| Parent Class   | Field  | Ontology Identifier              | Definition   | Guidance   | Examples  | Deprecated Label   | Deprecated ID | Version<br>Tracking<br>Label | ID    | Description/Gui |
|--|--|----------------------------------|--|--|---|--|---------------|------------------------------|-------|-----------------|
|  | Colour Code Legend field name in yellow = required |                                  |  |  |   | IMPORTANT: Only<br>labels and/or IDs will be<br>deprecated, always with<br>replacement version<br>provided. If a term<br>changes in its meaning,<br>a new term will be<br>created. |               | Laber                        | U     | dance           |
|  | field name in purple = recommended                 |                                  |  |  |   |  |               |                              |       |                 |
|  | field name in white = optional                     |                                  |  |  |   |  |               |                              |       |                 |
| Sample collection and processing  Sample collection and processing | sample_collector_sample_ID                         | GENEPIO:0001150  GENEPIO:0001123 | The user-defined name for the sample.  | The sample_ID should represent the identifier assigned to the sample at time of collection, for which all the descriptive information applies. If the original sample_ID is unknown or cannot be provided, leave blank or provide a null value.  | ABCD123   |  |               | 3.0.0                        | 3.0.0 | 3.0.0           |
| Sample collection and processing                                   | alternative_sample_ID                              | GENEPIO:0100427                  | An alternative sample_ID assigned to the sample by another organization.             | Alternative identifiers assigned to the sample should be tracked along with original IDs to establish chain of custody. Alternative sample 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]. Agency short forms include the following: Public Health Agency of Canada: PHAC Canadian Food Inspection Agency: CFIA Agriculture and Agri-Food Canada: AAFC Fisheries and Oceans Canada: DFO Environment and Climate Change Canada: ECCC Health Canada: HC Multiple identifiers can be provided and separated by semi-colons. If the information is unknown or cannot be provided, leave blank or provide a null value. |   |  |               | 3.0.0                        | 3.0.0 | 3.0.0           |
| Sample collection and processing                                   | sample_collected_by                                | GENEPIO:0001153                  | The name of the organization with which the sample collector is affiliated.          | Provide the name of the agency, organization or institution that collected the sample in full (avoid abbreviations). If the information is unknown or cannot be provided, leave blank or provide a null value.   | Public Health Agency of<br>Canada (PHAC)<br>[GENEPIO:0100551] |  |               | 3.0.0                        | 3.0.0 | 3.0.0           |
| 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  |  |               | 3.0.0                        | 3.0.0 | 3.0.0           |
|  |  |                                  | The name of the project/initiative/program for                                       | Provide the name of the project and/or the project ID here. If the information is unknown or cannot be provided, leave blank or provide  | Watershed Project (HA-120)                                    |  |               |                              |       |                 |
| Sample collection and processing  Sample collection and processing | sample_collection_project_name sample_plan_name    | GENEPIO:0100429  GENEPIO:0100430 | ,  | a null value.  Provide the name of the sample plan used for sample collection. If the information is unknown or cannot be provided, leave blank or provide a null value.   | Baseline Study in Broiler                                     |  |               | 3.0.0                        | 3.0.0 | 3.0.0           |
| Sample collection and processing                                   | sample_plan_ID                                     | GENEPIO:0100431                  | The identifier of the study design for a surveillance project.                       | Provide the identifier of the sample plan used for sample collection. If the information is unknown or cannot be provided, leave blank or provide a null value.  | _   |  |               | 3.0.0                        | 3.0.0 | 3.0.0           |
| 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.  | Enterics Lab Manager  |  |               | 3.0.0                        | 3.0.0 | 3.0.0           |
| Sample collection and processing                                   | sample_collector_contact_email                     | GENEPIO:0001156                  | The email address of the contact responsible for follow-up regarding the sample.     | Provide the email associated with the listed contact. As personnel turnover may render an individual's email obsolete, it is more preferable 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.  |   |  |               | 3.0.0                        | 3.0.0 | 3.0.0           |

