| Parent Class | Field | Ontology Identifier | Definition | Guidance | Examples | Deprecated De Label | eprecated ID Version Tracking | ID | Description/Gui |
|--------------------------------------------|----------------------------------------------------------------------|------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------|----------------------------------|-------|-----------------|
| | Colour Code Legend field name in yellow = required | | | | | IMPORTANT: Only labe will be deprecated, alwa replacement version pro term changes in its mea term will be created. | ays with ovided. If a | | dance |
| | field name in purple = recommended field name in white = optional | | | | | | | | |
| | | | | | | | | | |
| | Database Identifiers | GENEPIO:0001122 | | | | | | | |
| | | | | 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 | | | | | |
| Database identifiers | specimen collector sample ID | GENEPIO:0001123 | The user-defined name for the sample. | concise, unique and consistent within your lab. | ASDFG123 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Database identifiers | specimen collector subsample ID | GENEPIO:0100752 | The user-defined identifier assigned to a portion of the original sample. | Store the ID for the subsample/aliquot. | ASDFG123_12 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Database identifiers | pooled sample ID | GENEPIO:0100996 | The user-defined identifier assigned to a combined (pooled) set of samples. | If the sample being analyzed is the result of pooling individual samples, rename the pooled sample with a new identifier. Store the pooled sample ID. | 12345AYZ | | 1.0.0 | 1.0.0 | 1.0.0 |
| Database identifiers | metagenome-assembled genome (MAG) ID | GENEPIO:0100753 | The user-defined identifier assigned to a genome reconstructed from metagenomic data. | Store the MAG ID. | XYZ1234.1 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Database identifiers | specimen collector project ID | GENEPIO:0100918 | The user-defined project name assigned to a sequencing project. | If the sample was collected or analyzed under the umbrella of a specific project, include the name of that project here. | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Database identifiers | BioProject accession | GENEPIO:0001136 | The INSDC (i.e., ENA, NCBI, or DDBJ) accession number of the BioProject(s) to which the BioSample belongs. | Store the BioProject accession number. BioProjects are an organizing tool that links together raw sequence data, assemblies, and their associated metadata. Each province will be assigned a different bioproject accession number by the National Microbiology Lab. A valid NCBI BioProject accession has prefix PRJN e.g., PRJNA12345, and is created once at the beginning of a new sequencing project. | PR.INARORRE1 | | 1.0.0 | 1.0.0 | 1.0.0 |
| | | | The identifier assigned to a BioSample in | Store the accession returned from the BioSample submission. NCBI BioSamples will have the prefix SAMN, | SAMN14180202, SAMEA00000002, | | | | |
| Database identifiers Database identifiers | BioSample accession INSDC sequence read accession | GENEPIO:0001139 GENEPIO:0101203 | INSDC (i.e., ENA, NCBI, or DDBJ) archives. The identifier assigned to a sequence in one of the International Nucleotide Sequence Database Collaboration (INSDC) repositories. | ENA have the prefix SAMEA, DDBJ have SAMD 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. | SAMD00000001 ERR123456, DRR123456, CRR123456 | | 3.1.1 | 3.1.1 | 3.1.1 |
| Database identifiers | Enterobase accession | GENEPIO:0100759 | The identifier assigned to a sequence in Enterobase archives. | Store the barcode assigned to the submitted sequence. Enterobase barcodes start with different 3 letter codes depending on the organism. | SAL_AA0019AA_ST | | 1.0.0 | 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 | | 3.1.1 | 3.1.1 | 3.1.1 |
| Database identifiers | GISAID accession | GENEPIO:0001147 | The identifier assigned to a sequence in GISAID (the Global Initiative on Sharing All Influenza Data) archives. | Store the accession assigned to the submitted sequence. GISAID accessions start with EPI. | EPI ISL 402131 | | 1.0.0 | 1.0.0 | 1.0.0 |
| | | GENEPIO:0100282 | The user-defined GISAID virus name | GISAID virus names should be in the format | | | | | |
| Database identifiers Database identifiers | GISAID virus name | GENEPIO:0100282 | assigned to the sequence. The user-defined identifier assigned to a specific location from which samples are taken. | "hCoV-19/Country/Identifier/year". Store the ID for the site from which a sample was taken. The "site" is user defined (e.g. It may be a building and its environs, a specific entity within an environment). Please use the same site ID for all samples from a given site, regardless of when these samples were taken. Any important changes in site location, should be represented with a new site ID. | hCoV-19/Canada/prov_rona_99/2020 Site 12A | | 1.0.0 | 1.0.0 | 1.0.0 |
| | | | The user-defined identifier assigned to a specific event during which one or more | Store the ID for the event during which a sample or samples were taken. For example, an event could be one person taking samples from multiple sites, or multiple | | | | | |
| Database identifiers | sampling event ID Sample collection and processing | GENEPIO:0100761 GENEPIO:0001150 | samples are taken, from one or more sites. | people taking samples from one site. | Event 120522.1 | | 1.0.0 | 1.0.0 | 1.0.0 |

| Parent Class | Field | Ontology Identifier | Definition | Guidance | Examples | Deprecated Label | Deprecated ID | Tracking | 15 | 5 |
|-------------------------------------|----------------------------------------------------------------------|---------------------|------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|----------------------------------|-----------------------------------------|--------------------|----------|-------|--------------------------|
| | Colour Code Legend field name in yellow = required | | | | | will be deprecated replacement versi | ion provided. If a | Label | ID | Description/Gui dance |
| | field name in purple = recommended field name in white = optional | | | | | | | | | |
| Sample collection and processing | sample collection data steward name | GENEPIO:0100762 | The name of the individual responsible for the data governance, (meta)data usage and distribution of the sample. | Provide the name of the sample collection data steward. | Joe Bloggs | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | sample collected by laboratory name | GENEPIO:0100428 | The specific laboratory affiliation of the sample collector. | Provide the name of the specific laboratory that collected the sample (avoid abbreviations). If the information is unknown or cannot be provided, leave blank or provide a null value. | Topp Lab | | | 3.1.1 | 3.1.1 | 3.1.1 |
| Sample collection and processing | sample collected by | GENEPIO:0001153 | The name of the organization with which the sample collector is affiliated. | The name of the agency should be written out in full, (with | Public Health Agency of Canada | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | sample collector contact name | GENEPIO:0100432 | The name or job title of the contact responsible for follow-up regarding the sample. | Provide the name of an individual or their job title. As personnel turnover may render the contact's name obsolete, it is more preferable to provide a job title for ensuring accuracy of information and institutional memory. If the information is unknown or cannot be provided, leave blank or provide a null value. | Joe Bloggs, Enterics Lab Manager | | | 3.1.1 | 3.1.1 | 3.1.1 |
| Sample collection and processing | sample collector contact email | GENEPIO:0001156 | The email address of the contact responsible for follow-up regarding the sample. | The email address can represent a specific individual or lab e.g. johnnyblogs@lab.ca, or RespLab@lab.ca | WaterTester@facility.ca | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | geo loc name (country) | GENEPIO:0001181 | The country of origin of the sample. | If known, select a value from the pick list. | 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 of origin of the sample. | Provide the state/province/territory name from the GAZ geography ontology. Search for geography terms here: https://www.ebi.ac.uk/ols/ontologies/ga | Western Cape | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | geo loc name (county/region) | GENEPIO:0100280 | The county/region of origin of the sample. | Provide the county/region name from the GAZ geography ontology. Search for geography terms here: https://www.ebi.ac.uk/ols/ontologies/gaz | Derbyshire | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | geo loc name (city) | GENEPIO:0001189 | The city of origin of the sample. | Provide the city name from the GAZ geography ontology. Search for geography terms here: https://www.ebi.ac.uk/ols/ontologies/gaz | Vancouver | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | geo loc name (site) | GENEPIO:0100436 | The name of a specific geographical location e.g. Credit River (rather than river). | Provide the name of the specific geographical site using a specific noun (a word that names a certain place, thing). | Credit River | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | geo loc latitude | GENEPIO:0100309 | The latitude coordinates of the geographical location of sample collection. | Provide latitude coordinates if available. Do not use the centre of the cityregion/province/state/country or the location of your agency as a proxy, as this implicates a real location and is misleading. Specify as degrees latitude in format "d(d.odd) NIS". | 38.98 N | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | geo loc longitude | GENEPIO:0100310 | The longitude coordinates of the geographical location of sample collection. | Provide longitude coordinates if available. Do not use the centre of the cityregion/province/state/country or the location of your agency as a proxy, as this implicates a real location and is misleading. Specify as degrees longitude in format "fd(d.ddd) VIE". | 77.11 W | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | watershed shapefile availability | GENEPIO:0100919 | The availability status of a shapefile descriping the catchment contributing to a watershed. | Select a value from the picklist to describe whether or not a watershed shapefile would be available upon request. | watershed shapefile available | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | organism | GENEPIO:0001191 | Taxonomic name of the organism. | Provide the official nomenclature for the organism(s) present in the sample. Multiple organisms can be entered, separated by semicolons. Avoid abbreviations. Search for taxonomic names here: ncbi.nlm.nlh.gov/taxonomy. | Vibrio cholerae | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | organism | GENEPIO:0001191 | Taxonomic name of the organism. | Provide the official nomenclature for the organism(s) present in the sample if AMR profiles are assigned to organisms. Multiple organisms can be entered, separated by semicolons. Avoid abbreviations. Search for taxonomic names here: ncbi.nlm.nih.gov/taxonomy. | Vibrio cholerae | | | 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. | The reason a sample was collected may provide information about potential biases in sampling strategy. Provide the purpose of sampling from the picklist in the template. Most likely, the sample was collected for Public health surveillance. 