.0	1.0.0	1.0.0
ost exposure information	location of exposure geo_loc name (country)	GENEPIO:0001410	agent of the illness. The country where the host was likely exposed to the causative	template. Select the country name from the pick list provided in the	Canada	1.0.0	1.0.0	1.0.0
ost exposure information	location of exposure geo_loc name (country)	GENEPIO:0001410	agent of the illness.	template.	Canada [GAZ:00002560]	1.0.0	1.0.0	1.0.0
lost exposure information	destination of most recent travel (city)	GENEPIO:0001411	The name of the city that was the destination of most recent travel.	Provide the name of the city that the host travelled to. Use this look-up service to identify the standardized term: https://www.ebi.ac.uk/ols/ontologies/gaz	New York City	1.0.0	1.0.0	1.0.0
			The name of the state/province/territory that was the destination of		,			
lost exposure information	destination of most recent travel (state/province/territory)	GENEPIO:0001412	most recent travel. The name of the state/province/territory that was the destination of	template. Provide the name of the state/province/territory that the flost travelled to. Use this look-up service to identify the		1.0.0	1.0.0	1.0.0
lost exposure information	destination of most recent travel (state/province/territory)	GENEPIO:0001412	most recent travel.	standardized term: https://www.ebi.ac.uk/ols/ontologies/gaz	California	1.0.0	1.0.0	1.0.0
lost exposure information	destination of most recent travel (country)	GENEPIO:0001413	The name of the country that was the destination of most recent travel. The name of the country that was the destination of most recent	Select the country name from the pick list provided in the template. Select the country name from the pick list provided in the	Canada	1.0.0	1.0.0	1.0.0
lost exposure information	destination of most recent travel (country)	GENEPIO:0001413	travel.	select the country name from the pick list provided in the template.	United Kingdom [GAZ:00002637]	1.0.0	1.0.0	1.0.0
			The date of a person's most recent departure from their primary residence (at that time) on a journey to one or more other					
ost exposure information	most recent travel departure date	GENEPIO:0001414	locations.	Provide the travel departure date.	2020-03-16	1.0.0	1.0.0	1.0.0
ost exposure information	most recent travel return date	GENEPIO:0001415	The date of a person's most recent return to some residence from a journey originating at that residence.	Provide the travel return date. Specify the countries (and more granular locations if known, separated by a comma) travelled in the last six	2020-04-26	1.0.0	1.0.0	1.0.0
ost exposure information	travel history	GENEPIO:0001416	Travel history in last six months.	months; can include multiple travels. Separate multiple travel events with a semi-colon. List most recent travel first. Select an exposure event from the pick list provided in the	Canada, Vancouver; USA, Seattle; Italy, Milan	1.0.0	1.0.0	1.0.0
ost exposure information	exposure event	GENEPIO:0001417	Event leading to exposure.	template. If the desired term is missing, contact the DataHarmonizer curation team.	Party	1.0.0	1.0.0	1.0.0
ost exposure information	exposure event	GENEPIO:0001417	Event leading to exposure.	Select an exposure event from the pick list provided in the template. If the desired term is missing, contact the DataHarmonizer curation team.	Party [PCO:0000035]	1.0.0	1.0.0	1.0.0
ost exposure information	exposure contact level	GENEPIO:0001418	The exposure transmission contact type.	Select exposure contact level from the pick-list.	Contact with infected individual	1.0.0	1.0.0	1.0.0
ost exposure information	exposure contact level	GENEPIO:0001418	The exposure transmission contact type.	Select exposure contact level from the pick-list.	Contact with infected individual [GENEPIO:0100357]	1.0.0	1.0.0	1.0.0
ost exposure information	host role	GENEPIO:0001419	The role of the host in relation to the exposure setting.	Select the host's personal role(s) from the pick list provided in the template. If the desired term is missing, contact the DataHarmonizer curation team.	Acquaintance of case	1.0.0	1.0.0	1.0.0
lost exposure information	host role	GENEPIO:0001419	The role of the host in relation to the exposure setting.	Select the host's personal role(s) from the pick list provided in the template. If the desired term is missing, contact the DataHarmonizer curation team.		100	1.0.0	100
				Select the host exposure setting(s) from the pick list provided in the template. If a desired term is missing,				
lost exposure information	exposure setting	GENEPIO:0001428	The setting leading to exposure.	contact the DataHarmonizer curation team. Select the host exposure setting(s) from the pick list provided in the template. If a desired term is missing,	Healthcare Setting	1.0.0	1.0.0	1.0.0
ost exposure information	exposure setting	GENEPIO:0001428	The setting leading to exposure.	contact the DataHarmonizer curation team.	Healthcare Setting [GENEPIO:0100201]	1.0.0	1.0.0	1.0.0
ost exposure information	exposure details	GENEPIO:0001431	Additional host exposure information.	Free text description of the exposure.	Large party, many contacts	1.0.0	1.0.0	1.0.0
lost reinfection information	Host reinfection information	GENEPIO:0001434 GENEPIO:0100532	The absence or presence of a prior Mpox infection.	If known, provide information about whether the individual had a previous Mpox infection. Select a value from the pick list	Prior infection	1.0.0	1.0.0	1.0.0
ost reinfection information	prior Mpox infection	GENEPIO:0100532	The absence or presence of a prior Mpox infection.	If known, provide information about whether the individual had a previous Mpox infection. Select a value from the pick list		1.0.0	1.0.0	1.0.0
				Provide the date that the most recent prior infection was diagnosed. Provide the prior Mpox infection date in ISO				