| Parent Class                     | Field                                      | Ontology Identifier | Definition  | Guidance   | Examples  | Deprecated Label | Deprecated ID | Version<br>Tracking |        |                |
|----------------------------------|--|---------------------|---|--|---|------------------|---------------|---------------------|--------|----------------|
|                                  |  |                     |   |  |   |                  |               | Label               | ID     | Description/Gu |
|                                  | Colour Code Legend                         |                     |   | The reason a completure collected may  | Surveillance  |                  |               |                     |        | dance          |
|                                  |  |                     |   | 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 Diagnostic testing. The reason why a sample was originally collected may differ from the   | [GENEPIO:0100004]   |                  |               |                     |        |                |
|                                  |  |                     |   | reason why it was selected for sequencing,<br>which should be indicated in the "purpose of   |   |                  |               |                     |        |                |
| Sample collection and processing | purpose_of_sampling                        | GENEPIO:0001198     | collected.  | sequencing" field.  If there was presampling activity that would   | Antimicrobial pre-treatment   |                  |               | 3.0.0               | 3.0.0  | 3.0.0          |
| Sample collection and processing | presampling_activity                       | GENEPIO:0100433     | The activities or variables introduced upstream of sample collection that may affect the sample collected.              | If thele was presamping activity that would affect the sample prior to collection (this is different than sample processing which happens after the sample is collected), provide the experimental activities by selfecting one or more values from the template pick list. If the information is unknown or cannot be provided, leave blank or provide a null value.  | [GENEPIO:0100537]   |                  |               | 3.0.0               | 3.0.0  | 3.0.0          |
|                                  |  |                     | The details of the activities or  |  | Chicken feed containing X   |                  |               |                     |        |                |
| Sample collection and processing | presampling_activity_details               | GENEPIO:0100434     | variables introduced upstream of<br>sample collection that may affect<br>the sample collected.                          | Briefly describe the presampling activity details using free text.   | amount of novobiocin was fed<br>to chickens for 72 hours prior<br>to collection of litter.  |                  |               | 3.0.0               | 3.0.0  | 3.0.0          |
|                                  | experimental _protocol_field               | GENEPIO:0101029     | The name of the overarching   | Provide the name of the methodology used in  | OneHealth2024_protocol  |                  |               |                     |        |                |
| Sample collection and processing |  |                     | experimental methodology that<br>was used to process the<br>biomaterial.  | your study. If available, provide a link to the protocol.  |   |                  |               | 11.1.1              | 11.1.1 | 11.1.1         |
| Sample collection and processing | experimental_specimen_role_type            | GENEPIO:0100921     | The type of role that the sample  | Samples can play different types of roles in   | Positive experimental control   |                  |               | 11.1.1              | 11.1.1 | 11.1.1         |
| Sample collection and processing |  |                     | represents in the experiment.   | experiments. A sample under study in one experiment may act as a control or be a replicate of another sample in another  | [GENEPIO:0101018]   |                  |               | 11.1.1              | 11.1.1 | 11.1.1         |
| Sample collection and processing | specimen_processing                        | GENEPIO:0100435     | The processing applied to samples post-collection, prior to further testing, characterization, or isolation procedures. | Provide the sample processing information by selecting a value from the template pick list. If the information is unknown or cannot be provided, leave blank or provide a null value.  |   |                  |               | 3.0.0               | 3.0.0  | 3.0.0          |
| bampic collection and processing | nucleic acid extraction method             | GENEPIO:0100939     | The process used to extract   | Briefly describe the extraction method used.   | Direct wastewater RNA   |                  |               | 0.0.0               | 0.0.0  | 0.0.0          |