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. | Public health surveillance | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | scale of sampling | GENEPIO:0100877 | The range of locations or entities sampled expressed in general terms. | Provide the scale of wastewater sampling by selecting a value from the picklist. | Community-level surveillance | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | sample received date | GENEPIO:0001179 | The date on which the sample was received. | Provide the sample received date in ISO 8601 format, i.e. "YYYY-MM-DD". | 2020-03- | 28 | | 1.0.0 | 1.0.0 | 1.0.0 |

| Parent Class | Field | Ontology Identifier | Definition | Guidance | Examples | Deprecated Label | Deprecated ID | Version Tracking | | |
|-----------------------------------|----------------------------------------------------------------------|---------------------|------------------------------------------------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----------------------------------------------------------------------------------------------------------|----------------------------------|---------------------|-------|-----------------|
| | Colour Code Legend | 37 | | | | | | Label | ID | Description/Gui |
| | field name in yellow = required | | | | | IMPORTANT: Only will be deprecated replacement versic term changes in its term will be create | on provided. If a meaning, a new | | | dance |
| | field name in purple = recommended field name in white = optional | | | | | | | | | |
| Sample collection and processing | sample collection start date | GENEPIO:0001174 | | If your sample is a continuous sample please use this field to capture your start date. Sample collection date is critical for surveillance and many types of analyses. Required granularity includes year, month and day. The Yate should be provided in ISO 8601 standard format "YYYY-MM-DD". | 2020-03-1 | 5 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and | • | | The date on which sample collection ended | Provide the date that sample collection ended in ISO 8601 | | | | | | |
| processing | sample collection end date | GENEPIO:0101071 | for a continuous sample. | format i.e. YYYY-MM-DD Provide the sample processed date in ISO 8601 format, i.e. | 2020-03-1 | 3 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | sample processing date | GENEPIO:0100763 | The date on which the sample was processed. | "YYYY-MM-DD". The sample may be collected and processed (e.g. filtered, extraction) on the same day, or on different dates. | 2020-03-1 | 3 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | sample collection start time | GENEPIO:0101072 | The time at which sample collection began. | Provide this time in ISO 8601 24hr format, in your local time. | 17:15 PST | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | sample collection end time | GENEPIO:0101073 | The time at which sample collection ended. | Provide this time in ISO 8601 24hr format, in your local time. | 19:15 PST | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | sample collection time of day | GENEPIO:0100765 | The descriptive time of day during which the sample was collected. | If known, select a value from the pick list. The time of sample processing matters especially for grab samples, as fecal concentration in wastewater fluctuates over the course of the day. | Morning | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | sample collection time duration value | GENEPIO:0100766 | The amount of time over which the sample was collected. | Provide the numerical value of time. | | 1 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | sample collection time duration unit | GENEPIO:0100767 | The units of the time duration measurement of sample collection. | Provide the units from the pick list. | Hour | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and | | | The activities or variables upstream of | If there was an activity that would affect the sample prior to collection (this is different than sample processing), provide the activities by selecting one or more values from the template pick list. If the information is unknown or cannot be | | | | | | |
| processing Sample collection and | presampling activity | GENEPIO:0100433 | The details of the activities or variables that | provided, leave blank or provide a null value. | Agricultural activity Agricultural waste from large farm | | | 1.0.0 | 1.0.0 | 1.0.0 |
| processing Sample collection and | presampling activity details | GENEPIO:0100434 | affected the sample collected. The numerical value of the volume | Briefly describe the presampling activities using free text. | contributes waste to the site sampled. | | | 1.0.0 | 1.0.0 | 1.0.0 |
| processing Sample collection and | sample volume measurement value | GENEPIO:0100768 | measurement of the sample collected. The units of the volume measurement of the | Provide the numerical value of volume. | | 5 | | 1.0.0 | 1.0.0 | 1.0.0 |
| processing | sample volume measurement unit | GENEPIO:0100769 | sample collected. | Provide the units from the pick list. | mL The sample was placed in a tube in a | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | sample storage method | GENEPIO:0100448 | The process used to store the sample. | Provide details of how the sample was stored from time of collection until time of processing, if there were issues with the cold chain storage, note those here. | cooler bag during transportation (~3 hours) to the lab site. At this point the sample was placed in storage medium and put in a -10C freezer until it was processed and extracted 5 days later. | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and | | | | Provide the name of the transport medium or storage medium used for this sample. If none was used, leave blank | , | | | | | |
| processing | sample storage medium | GENEPIO:0100449 | The medium in which a sample is stored. The numerical value of the time | or write "None". | Cary-Blair transport medium | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | sample storage duration value | GENEPIO:0101014 | measurement during which a sample is in storage. | Provide the numerical value of time. | | 5 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | sample storage duration unit | GENEPIO:0101015 | The units of a measured sample storage duration. | Provide the units from the pick list. | Day | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | specimen processing | GENEPIO:0001253 | Any processing applied to the sample during or after receiving the sample. | | Centrifugation | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | specimen processing details | GENEPIO:0100311 | The details of the processing applied to the sample during or after receiving the sample. | Briefly describe the processes applied to the sample. | 25 samples were pooled and further prepared as a single sample during librar prep. | , | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and | | | The name of the overarching experimental methodology that was used to process the | Provide the name of the methodology used in your study. If | | | | | | |
| processing Sample collection and | experimental protocol field | GENEPIO:0101029 | biomaterial. An environmental location may describe a site in the natural or built environment e.g. contact surface, metal can, hospital, wet | available, provide a link to the protocol. Provide a descriptor of the environmental site sampled. Use the picklist provided in the template. If not applicable | | | | 1.0.0 | 1.0.0 | 1.0.0 |
| processing | environmental site | GENEPIO:0001232 | market, bat cave. An environmental location in the natural or built environment, that is proximal to a | choose a null value. Provide a descriptor of the environmental site close to the | Meat processing plant | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | proximal environmental site | GENEPIO:0101205 | sampling location and which can impact a sample. | sampling site. Use the picklist provided in the template. If not applicable, choose a null value. | Farm | | | 3.1.1 | 3.1.1 | 3.1.1 |

| Parent Class | Field | Ontology Identifier | Definition | Guidance | Examples | Deprecated Label | Deprecated ID | Tracking | | |
|-------------------------------------|----------------------------------------------------------------------------------------|---------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------|----------------------------------|----------|-------|--------------------------|
| | Colour Code Legend field name in yellow = required field name in purple = recommended | | | | | IMPORTANT: Only will be deprecated, replacement versic term changes in its term will be created | on provided. If a meaning, a new | Label | ID | Description/Gui dance |
| | field name in white = optional | | | | | | | | | |