ost reinfection information	prior Mpox infection date	GENEPIO:0100533	The date of diagnosis of the prior Mpox infection.	8601 standard format "YYYY-MM-DD". If known, provide information about whether the individual	2022-06-20	1.0.0	1.0.0	1.0.0
ost reinfection information	prior Mpox antiviral treatment	GENEPIO:0100534	The absence or presence of antiviral treatment for a prior Mpox infection.	had a previous Mpox antiviral treatment. Select a value from the pick list. If known provide information about whether the individual.	Prior antiviral treatment	1.0.0	1.0.0	1.0.0
			The absence or presence of antiviral treatment for a prior Mpox	had a previous Mpox antiviral treatment. Select a value				
ost reinfection information	prior Mpox antiviral treatment	GENEPIO:0100534	infection.	from the pick list. Provide a description of any antiviral treatment administered for viral infections (not including Mpox	Prior antiviral treatment [GENEPIO:0100037]	1.0.0	1.0.0	1.0.0
				treatment) during the prior Mpox infection period. This field	I			

						1.0.0	1.0.0	1.0.0
	Sequencing	GENEPIO:0001441				1.0.0	1.0.0	1.0.0
equence information	sequencing project name	GENEPIO:0100472	The name of the project/initiative/program for which sequencing was performed.	Provide the name of the project and/or the project ID here. If the information is unknown or cannot be provided, leave blank or provide a null value.	MPOX-1356	1.0.0	1.0.0	1.0.0
				The name of the agency should be written out in full, (with minor exceptions) and be consistent across multiple submissions. If submitting specimens rather than sequencing data, please put the "National Microbiology				
Sequencing	sequenced by	GENEPIO:0100416	The name of the agency that generated the sequence.	Laboratory (NML)*. The name of the agency should be written out in full, (with minor exceptions) and be consistent across multiple	Public Health Ontario (PHO)	1.0.0	1.0.0	1.0.0
Sequencing	sequenced by	GENEPIO:0100416	The name of the agency that generated the sequence.	Provide the name of the specific laboratory that that performed the sequencing in full (avoid abbreviations). If	Public Health Ontario (PHO)	1.0.0	1.0.0	1.0.0
Sequence information	sequenced by laboratory name	GENEPIO:0100470	The specific laboratory affiliation of the responsible for sequencing the isolate's genome.	blank or provide a null value. Provide the name of an individual or their job title. As	Topp Lab	1.0.0	1.0.0	1.0.0
			The name or title of the contact responsible for follow-up regarding	personnel turnover may render the contact's name obsolete, it is more prefereable to provide a job title for ensuring accuracy of information and institutional memory. If the information is unknown or cannot be provided, leave				
Sequence information	sequenced by contact name	GENEPIO:0100471	the sequence. The email address of the contact responsible for follow-up	blank or provide a null value. The email address can represent a specific individual or lab	Joe Bloggs, Enterics Lab Manager	1.0.0	1.0.0	1.0.0
equencing	sequenced by contact email	GENEPIO:0100422	regarding the sequence.	e.g. johnnyblogs@lab.ca, or RespLab@lab.ca The mailing address should be in the format: Street number	RespLab@lab.ca 123 Sunnybrooke St, Toronto, Ontario, M4P	1.0.0	1.0.0	1.0.0
Sequencing	sequenced by contact address	GENEPIO:0100423	The mailing address of the agency submitting the sequence.		1L6, Canada	1.0.0	1.0.0	1.0.0
Sequencing	sequence submitted by	GENEPIO:0001159	The name of the agency that submitted the sequence to a database.	submissions. If submitting specimens rather than sequencing data, please put the "National Microbiology Laboratory (NML)".	Public Health Ontario (PHO)	1.0.0	1.0.0	1.0.0
Sequencing	sequence submitted by	GENEPIO:0001159	The name of the agency that submitted the sequence to a database.	The name of the agency should be written out in full, (with minor exceptions) and be consistent across multiple submissions. If submitting specimens rather than sequencing data, please put the "National Microbiology Laborator, (MNL)".	Public Health Ontario (PHO)	1.0.0	1.0.0	1.0.0
Sequencing	sequence submitter contact email	GENEPIO:0001165	The email address of the agency responsible for submission of the sequence.			1.0.0	1.0.0	1.0.0
Sequencing	sequence submitter contact address	GENEPIO:0001167	The mailing address of the agency responsible for submission of the sequence.	The mailing address should be in the format: Street number	123 Sunnybrooke St, Toronto, Ontario, M4P 1L6. Canada	1.0.0	1.0.0	1.0.0
Sequencing	purpose of sequencing	GENEPIO,0001445	The reason that the sample was sequenced.	The reason why a sample was originally collected may differ from the reason why it was selected for sequencing. The reason a sample was sequenced may provide information about potential biases in sequencing strategy. Provide the purpose of sequencing from the picklist in the	individuals with no known intimate contacts to positive cases. Specimens attributed to youthrimors <18 yrs. Specimens attributed to youthrimors <18 yrs. Specimens attributed to vulnerable persons living in transient shelters or congregant settings. Specimens attributed to individuals self-identifying as "female". For specimens with a recent international and/or domestic travel history, please select the most appropriate tag from the following three options: Domestic travel surveillance, Travel-associated surveillance. For specimens targeted for sequencing as part of an outbreak investigation, please select. Cluster/Outbreak investigation, in all other cases use: Baseline surveillance (random sampling).	1.00	1.0.0	1.0.0
Sequencing	purpose of sequencing		The reason that the sample was sequenced.	The reason why a sample was originally collected may differ from the reason why it was selected for sequencing. The reason a sample was sequenced may provide information about potential biases in sequencing strategy. Provide the purpose of sequencing from the picklist in the	Baseline surveillance (random sampling) [GENEPIO:0100005]	1.0.0	1.0.0	1.0.0