|                                  |  |                     | genomic material from a sample.   | ,  | capture and purification via the<br>"Sewage, Salt, Silica and<br>Salmonella (4S)" method v4 |                  |               |                     |        |                |
| Sample collection and processing |  | OENEDIO 0400770     | The life word to entered account  | Described the second of the se | found at  |                  |               | 11.1.1              | 11.1.1 | 11.1.1         |
|                                  | nucleic acid extraction kit                | GENEPIO:0100772     | The kit used to extract genomic material from a sample  | Provide the name of the genomic extraction kit used.   | Kit   |                  |               |                     |        |                |
| Sample collection and processing |  |                     |   |  |   |                  |               | 11.1.1              | 11.1.1 | 11.1.1         |
| Sample collection and processing | geo_loc_name (country)                     | GENEPIO:0001181     | The country of origin of the sample.  | Provide the name of the country where the<br>sample was collected. Use the controlled<br>vocabulary provided in the template pick list. It<br>the information is unknown or cannot be<br>provided, provide a null value.   | Canada [GAZ:00002560]<br>f  |                  |               | 3.0.0               | 3.0.0  | 3.0.0          |
|                                  |  |                     | The state/province/territory of   | Provide the name of the province/state/region where the sample was collected. If the information is unknown or cannot be provided,   | British Columbia<br>[GAZ:00002562]  |                  |               |                     |        |                |
| Sample collection and processing | geo_loc_name (state/province/region)       | GENEPIO:0001185     | origin of the sample. The name of a specific  | provide a null value.  Provide the name of the specific geographical   | Credit River  |                  |               | 3.0.0               | 3.0.0  | 3.0.0          |
| Sample collection and processing | geo_loc_name (site)                        | GENEPIO:0100436     | geographical location e.g. Credit<br>River (rather than river).   | site using a specific noun (a word that names a certain place, thing).   | Credit River  |                  |               | 3.0.0               | 3.0.0  | 3.0.0          |
| Sample collection and processing | food_product_origin geo_loc_name (country) | GENEPIO:0100437     | The country of origin of a food product.  | If a food product was sampled and the food product was sampled and outside of Canada, provide the name of the country where the food product originated by selecting a value from the template pick list. If the information is unknown or cannot be provided, leave blank or provide a null value.  |   |                  |               | 3.0.0               | 3.0.0  | 3.0.0          |
| Sample collection and processing | host_origin geo_loc_name (country)         | GENEPIO:0100438     | The country of origin of the host.  | If a sample is from a human or animal host that originated from outside of Canada, provide the the name of the country where the host originated by selecting a value from the template pick list. If the information is unknown or cannot be provided, leave blank or provide a null value.   |   |                  |               | 3.0.0               | 3.0.0  | 3.0.0          |
|                                  |  |                     | The latitude coordinates of the geographical location of sample   | If known, provide the degrees latitude. Do NOT simply provide latitude of the institution if this is not where the sample was collected, not the centre of the city/region where the sample was collected as this falsely implicates an existing geographical location and creates data inaccuracies. If the information is unknown or cannot be provided, leave blank or  |   |                  |               |                     |        |                |
| Sample collection and processing | geo_loc latitude                           | GENEPIO:0100309     | collection.   | provide a null value.  |   |                  |               | 3.0.0               | 3.0.0  | 3.0.0          |

| Parent Class                     | Field                             | Ontology Identifier              | Definition  | Guidance  | Examples  | Deprecated Label | Deprecated ID | Version<br>Tracking | ID    | Description (C. d        |
|----------------------------------|-----------------------------------|----------------------------------|---|---|---|------------------|---------------|---------------------|-------|--------------------------|
|                                  | Colour Code Legend                |                                  |   |   |   |                  |               | Label               | ID    | Description/Gui<br>dance |
| Sample collection and processing | geo_loc longitude                 | GENEPIO:0100310                  | The longitude coordinates of the geographical location of sample collection.  | If known, provide the degrees longitude. Do NOT simply provide longitude of the institution if this is not where the sample was collected, nor the centre of the city/region where the sample was collected as this falsely implicates an existing geographical location and creates data inaccuracies. If the information is unknown or cannot be provided, leave blank or provide a null value.   |   |                  |               | 3.0.0               | 3.0.0 | 3.0.0                    |