| Sample collection and processing | environmental material | GENEPIO:0001223 | A substance obtained from the natural or man-made environment e.g. soil, water, sewage. | Provide a descriptor of the environmental material sampled. Use the picklist provided in the template. If not applicable, choose a null value. | Raw wastewater | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | environmental material properties | GENEPIO:0100770 | The properties, characteristics and qualities of a substance obtained from the natural or man-made environment. | Provide the environmental material properties by selecting descriptors from the pick list. | Stagnant | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | wastewater system type | GENEPIO:0100771 | The type or classification of a wastewater system e.g. sanitary sewer, combined sewer, latrine | Provide the classification of the wastewater system by selecting from the provided pick list. | Sanitary sewer | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | experimental specimen role type | GENEPIO:0100921 | The type of role that the sample represents in the experiment. | 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 the picklist. If the sample was not a control, leave blank or select "Not Applicable". | | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | experimental control details | GENEPIO:0100922 | The details regarding the experimental control contained in the sample. | Provide details regarding the nature of the reference strain used as a control, or what is was used to monitor. | Human coronavirus 229E (HCoV-229E) spiked in sample as process control | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | collection device | GENEPIO:0001234 | The instrument or container used to collect the sample e.g. grab sampler. | Provide a descriptor of the device used for sampling. Use the picklist provided in the template. If not applicable, choose a null value. | Automatic flow-proportional sampler | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | collection method | GENEPIO:0001241 | The process used to collect the sample. | Provide a descriptor of the collection method used for sampling. Use the picklist provided in the template. If not applicable, choose a null value. | Automatic composite sampling | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | nucleic acid extraction method | GENEPIO:0100939 | The process used to extract genomic material from a sample. | Briefly describe the extraction method used. | Direct wastewater RNA capture and purification via the "Sewage, Salt, Silica and SARS-CoV-2 (4S)" method v4 found at https://www.protocols.io/view/v-4-direct-wastewater-rna-capture-and-purification-36/wq581y4K5/v4 | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and | | | The kit used to extract genomic material from | | 0.70 | | | | | |
| processing | nucleic acid extraction kit | GENEPIO:0100772 | a sample | Provide the name of the genomic extraction kit used. Provide the names of endogenous controls that were used | QIAamp PowerFecal Pro DNA Kit | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | endogenous control details | GENEPIO:0100923 | The description of the endogenous controls included when extracting a sample. | as a reference during extraction. If relevant, include titers of these controls, as well as whether any controls were expected but not identified in the sample. | | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | extraction recovery efficiency measurement value | GENEPIO:0100924 | The recovery efficiency of an extraction, calculated as the amount of a synthetic or endogenous compound identified in the sample relative to the amount expected. | Provide value as a percent. | | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sample collection and processing | extraction recovery efficiency measurement method | GENEPIO:0100925 | The method by which recovery efficiency of an extraction was calculated. | Provide a brief description of how extraction recovery efficiency was measured or estimated. | | | | 1.0.0 | 1.0.0 | 1.0.0 |
| | Strain and isolation information | GENEPIO:0100453 | | , | | | | | | |
| Strain and isolation information | microbiological method | GENEPIO:0100454 | The laboratory method used to grow, prepare, and/or isolate the microbial isolate. | Provide the name and version number of the microbiological method. The ID of the method is also acceptable if the ID can be linked to the laboratory that created the procedure. | MFHPB-30 | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Strain and isolation information | strain | GENEPIO:0100455 | The strain identifier. | If the isolate represents or is derived from, a lab reference strain or strain from a type culture collection, provide the strain identifier. | K12 | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Strain and isolation information | isolate ID | GENEPIO:0100456 | The user-defined identifier for the isolate, as provided by the laboratory that originally isolated the isolate. | Provide the isolate_ID created by the lab that first isolated the isolate (i.e. the original isolate ID). If the information is unknown or cannot be provided, leave blank or provide a null value. If only an alternate isolate ID is known (e.g. the ID from your lab, if your lab did not isolate the isolate from the original sample), make a sure to include it in the alternative_isolate_ID field. | SA20131043 | | | 1.0.0 | 1.0.0 | 1.0.0 |

| Parent Class | Field | Ontology Identifier | Definition | Guidance | Examples | Deprecated Label | Deprecated ID | Version Tracking | | |
|----------------------------------------------|-----------------------------------------------------------------------|---------------------|---------------------------------------------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------|-----------------------------------------|---------------------|-------|-----------------|
| . arom siaco | | omology racinates | | | | | | Label | ID | Description/Gui |
| | Colour Code Legend field name in yellow = required | | | | | IMPORTANT: Only will be deprecated, replacement versic term changes in its term will be created | on provided. If a seemed meaning, a new | | | dance |
| | field name in purple = recommended field name in white = optional | | | | | | | | | |
| Strain and isolation information | alternative isolate ID | GENEPIO:0100457 | An alternative isolate_ID assigned to the isolate by another organization. | Alternative isolate IDs should be provided in the in a prescribed format which consists of the ID followed by square brackets (no space in between the ID and bracket) containing the short form of ID provider's agency name i.e. ID[short organization code]. An example of a properly formatted alternative_isolate_identifier would be e.g. XYZ4567(CFIA). Multiple alternative isolate IDs can be provided, separated by semi-colons. | GHIF3456[PHAC]; QWICK222[CFIA] | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Strain and isolation | | | The identifier assigned to a progenitor isolate derived from an isolate that was directly | lf your sequence data pertains to progeny of an original | | | | | | |
| information | progeny isolate ID | GENEPIO:0100458 | obtained from a sample. | isolate, provide the progeny_isolate_ID. | SUB_ON_1526 | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Strain and isolation information | isolated by | GENEPIO:0100461 | The name of the agency, organization or institution with which the individual who performed the isolation procedure is affiliated. | Provide the name of the agency, organization or institution that isolated the original isolate in full (avoid abbreviations). If the information is unknown or cannot be provided, leave blank or provide a null value. | Public Health Agency of Canada (PHAC) [GENEPIO:0100551] | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Strain and isolation information | isolated by laboratory name | GENEPIO:0100462 | The specific laboratory affiliation of the individual who performed the isolation procedure. | Provide the name of the specific laboratory that that isolated the original isolate (avoid abbreviations). If the information is unknown or cannot be provided, leave blank or provide a null value. | Topp Lab | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Strain and isolation information | isolated by contact name | GENEPIO:0100463 | The name or title of the contact responsible for follow-up regarding the isolate. | Provide the name of an individual or their job title. As personnel turnover may render the contact's name obsolete, it is prefereable to provide a job title for ensuring accuracy of information and institutional memory. If the information is unknown or cannot be provided, leave blank or provide a null value. | Enterics Lab Manager | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Strain and isolation information | isolated by contact email | GENEPIO:0100464 | | Provide the email associated with the listed contact. As personnel turnover may render an individual's email obsolete, it is more prefereable to provide an address for a position or lab, to ensure accuracy of information and institutional memory. If the information is unknown or cannot be provided, leave blank or provide a null value. | enterics@lab.ca | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Strain and isolation information | isolation date | GENEPIO:0100465 | The date on which the isolate was isolated from a sample. | Provide the date according to the ISO 8601 standard "YYYY-MM-DD", "YYYY-MM" or "YYYY". | 2020-10-3 | 0 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Strain and isolation information | isolate received date | GENEPIO:0100466 | The date on which the isolate was received by the laboratory. | Provide the date according to the ISO 8601 standard "YYYY-MM-DD", "YYYY-MM" or "YYYY". | 2020-11-1 | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Strain and isolation information | serovar | GENEPIO:0100467 | The serovar of the organism. | Only include this information if it has been determined by traditional serological methods or a validated in silico prediction tool e.g. SISTR. | Heidelberg | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Strain and isolation information | serotyping method | GENEPIO:0100468 | The method used to determine the serovar. | If the serovar was determined via traditional serotyping methods, put "Traditional serotyping". If the serovar was determined via in silico methods, provide the name and version number of the software | SISTR 1.0.1 | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Strain and isolation | | | | | | _ | | | | |