· ·			The description of why the sample was sequenced providing	Provide an expanded description of why the sample was sequenced using free text. The description may include the importance of the sequences for a particular public health investigation/surveillance activity/research question. Suggested standardized descriptions include: Screened due to travel history, Screened due to travel history, Screened due to close contact with				
Sequencing	purpose of sequencing details	GENEPIO:0001446	specific details.	infected individual. ISO 8601 standard "YYYY-MM-DD"	Outbreak in MSM community 2020-06-22	1.0.0	1.0.0	1.0.0
equencing equencing	sequencing date sequencing date		The date the sample was sequenced. The date the sample was sequenced.	ISO 8601 standard "YYYY-MM-DD". ISO 8601 standard "YYYY-MM-DD".	2020-06-22 2020-06-22	1.0.0	1.0.0	1.0.0
equencing	library ID	GENEPIO:0001448	The user-specified identifier for the library prepared for sequencing.	The library name should be unique, and can be an autogenerated ID from your LIMS, or modification of the	XYZ 123345	1.0.0	1.0.0	1.0.0
equencing	library preparation kit	GENEPIO:0001450	The name of the DNA library preparation kit used to generate the library being sequenced.	Provide the name of the library preparation kit used.	Nextera XT	1.0.0	1.0.0	1.0.0
equence information	sequencing assay type	GENEPIO:0100997	The overarching sequencing methodology that was used to determine the sequence of a biomaterial.	Example Guidance: Provide the name of the DNA or RNA sequencing technology used in your study. If unsure refer to the protocol documentation, or provide a null value.	whole genome sequencing assay	1.0.0	1.0.0	1.0.0
Sequence information	sequencing assay type	GENEPIO:0100997	The overarching sequencing methodology that was used to determine the sequence of a biomaterial.	Example Guidance: Provide the name of the DNA or RNA sequencing technology used in your study. If unsure refer to the protocol documentation, or provide a null value.	whole genome sequencing assay [OBI:0002117]	1.0.0	1.0.0	1.0.0
Sequencing	sequencing instrument	GENEPIO:0001452	The model of the sequencing instrument used.	Select a sequencing instrument from the picklist provided in the template.	Oxford Nanopore MinION	1.0.0	1.0.0	1.0.0
				Select a sequencing instrument from the picklist provided in				

				Flow cells can vary in terms of design, chemistry, capacity,				
			The version number of the flow cell used for generating sequence	etc. The version of the flow cell used to generate sequence data can affect sequence quantity and quality. Record the				
Sequence information	sequencing flow cell version	GENEPIO:0101102	data.	not include "version" or "v" in the version number.	R.9.4.1	3.0.0	3.0.0	3.0.0
				approach. Sequencing was performed using a <fill in=""></fill>	Viral sequencing was performed following a metagenomic shotgun sequencing approach. Libraries were created using Illumina DNA Prep kits, and sequence data was produced using			
Sequencing	sequencing protocol	GENEPIO:0001454	The protocol used to generate the sequence.	in> library kit. "	Miseq Micro v2 (500 cycles) sequencing kits.	3.0.0	3.0.0	3.0.0
Sequencing	sequencing kit number	GENEPIO:0001455	The manufacturer's kit number.	Alphanumeric value.	AB456XYZ789	3.0.0	3.0.0	3.0.0
			The length of the DNA fragment generated by mechanical					
Sequencing	DNA fragment length	GENEPIO:0100843	shearing or enzymatic digestion for the purposes of library preparation. The molecular technique used to selectively capture and amplify	Provide the fragment length in base pairs (do not include the units).	400	1.0.0	1.0.0	1.0.0
Sequencing	genomic target enrichment method	GENEPIO:0100966	specific regions of interest from a genome.	Provide the name of the enrichment method	hybrid selection method	1.0.0	1.0.0	1.0.0
Sequencing	genomic target enrichment method	GENEPIO:0100966	The molecular technique used to selectively capture and amplify specific regions of interest from a genome.	Provide the name of the enrichment method	hybrid selection method	1.0.0	1.0.0	1.0.0
			Details that provide additional context to the molecular technique used to selectively capture and amplify specific regions of interest		enrichment was done using Illumina Target Enrichment methodology with the Illumina DNA			
Sequencing	genomic target enrichment method details	GENEPIO:0100967	from a genome. The specifications of the primers (primer sequences, binding		Prep with enrichment kit.	1.0.0	1.0.0	1.0.0
Sequencina	amplicon pcr primer scheme	GENEPIO:0001456	positions, fragment size generated etc) used to generate the amplicons to be sequenced.	Provide the name and version of the primer scheme used to generate the amplicons for sequencing.	MPXV Sunrise 3.1	1.0.0	1.0.0	1.0.0
Sequencing	amplicon size	GENEPIO:0001449	The length of the amplicon generated by PCR amplification.	Provide the amplicon size expressed in base pairs.	300bp	1.0.0	1.0.0	1.0.0
	Bioinformatics and QC metrics	GENEPIO:0001457				1.0.0	1.0.0	1.0.0
	DOMINIMATES AND STREET	SENE COMMON	The name of the method used to assess whether a sequence	Providing the name of the method used for quality control is very important for interpreting the rest of the QC information. Method names can be provided as the name of a pipeline or a link to a GitHub repository. Multiple methods should be listed and separated by a semi-colon. Do not include QC tags in other fields if no method name is				
Bioinformatics and QC metrics	quality control method name	GENEPIO:0100557	passed a predetermined quality control threshold.	provided.	ncov-tools	1.0.0	1.0.0	1.0.0