| Sample collection and processing | sample_collection_date            | GENEPIO:0001174                  | The date on which the sample was collected.   | Provide the date according to the ISO 8601 standard "YYYY-MM-DD", "YYYY-MM" or "YYYY".  | 2020-10-30  |                  |               | 3.0.0               | 3.0.0 | 3.0.0                    |
| Sample collection and processing | sample_collection_date_precision  | GENEPIO:0001177                  | The precision to which the "sample collection date" was provided.   | Provide the precision of granularity to the<br>"day", "month", or "year" for the date provided<br>in the "sample collection date" field. The<br>"sample collection date" will be truncated to<br>the precision specified upon export, "day" for<br>"YYYY-MM-DD", "month" for "YYYY-MM", or<br>"year" for "YYYY".  | day [UO:0000033]  |                  |               | 7.6.4               | 7.6.4 | 7.6.4                    |
| Sample collection and processing | sample_received_date              | GENEPIO:0001179                  | The date on which the sample was received.  | Provide the date according to the ISO 8601 standard "YYYY-MM-DD", "YYYY-MM" or "YYYY".  | 2020-11-15  |                  |               | 3.0.0               | 3.0.0 | 3.0.0                    |
| Sample collection and processing | original_sample_description       | GENEPIO:0100439                  | The original sample description provided by the sample collector.   | Provide the sample description provided by<br>the original sample collector. The original<br>description is useful as it may provide further<br>details, or can be used to clarify higher level<br>classifications.   | RTE Prosciutto from deli  |                  |               | 3.0.0               | 3.0.0 | 3.0.0                    |
| Sample collection and processing | environmental_site                | GENEPIO:0001232                  | An environmental location may describe a site in the natural or built environment e.g. hospital, wet market, bat cave.                    | If applicable, select the standardized term and ontology ID for the environmental site from the picklist provided. Multiple values can be provided, separated by a semi-colon.  |   |                  |               | 3.0.0               | 3.0.0 | 3.0.0                    |
| Sample collection and processing | animal_or_plant_population        | GENEPIO:0100443                  | The type of animal or plant population inhabiting an area.  | This field should be used when a sample is taken from an environmental location inhabited by many individuals of a specific type, rather than describing a sample taken from one particular host. If applicable, provide the standardized term and ontology ID for the animal or plant population name. The standardized term can be sourced from this look-up service: https://www.ebi.ac.uk/ols/ontologies/genepio. If not applicable, leave blank. |   |                  |               | 3.0.0               | 3.0.0 | 3.0.0                    |
| Sample collection and processing | environmental_material            | GENEPIO:0001223                  | A substance obtained from the<br>natural or man-made<br>environment e.g. soil, water,<br>sewage, door handle, bed<br>handrail, face mask. | If applicable, select the standardized term and ontology ID for the environmental material from the picklist provided. Multiple values can be provided, separated by a semi-colon.  | [ENVO:00002001]; Broom  |                  |               | 3.0.0               | 3.0.0 | 3.0.0                    |
| Sample collection and processing | anatomical_material               | GENEPIO:0001223                  | A substance obtained from an anatomical part of an organism e.g. tissue, blood.   | An anatomical material is a substance taken<br>from the body. If applicable, select the<br>standardized term and ontology ID for the<br>anatomical material from the picklist provided.<br>Multiple values can be provided, separated by<br>a semi-colon  | Tissue [UBERON:0000479];<br>Blood [UBERON:0000178]  |                  |               | 3.0.0               | 3.0.0 | 3.0.0                    |
| Sample collection and processing | body_product                      | GENEPIO:0001216                  | A substance excreted/secreted from an organism e.g. feces, urine, sweat.  | A body product is a substance produced by the body but meant to be excreted/secreted (i.e. not part of the body). If applicable, select the standardized term and ontology ID for the body product from the picklist provided. Multiple values can be provided, separated by a semi-colon.  | Feces [UBERON:0001988];<br>Urine [UBERON:0001088]   |                  |               | 3.0.0               | 3.0.0 | 3.0.0                    |