| information | phagetype Environmental conditions and | GENEPIO:0100469 | The phagetype of the organism. | Provide if known. If unknown, put "missing". | 4 | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | measurements water catchment area human population measurement value | GENEPIO:0100773 | The numerical value of the human population measurement that contributes to the composition of water in a catchment area. | Where known, provide the numerical value of population size, i.e. the number of people. | 10,50 | 0 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | water catchment area human population range | GENEPIO:0100774 | The human population range of the water catchment that contributes effluent to a wastewater site. | Where catchment oppulation is not well known, provide an estimation of population size by selecting a value from the picklist. | 1,000 - 10,000 people | - | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | water catchment area human population measurement method | GENEPIO:0100775 | The method by which a water catchment 's human population size was measured or estimated | Provide a brief description of how catchment population size was measured or estimated. | population of jurisdiction encompassing the wastewater service area | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | water catchment area human population density value | GENEPIO:0100776 | The numerical value describing the number of humans per geographical area in a water catchment. | Provide the numerical value of the population density in the catchement area. | | 4 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | water catchment area human population density unit | GENEPIO:0100777 | The unit describing the number of humans per geographical area in a water catchment. | Provide the unit of the population density in the catchement area. | persons per Km^2 | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | populated area type | GENEPIO:0100778 | A type of area that is populated by humans to different degrees. | Provide the populated area type from the pick list. | Urban area | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | sampling weather conditions | GENEPIO:0100779 | The state of the atmosphere at a place and time as regards heat, dryness, sunshine, wind, rain, etc. | Provide the weather conditions at the time of sample collection. | Rain | | | 1.0.0 | 1.0.0 | 1.0.0 |

| Barrett Olava | Et al. | Outstand Marking | B. C. W. | 2.11 | Formula | | Deprecated ID | Version Tracking | | |
|--------------------------------------------------------------------------|------------------------------------------------------------------|---------------------|--------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------|-------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------|--------------------------------------------------------|---------------------|-------|-----------------|
| Parent Class | Field | Ontology Identifier | Definition | Guidance | Examples | Label | | Label | ID | Description/Gui |
| | Colour Code Legend field name in yellow = required | | | | | IMPORTANT: Only will be deprecated, replacement versic term changes in its term will be created | , always with on provided. If a s meaning, a new | | | dance |
| | field name in purple = recommended | | | | | | | | | |
| | field name in white = optional | | | | | | | | | |
| Environmental conditions and measurements | presampling weather conditions | GENEPIO:0100780 | Weather conditions prior to collection that may affect the sample. | Provide the weather conditions prior to sample collection. | Drizzle | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | precipitation measurement value | GENEPIO:0100911 | | Provide the quantity of precipitation in the area leading up to the time of sample collection. | 1 | 0 | | 1.0.0 | 1.0.0 | 1.0.0 |
| and measurements | precipitation measurement value | GENEPIO.0100911 | a precipitation process. The units of measurement for the amount of | to the time of sample collection. | | 2 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | precipitation measurement unit | GENEPIO:0100912 | water which has fallen during a precipitation process. | Provide the units of precipitation by selecting a value from the pick list. | inch | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | precipitation measurement method | GENEPIO:0100913 | The process used to measure the amount of water which has fallen during a precipitation process. | Provide the name of the procedure or method used to measure precipitation. | Rain gauge over a 12 hour period prior to sample collection | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions | precipitation measurement metrod | | The numerical value of a measurement of | measure precipitation. | | | | 1.0.0 | 1.0.0 | 1.0.0 |
| and measurements Environmental conditions | ambient temperature measurement value | GENEPIO:0100935 | the ambient temperature. The units of a measurement of the ambient | Provide the numerical value of the measured temperature. | 7 | 0 | | 1.0.0 | 1.0.0 | 1.0.0 |
| and measurements | ambient temperature measurement unit | GENEPIO:0100936 | temperature. | Provide the units of the measured temperature. | degree Celsius (C) | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | pH measurement value | GENEPIO:0001736 | The measured pH value indicating the acidity or basicity(alkalinity) of an aqueous solution. | | 7. | 4 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | pH measurement method | GENEPIO:0100781 | The process used to measure pH value. | Provide the name of the procedure or technology used to measure pH. | pH test strip (litmus test) | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions | · | | The numerical value of a measured fluid flow | | | _ | | | | |
| and measurements Environmental conditions | total daily flow rate measurement value | GENEPIO:0100905 | rate over the course of a day. The units of a measured fluid flow rate over | Provide the numerical value of the measured flow rate. Provide the units of the measured flow rate by selecting a | 1 | 0 | | 1.0.0 | 1.0.0 | 1.0.0 |
| and measurements | total daily flow rate measurement unit | GENEPIO:0100906 | the course of a day. | value from the pick list. | million gallons per day (MGD) | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements Environmental conditions | total daily flow rate measurement method | GENEPIO:0100907 | flow rate. The numerical value of a measured | Provide the name of the procedure or technology used to measure flow rate. | Flow meter | | | 1.0.0 | 1.0.0 | 1.0.0 |
| and measurements | instantaneous flow rate measurement value | GENEPIO:0100908 | instantaneous fluid flow rate. | Provide the numerical value of the measured flow rate. | 2 | 5 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | instantaneous flow rate measurement unit | GENEPIO:0100909 | The units of a measured instantaneous fluid flow rate. | Provide the units of the measured flow rate by selecting a value from the pick list. | cubic meter per hour (m^3/h) | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | instantaneous flow rate measurement method | GENEPIO:0100910 | The process used to measure instantaneous fluid flow rate. | Provide the name of the procedure or technology used to measure flow rate. | Flow meter | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | turbidity measurement value | GENEPIO:0100783 | The numerical value of a measurement of turbidity. | Provide the numerical value of the measured turbidity. | 0.0 | 2 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | turbidity measurement unit | GENEPIO:0100914 | The units of a measurement of turbidity. | Provide the units of the measured turbidity by selecting a value from the pick list. | nephelometric turbidity unit (NTU) | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | turbidity measurement method | GENEPIO:0101013 | The process used to measure turbidity. | Provide the name of the procedure or technology used to measure turbidity. | Nephelometric method | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | dissolved oxygen measurement value | GENEPIO:0000035 | The numerical value of a measurement of dissolved oxygen. | Provide the numerical value of the measured dissolved oxygen. | | 5 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | dissolved oxygen measurement unit | GENEPIO:0100784 | The units of a measurement of dissolved oxygen. | Provide the units of the measured dissolved oxygen by selecting a value from the pick list. | part per million (ppm) | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | dissolved oxygen measurement method | GENEPIO:0100785 | The method used to measure dissolved oxygen. | Provide the name of the procedure or technology used to measure dissolved oxygen. | Dissolved oxygen meter in vertical direction | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions | oxygen reduction potential (ORP) | | The numerical value of a measurement of | Provide the numerical value of the measured oxygen | | | | | | |
| and measurements Environmental conditions | measurement value oxygen reduction potential (ORP) | FIX:0000278 | oxygen reduction potential (ORP). The units of a measurement of oxygen | reduction potential. Provide the units of the measured oxygen reduction | -5 | 0 | | 1.0.0 | 1.0.0 | 1.0.0 |
| and measurements | measurement unit | GENEPIO:0100786 | reduction potential (ORP). | potential by selecting a value from the pick list. | milliVolt (mV) | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | oxygen reduction potential (ORP) measurement method | GENEPIO:0100787 | The method used to measure oxygen reduction potential (ORP). | Provide the name of the procedure or technology used to measure oxygen reduction potential. | ORP sensor | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | chemical oxygen demand (COD) measurement value | GENEPIO:0100788 | The measured value from a chemical oxygen demand (COD) test. | Provide the numerical value of the COD test result. | 2 | 6 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | chemical oxygen demand (COD) measurement unit | GENEPIO:0100789 | The units associated with a value from a chemical oxygen demand (COD) test. | Provide the units of the COD test result. | milligram per liter (mg/L) | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | chemical oxygen demand (COD) measurement method | GENEPIO:0100790 | The method used to measure chemical oxygen demand (COD). | Provide the name of the procedure or technology used to measure COD. | Hach LCK test kit | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | carbonaceous biochemical oxygen demand (CBOD) measurement value | GENEPIO:0100791 | The numerical value of a measurement of carbonaceous biochemical oxygen demand (CBOD). | Provide the numerical value of the measured CBOD. | 2 | 0 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | carbonaceous biochemical oxygen demand (CBOD) measurement unit | GENEPIO:0100792 | The units of a measurement of | Provide the units of the measured CBOD by selecting a value from the pick list. | milligram per liter (mg/L) | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | carbonaceous biochemical oxygen demand (CBOD) measurement method | GENEPIO:0100793 | The method used to measure carbonaceous biochemical oxygen demand (CBOD). | Provide the name of the procedure or technology used to measure CBOD. | CBOD measurement by optical probe | | | 1.0.0 | 1.0.0 | 1.0.0 |