			The version number of the method used to assess whether a	Methods updates can make big differences to their outputs. Provide the version of the method used for quality control. The version can be expressed using whatever convention the developer implements (e.g. date, semantic versioning). If multiple methods were used, record the version numbers in the same order as the method names. Separate the				
Bioinformatics and QC metrics	quality control method version	GENEPIO:0100558	sequence passed a predetermined quality control threshold.	version numbers using a semi-colon.	1.2.3	1.0.0	1.0.0	1.0.0
				Select a value from the pick list provided. If a desired value is missing, submit a new term request to the PHA4GE QC				
Bioinformatics and QC metrics	quality control determination	GENEPIO:0100559	The determination of a quality control assessment.	Tag GitHub issuetracker using the New Term Request form.	sequence failed quality control	1.0.0	1.0.0	1.0.0
Bioinformatics and QC metrics	quality control determination	GENEPIO:0100559		Select a value from the pick list provided. If a desired value		1.0.0	1.0.0	1.0.0
Bioinformatics and QC metrics	quality control issues	GENEPIO:0100560	The reason contributing to, or causing, a low quality determination in a quality control assessment.	is missing, submit a new term request to the PHA4GE QC Tag GitHub issuetracker using the New Term Request form.	low average genome coverage	1.0.0	1.0.0	1.0.0
Bioinformatics and QC metrics	quality control issues	GENEPIO:0100560	in a quality control assessment.	tag Oil tub issued acker using the New Territ Nequest form.	low average genome coverage	1.0.0	1.0.0	1.0.0
Bioinformatics and QC metrics	quality control dataile	GENEPIO:0100561	The details surrounding a low quality determination in a quality control assessment.	Provide notes or details regarding QC results using free	CT value of 39. Low viral load. Low DNA concentration after amplification	100	100	100
	quality control details		The names of the software and version number used for raw data processing such as removing barcodes, adapter trimming, filtering	Provide the software name followed by the version e.g.				
Bioinformatics and QC metrics	raw sequence data processing method	GENEPIO:0001458	etc. The names of the software and version number used for raw data	Trimmomatic v. 0.38, Porechop v. 0.2.3	Porechop 0.2.3	1.0.0	1.0.0	1.0.0
Bioinformatics and QC metrics	raw sequence data processing method	GENEPIO:0001458	processing such as removing barcodes, adapter trimming, filtering etc.	Trimmomatic v. 0.38, Porechop v. 0.2.3	Porechop 0.2.3	1.0.0	1.0.0	1.0.0
Bioinformatics and QC metrics	dehosting method	GENEPIO:0001459	The method used to remove host reads from the pathogen sequence.	Provide the name and version number of the software used to remove host reads.	Nanostripper	3.0.0	3.0.0	3.0.0
Bioinformatics and QC metrics	dehosting method	GENEPIO:0001459	The method used to remove host reads from the pathogen sequence.	Provide the name and version number of the software used to remove host reads.	Nanostripper	3.0.0	3.0.0	3.0.0
Bioinformatics and QC metrics	deduplication method	GENEPIO:0100831	The method used to remove duplicated reads in a sequence read dataset.	Provide the deduplication software name followed by the version, or a link to a tool or method.	DeDup 0.12.8	1.0.0	1.0.0	1.0.0
	· ·			Provide the name and version number of the consensus				
Bioinformatics and QC metrics	consensus sequence name	GENEPIO:0001460	The name of the consensus sequence.	sequence. Provide the name and version number, with the file extension, of the processed genome sequence file e.g. a	mpxvassembly3	1.0.0	1.0.0	1.0.0
Bioinformatics and QC metrics	genome sequence file name	GENEPIO:0101715	The name of the consensus sequence file.	consensus sequence FASTA file or a genome assembly file.	mpxvassembly.fasta	1.0.0	1.0.0	1.0.0
Bioinformatics and QC metrics	genome sequence file path	GENEPIO:0101716	The filepath of the consensus sequence file.	Provide the filepath of the genome sequence FASTA file.	/User/Documents/ViralLab/Data/mpxvassembly. fasta	1.0.0	1.0.0	1.0.0
				Provide the name of the software used to generate the	iVar			
Bioinformatics and QC metrics	consensus sequence software name	GENEPIO:0001463	The name of software used to generate the consensus sequence. The version of the software used to generate the consensus	Provide the version of the software used to generate the		1.0.0	1.0.0	1.0.0
Bioinformatics and QC metrics	consensus sequence software version	GENEPIO:0001469	sequence.	consensus sequence. Provide the name of the software used to assemble the	1.3 SPAdes Genome Assembler, Canu, wtdbg2,	1.0.0	1.0.0	1.0.0
Bioinformatics and QC metrics	sequence assembly software name	GENEPIO:0100825	The name of the software used to assemble a sequence.	sequence. Provide the version of the software used to assemble the	velvet	1.0.0	1.0.0	1.0.0
Bioinformatics and QC metrics	sequence assembly software version	GENEPIO:0100826	The version of the software used to assemble a sequence.	sequence.	3.15.5	1.0.0	1.0.0	1.0.0
Bioinformatics and QC metrics	r1 fastq filename	GENEPIO:0001476	The user-specified filename of the r1 FASTQ file.	Provide the r1 FASTQ filename. This information aids in data management. Provide the r2 FASTQ filename. This information aids in	ABC123_S1_L001_R1_001.fastq.gz	1.0.0	1.0.0	1.0.0
Bioinformatics and QC metrics	r2 fastq filename	GENEPIO:0001477	The user-specified filename of the r2 FASTQ file.	data management.	ABC123_S1_L001_R2_001.fastq.gz /User/Documents/ViralLab/Data/ABC123_S1_L	1.0.0	1.0.0	1.0.0
Bioinformatics and QC metrics	r1 fasto filepath	GENEPIO:0001478	The location of the r1 FASTQ file within a user's file system.	aids in data management.	/User/Documents/ViraiLab/Data/ABC123_S1_L 001_R1_001.fastq.gz	100	100	100

				Provide the filepath for the r2 FASTQ file. This information	/User/Documents/ViralLab/Data/ABC123_S1_L			