| | | | | | | | | | | |

| Parent Class | Field | Ontology Identifier | Definition | Guidance | Examples | Deprecated Deprecated ID Label | Version Tracking | | |
|----------------------------------------------|----------------------------------------------------|---------------------|------------------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------|-------|--------------------------|
| | Colour Code Legend | | | | | | Label | ID | Description/Gui dance |
| | field name in yellow = required | | | | | IMPORTANT: Only labels and/or IDs will be deprecated, always with replacement version provided. If a term changes in its meaning, a new term will be created. | | | |
| | field name in purple = recommended | | | | | | | | |
| | field name in white = optional | | | | | | | | |
| Environmental conditions and measurements | total suspended solids (TSS) measurement value | GENEPIO:0100794 | The numerical value from a total suspended solids (TSS) test. | Provide the numerical value of the measured TSS. | | 8 | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | total suspended solids (TSS) measurement unit | GENEPIO:0100795 | The units associated with a value from a total suspended solids (TSS) test. | Provide the units of the measured TSS. | percent (%) | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | total suspended solids (TSS) measurement method | GENEPIO:0100796 | The method used to measure total suspended solids (TSS). | Provide the name of the procedure or technology used to measure TSS. | Vacuum filter through a 2-micron filter, then oven-dried and weighed sample | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | total dissolved solids (TDS) measurement value | GENEPIO:0100797 | The numerical value from a total dissolved solids (TDS) test. | Provide the numerical value of the measured TDS. | | 2 | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | total dissolved solids (TDS) measurement unit | GENEPIO:0100798 | The units associated with a value from a total dissolved solids (TDS) test. | Provide the units of the measured TDS. | percent (%) | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | total dissolved solids (TDS) measurement method | GENEPIO:0100799 | The method used to measure total dissolved solids (TDS). | Provide the name of the procedure or technology used to measure TDS. | Subtract calculated TSS from calculated TS | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions | | | The numerical value from a total solids (TS) | | | | | | |
| and measurements Environmental conditions | total solids (TS) measurement value | GENEPIO:0100800 | test. The units associated with a value from a total | Provide the numerical value of the measured TS. | 11 | U | 1.0.0 | 1.0.0 | 1.0.0 |
| and measurements Environmental conditions | total solids (TS) measurement unit | GENEPIO:0100801 | solids (TS) test. The method used to measure total solids | Provide the units of the measured TS. Provide the name of the procedure or technology used to | percent (%) Gravimetric method by oven drying, then | | 1.0.0 | 1.0.0 | 1.0.0 |
| and measurements Environmental conditions | total solids (TS) measurement method | GENEPIO:0100802 | (TS). | measure TS. | weighing | | 1.0.0 | 1.0.0 | 1.0.0 |
| and measurements | alkalinity measurement value | GENEPIO:0100878 | The numerical value of a measurement of alkalinity. | Provide the numerical value of the measured alkalinity. | | 3 | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | alkalinity measurement unit | GENEPIO:0100879 | The units of a measurement of alkalinity. | Provide the units of the measured alkalinity. | milligram per liter of calcium carbonate (mg/L CaCO3) | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | alkalinity measurement method | GENEPIO:0100880 | The process used to measure alkalinity. | Provide the name of the procedure or technology used to measure alkalinity. | Titration method | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | conductivity measurement value | GENEPIO:0100916 | The numerical value of a measurement of conductivity. | Provide the numerical value of the measured conductivity. | 141: | 2 | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | conductivity measurement unit | GENEPIO:0100803 | The units of a measurement of conductivity. | Provide the units of the measured conductivity. | microSiemen per centimeter (μS/cm) | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | conductivity measurement method | GENEPIO:0100804 | The method used to measure conductivity. | Provide the name of the procedure or technology used to measure conductivity. | Conductivity electrode and meter | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | salinity measurement value | GENEPIO:0100805 | The numerical value of a measurement of salinity. | Provide the numerical value of the measured salinity. | 3: | 5 | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | salinity measurement unit | GENEPIO:0100806 | The units of a measurement of salinity. | Provide the units of the measured salinity. | practical salinity unit (PSU) | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | salinity measurement method | GENEPIO:0100807 | The method used to measure salinity. | Provide the name of the procedure or technology used to measure salinity. | conductivity meter | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | total nitrogen (TN) measurement value | GENEPIO:0100808 | The numerical value of a measurement of total nitrogen (TN). | Provide the numerical value of the measured TN. | 12 | 0 | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | total nitrogen (TN) measurement unit | GENEPIO:0100809 | The units of a measurement of total nitrogen (TN). | Provide the units of the measured TN. | milligram per liter (mg/L) | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | total nitrogen (TN) measurement method | GENEPIO:0100810 | The method used to measure total nitrogen (TN). | Provide the name of the procedure or technology used to measure TN. | Hach total nitrogen spectrophotometric test | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | total phosphorus (TP) measurement value | GENEPIO:0100811 | The numerical value of a measurement of total phosphorus (TP). | Provide the numerical value of the measured TP. | | 2 | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | total phosphorus (TP) measurement unit | GENEPIO:0100812 | The units of a measurement of total phosphorus (TP). | Provide the units of the measured TP. | milligrams orthophosphate as phosphorus per liter (mg PO4-P/L) | S | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | total phosphorus (TP) measurement method | GENEPIO:0100813 | The method used to measure total phosphorus (TP). | Provide the name of the procedure or technology used to measure TP. | Merck phosphate spectrophotometric test kit | i | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | fecal contamination indicator | GENEPIO:0100814 | A gene, virus, bacteria, or substance used to measure the sanitary quality of water in regards to fecal contamination. | If a fecal contamination indicator was measured, select it from the picklist. | crAssphage | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | fecal contamination value | GENEPIO:0100815 | The numerical value of a measurement of fecal contamination. | Provide the numerical value of the measured fecal contamination. | 11 | 0 | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | fecal contamination unit | GENEPIO:0100816 | The units of a measurement of fecal contamination. | Provide the units of the measured fecal contamination. | cycle threshold (Ct) / quantification cycle (Cq) | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | fecal contamination method | GENEPIO:0100817 | The method used to measure fecal contamination. | Provide the name of the procedure or technology used to measure fecal contamination. | quantitative PCR assay | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | fecal coliform count value | GENEPIO:0100818 | The numerical value of a measurement of fecal coliforms within a sample. | Provide the numerical value of the measured fecal coliforms. | | 3 | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | fecal coliform count unit | GENEPIO:0100819 | The units of a measurement of fecal coliforms. | Provide the units of the measured fecal coliforms. | most probable number per milliliter (MPN/mL) | | 1.0.0 | 1.0.0 | 1.0.0 |

| Parent Class | Field | Ontology Identifier | Definition | Guidance | Examples | Deprecated Label | Deprecated ID | Version Tracking | | |
|----------------------------------------------|-------------------------------------------------------------------|---------------------|--------------------------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------|---------------------------------------------------|---------------------|-------|-----------------|
| Talent Olass | | Ontology Identifier | Definition | Guidance | Lxamples | Label | | Label | ID | Description/Gui |
| | Colour Code Legend field name in yellow = required | | | | | IMPORTANT: Only will be deprecated, replacement versio term changes in its term will be created | always with n provided. If a meaning, a new | | | dance |
| | field name in purple = recommended field name in white = optional | | | | | | | | | |
| Environmental conditions | nois name in mine opaona. | | | Provide the name of the procedure or technology used to | MPN method via serial dilutions until lack | | | | | |
| and measurements | fecal coliform count method | GENEPIO:0100820 | The method used to measure fecal coliforms | measure fecal coliforms. | of growth | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | urinary contamination indicator | GENEPIO:0100837 | A gene, virus, bacteria, or substance used to measure the sanitary quality of water in regards to urinary contamination. | If a urinary contamination indicator was measured, select it from the picklist. | urobilin | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | urinary contamination value | GENEPIO:0100838 | The numerical value of a measurement of urinary contamination. | Provide the numerical value of the measured urinary contamination. | | 3 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | urinary contamination unit | GENEPIO:0100839 | The units of a measurement of urinary contamination. | Provide the units of the measured urinary contamination. | nanograms per liter | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | urinary contamination method | GENEPIO:0100840 | The method used to measure urinary contamination. | Provide the name of the procedure or technology used to measure urinary contamination. | Urobilin Concentration Test | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | sample temperature value (at collection) | GENEPIO:0100821 | The numerical value of a measurement of temperature of a sample at collection. | Provide the numerical value of the measured temperature. | 2 | 0 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | sample temperature unit (at collection) | GENEPIO:0100822 | The units of a measurement of temperature of a sample at the time of collection. | Provide the units of the measured temperature. | degree Celsius (C) | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | sample temperature value (when received) | GENEPIO:0100823 | The numerical value of a measurement of temperature of a sample upon receipt. | Provide the numerical value of the measured temperature. | 2 | 2 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Environmental conditions and measurements | sample temperature unit (when received) | GENEPIO:0100824 | The units of a measurement of temperature of a sample at the time upon receipt. | Provide the units of the measured temperature. | degree Celsius (C) | | | 1.0.0 | 1.0.0 | 1.0.0 |
| | Sequence Information | GENEPIO:0001441 | | The reason why a sample was originally collected may | | | | | | |
| Sequence information | purpose of sequencing | GENEPIO:0001445 | The reason that the sample was sequenced. | 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 template. The reason for sample collection should be | Travel-associated surveillance | | | 2.0.0 | 2.0.0 | 2.0.0 |
| | | | The description of why the sample was | 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: Assessing public health control measures, Determining early introductions and spread, Investigating airline-related exposures, Investigating remote regions, Investigating | | | | | | |
| Sequence information | purpose of sequencing details | GENEPIO:0001446 | sequenced providing specific details. | health care workers, Investigating schools/universities. Provide the name of the agency, organization or institution | Investigating schools/universities | | | 2.0.0 | 2.0.0 | 2.0.0 |
| Sequence information | sequenced by | GENEPIO:0100416 | The name of the agency, organization or institution responsible for sequencing the isolate's genome. | that performed the sequencing in full (avoid abbreviations). If the information is unknown or cannot be provided, leave blank or provide a null value. | Public Health Agency of Canada (PHAC) [GENEPIO:0100551] | | | 3.1.1 | 3.1.1 | 3.1.1 |
| Sequence information | sequenced by laboratory name | GENEPIO:0100470 | The specific laboratory affiliation of the responsible for sequencing the isolate's genome. | Provide the name of the specific laboratory that that performed the sequencing in full (avoid abbreviations). If the information is unknown or cannot be provided, leave blank or provide a null value. | Topp Lab | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sequence information | sequenced by contact name | GENEPIO:0100471 | The name or title of the contact responsible for follow-up regarding the sequence. | Provide the name of an individual or their job title. As 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 blank or provide a null value. | Joe Bloggs, Enterics Lab Manager | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sequence information | sequenced by contact email | GENEPIO:0100471 | | Provide the email associated with the listed contact. As personnel turnover may render an individual's email obsolete, it is more prefersable to provide an address for a position or lab, to ensure accuracy of information and institutional memory. If the information is unknown or cannot be provided, leave blank or provide a null value. | | | | 1.0.0 | 1.0.0 | 1.0.0 |
| | | | The name of the agency that submitted the | The name of the agency should be written out in full, (with minor exceptions) and be consistent across multiple submissions. For Canadian institutions submitting specimens rather than sequencing data, please put the | | | | | | |
| Sequence information | sequence submitted by | GENEPIO:0001159 | sequence to a database. The email address of the contact responsible | "National Microbiology Laboratory (NML)". The email address can represent a specific individual or | Public Health Ontario (PHO) | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sequence information | sequence submitter contact email | GENEPIO:0001165 | for follow-up regarding the sequence. | laboratory. | RespLab@lab.ca | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sequence information | sequencing date | GENEPIO:0001447 | The date the sample was sequenced. | ISO 8601 standard "YYYY-MM-DD". | 2020-06-2 | 2 | | 1.0.0 | 1.0.0 | 1.0.0 |

| Parent Class | Field | Ontology Identifier | Definition | Guidance | Examples | Deprecated D Label | eprecated ID | Version Tracking Label | ID | Description/Gui |
|-------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------------------|-----------------------------|------------------------------|-------|-----------------|
| | Colour Code Legend field name in yellow = required field name in purple = recommended | | | | | IMPORTANT: Only la will be deprecated, al replacement version p term changes in its m term will be created. | ways with provided. If a | | | dance |
| | field name in white = optional | | | The library name should be unique, and can be an | | | | | | |
| Sequence information | library ID | GENEPIO:0001448 | The user-specified identifier for the library prepared for sequencing. | autogenerated ID from your LIMS, or modification of the isolate ID. | XYZ_123345 | | | 1.0.0 | 1.0.0 | 1.0.0 |
| | | | | Provide the name of the company that created the sequencing instrument by selecting a value from the template pick list. If the information is unknown or cannot be | • | | | | | |
| Sequence information | sequencing platform | GENEPIO:0100473 | sequencing. | provided, leave blank or provide a null value. Provide the model sequencing instrument by selecting a | Illumina [GENEPIO:0001923] | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sequence information | sequencing instrument | GENEPIO:0001452 | The model of the sequencing instrument used. | value from the template pick list. If the information is unknown or cannot be provided, leave blank or provide a null value. | Illumina HiSeq 2500 [GENEPIO:0100117 | 1 | | 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. | 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. | I whole genome sequencing assay | | | 1.0.0 | 1.0.0 | 1.0.0 |
| | | | The name of the DNA library preparation kit used to generate the library being | | | | | | | |
| Sequence information | library preparation kit | GENEPIO:0001450 | sequenced. | Provide the name of the library preparation kit used. Provide the name and version of the procedure or protocol | Nextera XT https://www.protocols.io/view/ncov-2019- | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sequence information | sequencing protocol | GENEPIO:0001454 | The protocol or method used for sequencing | used for sequencing. You can also provide a link to a protocol online. | equencing-protocol-bbmuik6w?version_v arning=no | 1 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sequence information | DNA fragment length | GENEPIO:0100843 | The length of the DNA fragment generated by mechanical shearing or enzymatic digestion for the purposes of library preparation. | Provide the fragment length in base pairs (do not include the units). | 40 | 0 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sequence information | 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 |
| Sequence information | genomic target enrichment method details | GENEPIO:0100967 | Details that provide additional context to the molecular technique used to selectively capture and amplify specific regions of interest from a genome. | | enrichment was done using Illumina Target Enrichment methodology with the Illumina DNA Prep with enrichment kit. | | | 1.0.0 | 1.0.0 | 1.0.0 |
| · | , and the second | | The specifications of the primers (primer sequences, binding positions, fragment size generated etc) used to generate the | Provide the name and version of the primer scheme used to | | | | | | |
| Sequence information | amplicon pcr primer scheme | GENEPIO:0001456 | amplicons to be sequenced. The length of the amplicon generated by | generate the amplicons for sequencing. | artic v3 | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Sequence information | amplicon size Bioinformatics and QC metrics | GENEPIO:0001449 GENEPIO:0001457 | PCR amplification. | Provide the amplicon size expressed in base pairs. | 30 | 0 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Bioinformatics and QC metrics | quality control method name | GENEPIO:0001457 | The name of the method used to assess whether a sequence passed a predetermined quality control threshold. | 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 Clittlub 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 provided. | r | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Bioinformatics and QC | | | The version number of the method used to assess whether a sequence passed 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 | | | | | | |
| metrics | quality control method version | GENEPIO:0100558 | predetermined quality control threshold. | version numbers using a semi-colon. Select a value from the pick list provided. If a desired value | 1.2.3 | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Bioinformatics and QC metrics | quality control determination | GENEPIO:0100559 | The determination of a quality control assessment. | is missing, submit a new term request to the PHA4GE QC 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 issues | GENEPIO:0100560 | The reason contributing to, or causing, a low quality determination in a quality control assessment. | Select a value from the pick list provided. If a desired value 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 | | | The details surrounding a low quality determination in a quality control | Provide notes or details regarding QC results using free | CT value of 39. Low viral load. Low DNA | | | | | |