Bioinformatics and QC metrics	r2 fastq filepath	GENEPIO:0001479	The location of the r2 FASTQ file within a user's file system.	aids in data management. Provide the FAST5 filename. This information aids in data	001_R2_001.fastq.gz	1.0.0	1.0.0	1.0.0
pinformatics and QC metrics	fast5 filename	GENEPIO:0001480	The user-specified filename of the FAST5 file.	management.	mpxv123seq.fast5	1.0.0	1.0.0	1.0.0
informatics and QC metrics	fast5 filepath	GENEPIO:0001481	The location of the FAST5 file within a user's file system.	Provide the filepath for the FAST5 file. This information aids in data management.	/User/Documents/RespLab/Data/mpxv123seq.f ast5	1.0.0	1.0.0	1.0.0
nformatics and OC metrics	number of total reads	GENEPIO:0100827	The total number of non-unique reads generated by the sequencing process.	Provide a numerical value (no need to include units).	423867	100	100	100
mornados ana qo mornos			The number of unique reads generated by the sequencing				1.0.0	
informatics and QC metrics	number of unique reads	GENEPIO:0100828	process. The threshold used as a cut-off for the minimum length of a read	Provide a numerical value (no need to include units).	248236	1.0.0	1.0.0	1.0.0
informatics and QC metrics	minimum post-trimming read length	GENEPIO:0100829	after trimming.	Provide a numerical value (no need to include units).	150	1.0.0	1.0.0	1.0.0
informatics and QC metrics	depth of coverage value	GENEPIO:0001474	The average number of reads representing a given nucleotide in the reconstructed sequence.	Provide value as a fold of coverage.	400x	1.0.0	1.0.0	1.0.0
informatics and QC metrics	depth of coverage threshold	GENEPIO:0001475	The threshold used as a cut-off for the depth of coverage.	Provide the threshold fold coverage.	100x	1.0.0	1.0.0	1.0.0
informatics and QC metrics	number of base pairs sequenced	GENEPIO:0001482	The number of total base pairs generated by the sequencing process.	Provide a numerical value (no need to include units).	2639019	1.0.0	1.0.0	1.0.0
informatics and QC metrics	consensus genome length	GENEPIO:0001483	Size of the reconstructed genome described as the number of base pairs.	Provide a numerical value (no need to include units).	197063	100	100	1.0.0
			The length of the genome generated by assembling reads using a	,			1.00	
informatics and QC metrics	sequence assembly length	GENEPIO:0100846	scaffold or by reference-based mapping. The number of contigs (contiguous sequences) in a sequence	Provide a numerical value (no need to include units).	34272	1.0.0	1.0.0	1.0.0
informatics and QC metrics	number of contigs	GENEPIO:0100937	assembly.	Provide a numerical value.	10	1.0.0	1.0.0	1.0.0
informatics and QC metrics	genome completeness	GENEPIO:0100844	(incompleteness) in the data.	Provide the genome completeness as a percent (no need to include units).	85	1.0.0	1.0.0	1.0.0
informatics and QC metrics	N50	GENEPIO:0100938	The length of the shortest read that, together with other reads, represents at least 50% of the nucleotides in a set of sequences.	Provide the N50 value in Mb.	150	1.0.0	1.0.0	1.0.0
informatics and QC metrics	percent Ns across total genome length	GENEPIO:0100830	The percentage of the assembly that consists of ambiguous bases (Ns).	Provide a numerical value (no need to include units).	2	3.0.0	3.0.0	3.0.0
			The number of ambiguous bases (Ns) normalized per 100	,				
informatics and QC metrics	Ns per 100 kbp	GENEPIO:0001484	kilobasepairs (kbp).	Provide a numerical value (no need to include units).	342	1.0.0	1.0.0	1.0.0
nformatics and QC metrics	reference genome accession	GENEPIO:0001485	A persistent, unique identifier of a genome database entry.	· · · · · · · · · · · · · · · · · · ·	NC_063383.1	1.0.0	1.0.0	1.0.0
informatics and QC metrics	bioinformatics protocol	GENEPIO:0001489	A description of the overall bioinformatics strategy used.	Further details regarding the methods used to process raw data, and/or generate assemblies, and/or generate consensus sequences can. This information can be provided in an SOP or protocol or pipeline/workflow. Provide the name and version number of the protocol, or a GitHub link to a pipeline or workflow.	https://dithub.com/ohac-nml/monkeyoox-nf	1.0.0	1.0.0	1.0.0
				Further details regarding the methods used to process raw data, and/or generate assemblies, and/or generate consensus sequences can. This information can be provided in an SOP or protocol or pipeline/workflow. Provide the name and version number of the protocol, or a				
pinformatics and QC metrics	bioinformatics protocol	GENEPIO:0001489 GENEPIO:0101082	A description of the overall bioinformatics strategy used.	GitHub link to a pipeline or workflow.	https://github.com/phac-nml/monkeypox-nf	1.0.0	1.0.0	1.0.0 1.0.0
	Taxonomic Identification Information	GENEFIO.0101082	The name of the software used to map sequence reads to a			1.0.0	1.0.0	
conomic identification information	read mapping software name	GENEPIO:0100832	reference genome or set of reference genes.	Provide the name of the read mapping software.	Bowtie2, BWA-MEM, TopHat	1.0.0	1.0.0	1.0.0
	5	GENEPIO:0100832 GENEPIO:0100833	reference genome or set of reference genes. The version of the software used to map sequence reads to a	•	Bowtie2, BWA-MEM, TopHat	1.0.0	1.0.0	1.0.0
conomic identification information	read mapping software version	GENEPIO:0100833	reference genome or set of reference genes. The version of the software used to map sequence reads to a reference genme or set of reference genes. The name of the taxonomic reference database used to identify	Provide the version number of the read mapping software.	2.5.1	1.0.0	1.0.0	1.0.0