| metrics | quality control details | GENEPIO:0100561 | assessment. The method used for raw data processing | text. Provide the software name followed by the version or a link | concentration after amplification. | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Bioinformatics and QC metrics | raw sequence data processing method | GENEPIO:0001458 | such as removing barcodes, adapter trimming, filtering etc. | to the github protocol e.g. Trimmomatic v. 0.38, Porechop v. 0.2.3 | Porechop 0.2.3 | | | 1.0.0 | 1.0.0 | 1.0.0 |

| | | | | | | Deprecated De | eprecated ID | Version | | |
|--------------------------------------|-------------------------------------------------------------------|---------------------|---------------------------------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------|-----------------------------|----------|-------|-------------------------|
| Parent Class | Field | Ontology Identifier | Definition | Guidance | Examples | Label | | Tracking | | |
| | Colour Code Legend | | | | | | | Label | ID | Description/Gu dance |
| | field name in yellow = required | | | | | IMPORTANT: Only lab will be deprecated, alw replacement version p term changes in its me term will be created. | vays with provided. If a | | | |
| | field name in purple = recommended field name in white = optional | | | | | | | | | |
| 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 | | | 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. | Provide the name and version number, with the file extension, of the processed genome sequence file e.g. a consensus sequence FASTA file or a genome assembly file | mpxyassembly fasta | | | 3.1.1 | 3.1.1 | 3.1.1 |
| Bioinformatics and QC | genome sequence file path | GENEPIO:0101716 | | Provide the filepath of the genome sequence FASTA file. | /User/Documents/ViralLab/Data/mpxvass embly.fasta | | | 311 | 3.1.1 | 3.1.1 |
| Bioinformatics and QC metrics | sequence assembly software name | GENEPIO:0100825 | The name of the software used to assemble a sequence. | | SPAdes Genome Assembler, Canu, wtdbg2, 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. | Provide the version of the software used to assemble the sequence. | 3.15.5 | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Bioinformatics and QC metrics | consensus sequence software name | GENEPIO:0100626 | The name of the software used to generate the consensus sequence. | Provide the name of the software used to generate the consensus sequence. | iVar | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Bioinformatics and QC metrics | consensus sequence software version | GENEPIO:0001469 | • | Provide the version of the software used to generate the consensus sequence. | 1. | 2 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Bioinformatics and QC metrics | breadth of coverage value | GENEPIO:0001409 | The percentage of the reference genome covered by the sequenced data, to a prescribed depth. | Provide value as a percent. | 9 | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Bioinformatics and QC metrics | depth of coverage value | GENEPIO:0001474 | The average number of reads representing a given nucleotide in the reconstructed sequence. | | 40 | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Bioinformatics 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. | 10 | 0 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Bioinformatics and QC | | | The percentage of expected genes identified in the genome being sequenced. Missing genes indicate missing genomic regions | Provide the genome completeness as a percent (no need to | | | | | | |
| metrics Bioinformatics and QC | genome completeness | GENEPIO:0100844 | (incompleteness) in the data. The number of total base pairs generated by | include units). | 8 | 5 | | 1.0.0 | 1.0.0 | 1.0.0 |
| metrics Bioinformatics and QC | number of base pairs sequenced | GENEPIO:0001482 | the sequencing process. The total number of non-unique reads | Provide a numerical value (no need to include units). | 38756 | 6 | | 1.0.0 | 1.0.0 | 1.0.0 |
| metrics Bioinformatics and QC | number of total reads | GENEPIO:0100827 | generated by the sequencing process. The number of unique reads generated by | Provide a numerical value (no need to include units). | 42386 | 7 | | 1.0.0 | 1.0.0 | 1.0.0 |
| metrics | number of unique reads | GENEPIO:0100828 | the sequencing process. | Provide a numerical value (no need to include units). | 24823 | 6 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Bioinformatics and QC metrics | minimum post-trimming read length | GENEPIO:0100829 | The threshold used as a cut-off for the minimum length of a read after trimming. | Provide a numerical value (no need to include units). | 15 | 0 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Bioinformatics and QC metrics | number of contigs | GENEPIO:0100937 | The number of contigs (contiguous sequences) in a sequence assembly. | Provide a numerical value. | 1 | 0 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Bioinformatics 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 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Bioinformatics and QC metrics | Ns per 100 kbp | GENEPIO:0001484 | The number of ambiguous bases (Ns) normalized per 100 kilobasepairs (kbp). | Provide a numerical value (no need to include units). | 34 | 2 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Bioinformatics and QC | | | The length of the shortest read that, together with other reads, represents at least 50% of | | | | | | | |
| metrics Bioinformatics and QC | N50 | GENEPIO:0100938 | the nucleotides in a set of sequences. The percent of the total number of reads identified as contamination (not belonging to | Provide the N50 value in Mb. Provide the percent contamination value (no need to | 15 | U | | 1.0.0 | 1.0.0 | 1.0.0 |
| metrics | percent read contamination | GENEPIO:0100845 | the target organism) in a sequence dataset. The length of the genome generated by | include units). | | 2 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Bioinformatics and QC metrics | sequence assembly length | GENEPIO:0100846 | assembling reads using a scaffold or by reference-based mapping. | Provide a numerical value (no need to include units). | 3427 | 2 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Bioinformatics and QC metrics | consensus genome length | GENEPIO:0001483 | The length of the genome defined by the most common nucleotides at each position. | Provide a numerical value (no need to include units). | 3867 | 7 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Bioinformatics and QC metrics | reference genome accession | GENEPIO:0001485 | A persistent, unique identifier of a genome database entry. | Provide the accession number of the reference genome. | NC_045512.2 | | | 1.0.0 | 1.0.0 | 1.0.0 |
| | Taxonomic identification information | GENEPIO:0101082 | | - | | | | | | |
| Taxonomic identification information | read mapping software name | GENEPIO:0100832 | The name of the software used to map sequence reads to a 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 |
| Taxonomic identification | | | The version of the software used to map sequence reads to a reference genome or | | | | | | | 1.0.0 |
| niormation | read mapping software version | GENEPIO:0100833 | set of reference genes. | Provide the version number of the read mapping software. | 2.0.1 | | | 1.0.0 | 1.0.0 | 1.0.0 |

| Parent Class | Field | Ontology Identifier | Definition | Guidance | Examples | Deprecated Label | Deprecated ID | Tracking | | |
|-----------------------------------------|----------------------------------------------------------------------|------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------------------------------------------------------|-------------------------------------------|---------------------------------------|----------|-------|--------------------------|
| | Colour Code Legend | | | | | | | Label | ID | Description/Gui dance |
| | field name in yellow = required | | | | | will be deprecated replacement version | on provided. If a s meaning, a new | | | |
| | field name in purple = recommended | | | | | | | | | |
| | field name in white = optional | | | | | | | | | |
| Taxonomic identification information | read mapping software name | GENEPIO:0100832 | The name of the software used to map sequence reads to a 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 |
| Taxonomic identification information | read mapping software version | GENEPIO:0100833 | The version of the software used to map sequence reads to a reference genome or set of reference genes. | Provide the version number of the read mapping software. | 2.5.1 | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Taxonomic identification information | taxonomic reference database name | GENEPIO:0100834 | The name of the taxonomic reference database used to identify the organism. | Provide the name of the taxonomic reference database. | NCBITaxon | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Taxonomic identification information | taxonomic reference database version | GENEPIO:0100835 | The version of the taxonomic reference database used to identify the organism. | Provide the version number of the taxonomic reference database | 1.3 | 2 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Taxonomic identification information | taxonomic analysis report filename | GENEPIO:0100033 | The filename of the report containing the results of a taxonomic analysis. | Provide the filename of the report containing the results of the taxonomic analysis. | WWtax report Feb1 2024.doc | 3 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Taxonomic identification information | , | GENEPIO:0101074 | The date a taxonomic analysis was performed. | Providing the date that an analyis was performed can help provide context for tool and reference database versions. Provide the date that the taxonomic analysis was performed | | 1 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Taxonomic identification | taxonomic analysis date | | A description of the criteria used to map | in ISO 8601 format, i.e. "YYYY-MM-DD". | | ı | | | | |
| information | read mapping criteria AMR detection information | GENEPIO:0100836 GENEPIO:0100479 | reads to a reference sequence. | Provide a description of the read mapping criteria. | Phred score >20 | | | 1.0.0 | 1.0.0 | 1.0.0 |