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onomic identification information onomic identification information onomic identification information	read mapping software version taxonomic reference database name taxonomic reference database version	GENEPIO:0100833 GENEPIO:0100834 GENEPIO:0100835	reference genome or set of reference genes. The version of the software used to map sequence reads to a reference genome or set of reference genes. The name of the taxonomic reference database used to identify the organism. The version of the taxonomic reference database used to identify the organism. The The flename of the report containing the results of a taxonomic	Provide the version number of the read mapping software. Provide the name of the taxonomic reference database. Provide the version number of the taxonomic reference database. Provide the flename of the report containing the results of the taxonomic analysis. Providing the date that an analyis was performed can help provide context for tool and reference database versions.	2.5.1 NCBITaxon	1.0.0	1.0.0 1.0.0 1.0.0	1.0.0
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nomic identification information onomic identification information onomic identification information onomic identification information onomic identification information	read mapping software version taxonomic reference database name taxonomic reference database version taxonomic analysis report filename taxonomic analysis date read mapping criteria	GENEPIO.0100833 GENEPIO.0100834 GENEPIO.0100835 GENEPIO.0101074 GENEPIO.0101075 GENEPIO.0100836	reference genome or set of reference genes. The version of the software used to map sequence reads to a reference genome or set of reference genes. The name of the taxonomic reference database used to identify the organism. The version of the taxonomic reference database used to identify the organism. The filename of the report containing the results of a taxonomic analysis. The date a taxonomic analysis was performed. A description of the criteria used to map reads to a reference	Provide the version number of the read mapping software. Provide the name of the taxonomic reference database. Provide the version number of the taxonomic reference database. Provide the version number of the taxonomic reference database. Provide the filename of the report containing the results of the taxonomic analysis was performed can help provide context for tool and reference database versions. Provide the date that the taxonomic analysis was performed in ISO 8601 format, i.e. "YYYY-MM-DD".	2.5.1 NCBITaxon 1.3 MPXV_report123.doc 2024-02-01	1.0.0 1.0.0 1.0.0 1.0.0	1.0.0 1.0.0 1.0.0 1.0.0	1.0.0 1.0.0 1.0.0 1.0.0 1.0.0
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onomic identification information informatio	read mapping software version taxonomic reference database name taxonomic reference database version taxonomic analysis report filename taxonomic analysis date read mapping criteria Pathogen diagnostic testing assay target name 1	GENEPIO:0100833 GENEPIO:0100834 GENEPIO:0100835 GENEPIO:0101074 GENEPIO:0101075 GENEPIO:0100836 GENEPIO:0100836 GENEPIO:0101206	reference genome or set of reference genes. The version of the software used to map sequence reads to a reference genome or set of reference genes. The name of the taxonomic reference database used to identify the organism. The version of the taxonomic reference database used to identify the organism. The flename of the report containing the results of a taxonomic analysis. The date a taxonomic analysis was performed. A description of the criteria used to map reads to a reference sequence. The name of the assay target used in the diagnostic RT-PCR test.	Provide the version number of the read mapping software. Provide the name of the taxonomic reference database. Provide the version number of the taxonomic reference database. Provide the version number of the taxonomic reference database. Provide the fliename of the report containing the results of the taxonomic analysis was performed can help provide context for tool and reference database versions. Provide the date that the taxonomic analysis was performed in ISO 8601 format, i.e. "YYYY-MM-DD". Provide a description of the read mapping criteria. The specific genomic region, sequence, or variant targeted by the assay in a diagnostic test. This may include parts of a gene, non-coding regions, or other genetic elements that serve as a marker for detecting the presence of a pathogen or other relevant entities. Select the above that are applicable to the assay used for the diagnostic test. Select the answer of the gene used for the diagnostic PCR from the standardized pick list.	2.5.1 NCBITaxon 1.3 MPXV_report123.doc 2024-02-01 Phred score >20	1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0	1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0	1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0
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conomic identification information	read mapping software version taxonomic reference database name taxonomic reference database version taxonomic analysis report filename taxonomic analysis date read mapping criteria Pathogon diagnostic testing assay target name 1 assay target details 1 gene name 1 gene symbol 1 diagnostic per protocol 1	GENEPIO-0100833 GENEPIO-0100834 GENEPIO-0100835 GENEPIO-0101074 GENEPIO-0101075 GENEPIO-0101075 GENEPIO-01010836 GENEPIO-0102045 GENEPIO-0102045 GENEPIO-0102045 GENEPIO-0102041 GENEPIO-0102041 GENEPIO-0001508	reference genome or set of reference genes. The version of the software used to map sequence reads to a reference genome or set of reference genes. The name of the taxonomic reference database used to identify the organism. The version of the taxonomic reference database used to identify the organism. The reference database used to identify the organism. The filename of the report containing the results of a taxonomic analysis. The date a taxonomic analysis was performed. A description of the criteria used to map reads to a reference sequence. The name of the assay target used in the diagnostic RT-PCR test. Describe any details of the assay target. The name of the gene used in the diagnostic RT-PCR test. The gene symbol used in the diagnostic RT-PCR test. The name and version number of the protocol used for diagnostic.	