| | AMR detection information | GENEPIO:0100479 | The name of the software used to perform ar | | | | | | | |
| AMR detection information | AMR analysis software name | GENEPIO:0101076 | in silico antimicrobial resistance determinant identification/analysis. | Provide the name of the software used for AMR analysis. | Resistance Gene Identifier | | | 1.0.0 | 1.0.0 | 1.0.0 |
| AMR detection information | AMR analysis software version | GENEPIO:0101077 | The version number of the software used to perform an in silico antimicrobial resistance determinant idenrtification/analysis. | Provide the version number of the software used for AMR analysis. | 6.0.3 | | | 1.0.0 | 1.0.0 | 1.0.0 |
| AMR detection information | AMR reference database name | GENEPIO:0101078 | determinant identification/analysis. | Provide the name of the reference database used for AMR analysis. | Comprehensive Antibiotic Resistance Database (CARD) | | | 1.0.0 | 1.0.0 | 1.0.0 |
| AMR detection information | AMR reference database version | GENEPIO:0101079 | The version number of the reference database used to perform an in silico antimicrobial resistance determinant identification/analysis. | Provide the version number of the reference database used for AMR analysis. | 3.2.9 | | | 1.0.0 | 1.0.0 | 1.0.0 |
| AMR detection information | AMR analysis report filename | GENEPIO:0101080 | The filename of the report containing the results of an in silico antimicrobial resistance analysis. | Provide the filename of the report containing the results of the AMR analysis. | WWAMR_report_Feb1_2024.doc | | | 1.0.0 | 1.0.0 | 1.0.0 |
| | Lineage/clade information | GENEPIO:0001498 | | | | | | | | |
| | | | | Provide the Pangolin or Nextstrain lineage/clade name. Multiple lineages/clades can be provided, separated by a | | | | | | |
| Lineage/clade information | lineage/clade name | GENEPIO:0001500 | The name of the lineage or clade. The name of the software used to determine | semicolon. Provide the name of the software used to determine the | B.1.1.7 | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Lineage/clade information | lineage/clade analysis software name | GENEPIO:0001501 | the lineage/clade. The version of the software used to | lineage/clade. Provide the version of the software used ot determine the | Freyja | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Lineage/clade information | lineage/clade analysis software version Pathogen diagnostic testing | GENEPIO:0001502 GENEPIO:0001506 | determine the lineage/clade. | lineage/clade. | 1.5.0 | | | 1.0.0 | 1.0.0 | 1.0.0 |
| | Fathogen diagnostic testing | GENEFIO.0001300 | The name of the assay target used in the | The specific genomic region, sequence, or variant targeted by the assay in a diagnostic RT-PCR test. This may include parts of a gene, non-coding regions, or other genetic elements that serve as a marker for detecting the presence | | | | | | |
| Pathogen diagnostic testing | assay target name 1 | GENEPIO:0101206 | diagnostic RT-PCR test. | of a pathogen or other relevant entities. | | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Pathogen diagnostic testing | assay target details | GENEPIO:0102040 | Describe any details of the assay target. | Provide details that are applicable to the assay used for the diagnostic test. | | | | 3.1.1 | 3.1.1 | 3.1.1 |
| Pathogen diagnostic testing | gene symbol 1 | GENEPIO:0102041 | The symbol of the gene used in the diagnostic RT-PCR test. | Select a gene name value from the pick list provided. | E gene (orf4) | | | 3.1.1 | 3.1.1 | 1.0.0 |
| Pathogen diagnostic testing | gene symbol 1 | GENEPIO:0102041 | The symbol of the gene used in the diagnostic RT-PCR test. | Provide the full name of the gene used in the test. Standardized gene names can be found in the Gene Ontology using this look-up service: https://bit.ly/2Sq1Lbl | gyrase A | | | 3.1.1 | 3.1.1 | 1.0.0 |
| Pathogen diagnostic testing | diagnostic target presence 1 | GENEPIO:0100962 | The binary value of the result from a diagnostic test. | Select a value from the pick list provided, to describe whether a target was determined to be present or absent within a sample. | diagnostic target present | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Pathogen diagnostic testing | diagnostic measurement value 1 | GENEPIO:0100963 | The value of the result from a diagnostic test | Provide the numerical result of a diagnostic test (no need to include units). | 100 | 0 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Pathogen diagnostic testing | diagnostic measurement unit 1 | GENEPIO:0100964 | The unit of the result from a diagnostic test. | Select a value from the pick list provided, to describe the units of the given diagnostic test. | cycle threshold (Ct) | | | 1.0.0 | 1.0.0 | 1.0.0 |

| Parent Class | Field | Ontology Identifier | Definition | Guidance | Examples | Deprecated Label | Deprecated ID | Version Tracking | | |
|-----------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------|------------------------------------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----------------------------------------|-------------------------------------------------------------------------------------------------------------|---------------------------------------------------|---------------------|-------|--------------------------|
| | Colour Code Legend | | | | | | | Label | ID | Description/Gui dance |
| | field name in yellow = required | | | | | IMPORTANT: Only will be deprecated, replacement versio term changes in its term will be created | always with n provided. If a meaning, a new | | | garice |
| | field name in purple = recommended | | | | | | | | | |
| Dathagan diagnostic testing | field name in white = optional diagnostic measurement method 1 | GENEPIO:0100965 | The method by which a diagnostic result was determined. | Select a value from the pick list provided to describe the method used for a given diagnostic test. | aPCR | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Pathogen diagnostic testing | , and the second | SENEI IS. S 100000 | The name of the assay target used in the diagnostic RT-PCR test. | The specific genomic region, sequence, or variant targeted by the assay in a diagnostic RT-PCR 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. | q. Giv | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Pathogen diagnostic testing | , , | | Describe any details of the assay target. | Provide details that are applicable to the assay used for the diagnostic test. | | | | 3.1.1 | 3.1.1 | 3.1.1 |
| Pathogen diagnostic testing | gene symbol 2 | | The symbol of the gene used in the diagnostic RT-PCR test. | Select a gene name value from the pick list provided. | E gene (orf4) | | | 3.1.1 | 3.1.1 | 1.0.0 |
| Pathogen diagnostic testing | gene symbol 2 | | The symbol of the gene used in the diagnostic RT-PCR test. | Provide the full name of the gene used in the test. Standardized gene names can be found in the Gene Ontology using this look-up service: https://bit.ly/2Sq1Lbl | gyrase A | | | 3.1.1 | 3.1.1 | 1.0.0 |
| | diagnostic target presence 2 | | The binary value of the result from a diagnostic test. | Select a value from the pick list provided, to describe whether a target was determined to be present or absent within a sample. | diagnostic target present | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Pathogen diagnostic testing | diagnostic measurement value 2 | | The value of the result from a diagnostic test | | 100 | 0 | | 1.0.0 | 1.0.0 | 1.0.0 |
| Pathogen diagnostic testing | diagnostic measurement unit 2 | | The unit of the result from a diagnostic test. | Select a value from the pick list provided, to describe the units of the given diagnostic test. | cycle threshold (Ct) | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Pathogen diagnostic testing | diagnostic measurement method 2 | | The method by which a diagnostic result was determined. | Select a value from the pick list provided to describe the method used for a given diagnostic test. | qPCR | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Pathogen diagnostic testing | assay target name 3 | | The name of the assay target used in the diagnostic RT-PCR test. | The specific genomic region, sequence, or variant targeted by the assay in a diagnostic RT-PCR 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. | | | | 1.0.0 | 1.0.0 | 1.0.0 |
| Pathogen diagnostic testing | assay target details 3 | | Describe any details of the assay target. | Provide details that are applicable to the assay used for the diagnostic test. | | | | 3.1.1 | 3.1.1 | 3.1.1 |
| Pathogen diagnostic testing | gene symbol 3 | | The symbol of the gene used in the diagnostic RT-PCR test. | Select a gene name value from the pick list provided. | E gene (orf4) | | | 3.1.1 | 3.1.1 | 1.0.0 |
| | - | | The symbol of the gene used in the | Provide the full name of the gene used in the test. Standardized gene names can be found in the Gene | | | | | | |
| Pathogen diagnostic testing | | | diagnostic RT-PCR test. The binary value of the result from a | Ontology using this look-up service: https://bit.ly/2Sq1Lbl Select a value from the pick list provided, to describe whether a target was determined to be present or absent | gyrase A | | | 3.1.1 | 3.1.1 | 1.0.0 |
| 0 0 0 | diagnostic target presence 3 diagnostic measurement value 3 | | diagnostic test. The value of the result from a diagnostic test | within a sample. Provide the numerical result of a diagnostic test (no need to include units) | diagnostic target present | 0 | | 1.0.0 | 1.0.0 | 1.0.0 |
| 0 0 0 | diagnostic measurement value 3 | | The unit of the result from a diagnostic test. | Select a value from the pick list provided, to describe the units of the given diagnostic test. | cycle threshold (Ct) | | | 1.0.0 | 1.0.0 | 1.0.0 |
| | diagnostic measurement method 3 | | | Select a value from the pick list provided to describe the method used for a given diagnostic test. | qPCR | | | 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 collection, sequence generation, analysis, and data submission. | Include the first and last names of all individuals that should be attributed, separated by a semicolon. | Tejinder Singh; Fei Hu; Joe Blogs | | | 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 automatically generated in this field after the user utilizes the "validate" function. This information will be generated regardless as to whether the row is valid of not. | DataHarmonizer v3.3.3, Influenza v1.0.0 | | | 1.0.0 | 1.0.0 | 1.0.0 |