Provide the version number of the read mapping software. Provide the name of the taxonomic reference database. Provide the version number of the taxonomic reference database. Provide the version number of the taxonomic reference database. Provide the flename of the report containing the results of the taxonomic analysis was performed can help provide context for tool and reference database versions. Provide the date that the taxonomic analysis was performed in ISO 8601 format, i.e. "YYYY-MM-DD". Provide a description of the read mapping criteria. The specific genomic region, sequence, or variant targeted by the assay in a diagnostic test. This may include parts of a gene, non-coding regions, or other genetic elements that serve as a marker for detecting the presence of a pathogen or other relevant entities. Provide details that are applicable to the assay used for the diagnostic test. Select the name of the gene used for the diagnostic PCR from the standardized pick list. Select the asbreviated representation or standardized symbol of the gene used in the diagnostic test from the pick list. The assay target or specific primer region should be added to assay target are select.	2.5.1 NCBITaxon 1.3 MPXV_report123.doc 2024-02-01 Phred score >20 MPX (orf B6R) MPX (orf B6R) opg190 gene (MPOX) B6R (Li et al., 2006)	1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 7.5.5 7.5.5 1.0.0	1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 7.5.5 7.5.5 1.0.0	1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 7.5.5 7.5.5 7.5.5
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onomic identification information information information information information information information onomic identification information in	read mapping software version taxonomic reference database name taxonomic reference database version taxonomic analysis report filename taxonomic analysis date read mapping criteria Pathogon diagnostic testing assay target name 1 assay target details 1 gene name 1 gene symbol 1 diagnostic pcr protocol 1 diagnostic pcr Ct value 1 assay target details 2 assay target details 2	GENEPIO-0100833 GENEPIO-0100834 GENEPIO-0100835 GENEPIO-0101074 GENEPIO-0101075 GENEPIO-0101075 GENEPIO-01010836 GENEPIO-0101206 GENEPIO-0102045 GENEPIO-0102041 GENEPIO-0001509 GENEPIO-0001509 GENEPIO-0102048	reference genome or set of reference genes. The version of the software used to map sequence reads to a reference genome or set of reference genes. The name of the taxonomic reference database used to identify the organism. The version of the taxonomic reference database used to identify the organism. The version of the taxonomic reference database used to identify the organism. The filename of the report containing the results of a taxonomic analysis. The date a taxonomic analysis was performed. A description of the criteria used to map reads to a reference sequence. The name of the assay target used in the diagnostic RT-PCR test. Describe any details of the assay target. The name of the gene used in the diagnostic RT-PCR test. The name and version number of the protocol used for diagnostic. The Ct value result from a diagnostic SARS-CoV-2 RT-PCR test. The name of the assay target used in the diagnostic RT-PCR test.	Provide the version number of the read mapping software. Provide the name of the taxonomic reference database. Provide the version number of the taxonomic reference database. Provide the version number of the taxonomic reference database. Provide the flename of the report containing the results of the taxonomic analysis was performed can help provide context for tool and reference database versions. Provide the date that the taxonomic analysis was performed in ISO 8601 format, i.e. "YYYY-MM-DD". Provide a description of the read mapping criteria. The specific genomic region, sequence, or variant targeted by the assay in a diagnostic test. This may include parts of a gene, non-coding regions, or other genetic elements that serve as a marker for detecting the presence of a pathogen or other relevant entities. Select the name of the gene used for the diagnostic PCR from the standardized pick list. The assay target or specific primer region should be added to assay target are used. The name and version number of the protocol used for carry Provide the CT value of the sample from the diagnostic rest. The name and version number of the protocol used for carry Provide the CT value of the sample from the diagnostic rest. The name and version number of the protocol used for carry Provide details that are applicable to the assay used for the Provide details that are applicable to the assay used for the Provide details that are applicable to the assay used for the Provide details that are applicable to the assay used for the Provide details that are applicable to the assay used for the Provide details that are applicable to the assay used for the	2.5.1 NCBITaxon 1.3 MPXV_report123.doc 2024-02-01 Phred score >20 MPX (orf B6R) MPX (orf B6R) MPX (orf B6R) 2024-02-01 MPX (orf B6R) MPX (orf B6R) 2024-02-01 MPX (orf B6R) 2024-02-01 MPX (orf B6R) 2024-02-01 MPX (orf B6R)	1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 7.5.5 7.5.5 7.5.5 7.5.5 7.5.5 7.5.5 7.5.5	1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 7.5.5 7.5.5 7.5.5 7.5.5 7.5.5 7.5.5 7.5.5	1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 7.5.5 7.5.5 7.5.5 7.5.5 7.5.5 7.5.5 7.5.5
xonomic identification information inf	read mapping software version taxonomic reference database name taxonomic reference database version taxonomic analysis report filename taxonomic analysis date read mapping criteria Pathogen diagnostic testing assay target name 1 assay target details 1 gene name 1 gene symbol 1 diagnostic protocol 1 diagnostic protocol 1 diagnostic pro Ct value 1 assay target name 2	GENEPIO.0100833 GENEPIO.0100834 GENEPIO.0100835 GENEPIO.0101074 GENEPIO.0101075 GENEPIO.0101075 GENEPIO.0101206 GENEPIO.0102045 GENEPIO.0001507 GENEPIO.0102041 GENEPIO.0102041 GENEPIO.0102049 GENEPIO.0102049 GENEPIO.0102049 GENEPIO.0102041	reference genome or set of reference genes. The version of the software used to map sequence reads to a reference genome or set of reference genes. The name of the taxonomic reference database used to identify the organism. The version of the taxonomic reference database used to identify the organism. The version of the taxonomic reference database used to identify the organism. The flename of the report containing the results of a taxonomic analysis. The date a taxonomic analysis was performed. A description of the criteria used to map reads to a reference sequence. The name of the assay target used in the diagnostic RT-PCR test. Describe any details of the assay target. The name of the gene used in the diagnostic RT-PCR test. The name and version number of the protocol used for diagnostic. The ct value result from a diagnostic SARS-CoV-2 RT-PCR test. The name of the assay target used in the diagnostic RT-PCR test.	Provide the version number of the read mapping software. Provide the name of the taxonomic reference database. Provide the version number of the taxonomic reference database. Provide the version number of the taxonomic reference database. Provide the flename of the report containing the results of the taxonomic analysis was performed can help provide context for tool and reference database versions. Provide the date that the taxonomic analysis was performed in ISO 8601 format, i.e. "YYYY-MM-DD". Provide a description of the read mapping criteria. The specific genomic region, sequence, or variant targeted by the assay in a diagnostic test. This may include parts of a gene, non-coding regions, or other genetic elements are serve as a marker for detecting the presence of a pathogen or other relevant entitles. Provide details that are applicable to the assay used for the diagnostic test. Select the abstrate for detecting the presence of a pathogen or other relevant entitles. Select the abstrate for detecting the presence of substrate diagnostic test. Select the abstrate traper of specific primer region should be added to assay target name. Provide the CT value of the sample from the diagnostic PCT. The specific genomic region, sequence, or variant targeted the content of the protocol used for carry.	2.5.1 NCBITaxon 1.3 MPXV_report123.doc 2024-02-01 Phred score >20 MPX (orf B6R) MPX (orf B6R) MPX (erf B6R) Ser (Li et al., 2006) 1.3 2024-02-01 Phred score >20 2024-02-01	1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 7.5.5 7.5.5 7.5.5 7.5.5 7.5.5 7.5.5 7.5.5	1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 7.5.5 1.0.0 7.5.5 7.5.5 7.5.5	1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 1.0.0 7.5.5 1.0.0 7.5.5 7.5.5 7.5.5

Pathogen diagnostic testing	diagnostic pcr Ct value 2	GENEPIO:0001512	The Ct value result from a diagnostic SARS-CoV-2 RT-PCR test.	Provide the CT value of the sample from the second diagnos	36	7.5.5	7.5.5	7.5.5
Pathogen diagnostic testing	assay target name 3	GENEPIO:0102039	The name of the assay target used in the diagnostic RT-PCR test.	The specific genomic region, sequence, or variant targeted b OPHA (orf B2R)		7.5.5	7.5.5	7.5.5
Pathogen diagnostic testing	assay target details 3	GENEPIO:0102047	Describe any details of the assay target.	Provide details that are applicable to the assay used for the diagnostic test.		7.5.5	7.5.5	7.5.5
Pathogen diagnostic testing	gene name 3	GENEPIO:0001513	The name of the gene used in the diagnostic RT-PCR test.	Select the name of the gene used for the diagnostic PCR fro OPHA (orf B2R)		1.0.0	1.0.0	1.0.0
Pathogen diagnostic testing	gene symbol 3	GENEPIO:0102043	The gene symbol used in the diagnostic RT-PCR test.	Select the abbreviated representation or standardized symbo opg188 gene (MPOX)		7.5.5	7.5.5	7.5.5
Pathogen diagnostic testing	diagnostic pcr protocol 3	GENEPIO:0001514	The name and version number of the protocol used for diagnostic	n The name and version number of the protocol used for carry G2R_G (Li et al., 2010) assay		7.5.5	7.5.5	7.5.5
Pathogen diagnostic testing	diagnostic pcr Ct value 3	GENEPIO:0001515	The Ct value result from a diagnostic SARS-CoV-2 RT-PCR test.	Provide the CT value of the sample from the second diagnos	19	7.5.5	7.5.5	7.5.5
Pathogen diagnostic testing	gene name 4	GENEPIO:0100576	The name of the gene used in the diagnostic RT-PCR test.	Select the name of the gene used for the diagnostic PCR fro G2R_G (TNFR)		1.0.0	1.0.0	1.0.0
Pathogen diagnostic testing	diagnostic pcr Ct value 4	GENEPIO:0100577	The Ct value result from a diagnostic SARS-CoV-2 RT-PCR test.	Provide the CT value of the sample from the second diagnos	27	1.0.0	1.0.0	1.0.0
Pathogen diagnostic testing	gene name 5	GENEPIO:0100578	The name of the gene used in the diagnostic RT-PCR test.	Select the name of the gene used for the diagnostic PCR fro RNAse P		1.0.0	1.0.0	1.0.0
Pathogen diagnostic testing	diagnostic pcr Ct value 5	GENEPIO:0100579	The Ct value result from a diagnostic SARS-CoV-2 RT-PCR test.	Provide the CT value of the sample from the second diagnos	30	1.0.0	1.0.0	1.0.0
	Contributor acknowledgement	GENEPIO:0001516						
Contributor acknowledgement	authors	GENEPIO:0001517	Names of individuals contributing to the processes of sample colle	c Include the first and last names of all individuals that should Tejinder Singh, Fei Hu, Joe Bloom	ogs	1.0.0	1.0.0	1.0.0
Contributor acknowledgement	DataHarmonizer provenance	GENEPIO:0001518	The DataHarmonizer software and template version provenance.	The current software and template version information will be DataHarmonizer v1.4.3, Mpox	v3.3.1	1.0.0	1.0.0	1.0.0