| Sample collection and processing | anatomical part                   | GENEPIO:0001214                  | An anatomical part of an organism e.g. oropharynx.  | An anatomical part is a structure or location in the body. If applicable, select the standardized term and ontology ID for the anatomical material from the picklist provided. Multiple values can be provided, separated by a semi-colon.  |   |                  |               | 3.0.0               | 3.0.0 | 3.0.0                    |
| Sample collection and processing | anatomical_part anatomical_region | GENEPIO:0001214  GENEPIO:0100700 | A 3D region in space without well-defined compartmental boundaries.   | semi-coion.  This field captures more granular spatial information on a host anatomical part e.g. dorso-lateral region vs back. Select a term from the picklist.  | Dorso-lateral region<br>[BSPO:0000080]  |                  |               | 7.6.4               | 7.6.4 | 7.6.4                    |
| Sample collection and processing | food_product                      | GENEPIO:0100444                  | A material consumed and digested for nutritional value or enjoyment.  | This field includes animal feed. If applicable, select the standardized term and ontology ID for the anatomical material from the picklist provided. Multiple values can be provided, separated by a semi-colon.  | Feather meal<br>[FOODON:00003927]; Bone<br>meal [ENVO:02000054];<br>Chicken breast<br>[FOODON:00002703] |                  |               | 3.0.0               | 3.0.0 | 3.0.0                    |

| Parent Class   | Field   | Ontology Identifier                | Definition   | Guidance  | Examples  | Deprecated Label D | eprecated ID | Version<br>Tracking<br>Label | ID     | Description/Cu           |
|--|---|------------------------------------|--|---|---|--------------------|--------------|------------------------------|--------|--------------------------|
|  | Colour Code Legend                                  |                                    |  |   |   |                    |              | Labei                        | ID     | Description/Gui<br>dance |
|  |   |                                    | Any characteristic of the food product pertaining to its state, processing, a label claim, or  | Provide any characteristics of the food product including whether it has been cooked, processed, preserved, any known information about its state (e.g. raw, ready-to-eat), any known information about its containment (e.g. canned), and any information about a label  | Food (chopped)<br>[FOODON:00002777];<br>Ready-to-eat (RTE)<br>[FOODON:03316636] |                    |              |                              |        |                          |
| Sample collection and processing                                     | food_product_properties                             | GENEPIO:0100445                    | implications for consumers.  | claim (e.g. organic, fat-free).  Provide the common name of the animal. If not  |   |                    |              | 3.0.0                        | 3.0.0  | 3.0.0                    |
| Sample collection and processing                                     | animal_source_of_food                               | GENEPIO:0100446                    | The animal from which the food<br>product was derived.   | applicable, leave blank. Multiple entries can be provided, separated by a comma.  |   |                    |              | 3.0.0                        | 3.0.0  | 3.0.0                    |
| Sample collection and processing                                     | food_product_production_stream                      | GENEPIO:0100699                    | A production pathway incorporating the processes, material entities (e.g. equipment, animals, locations), and conditions that participate in the generation of a food commodity. | Provide the name of the agricultural production stream from the picklist.   | Beef cattle production stream [FOODON:03000452]                                 | production_stream  |              | 8.8.7                        | 7.6.4  | 8.8.7                    |
| Sample collection and processing                                     | food_packaging                                      | GENEPIO:0100099                    | The type of packaging used to contain a food product.  | If known, provide information regarding how the food product was packaged.  | Plastic tray or pan<br>[FOODON:03490126]  | production_stream  |              | 3.0.0                        | 3.0.0  | 3.0.0                    |
| Sample collection and processing                                     | food_quality_date                                   | GENEPIO:0100615                    | A date recommended for the use   | This date is typically labeled on a food product as "best if used by", best by", "use by", or "freeze by" e.g. 5/24/2020. If the date is known, leave blank or provide a null value.  |   |                    |              | 6.4.2                        | 6.4.2  | 6.4.2                    |
| Sample collection and processing                                     | food_packaging_date                                 | GENEPIO:0100616                    | A food product's packaging date as marked by a food manufacturer or retailer.  | The packaging date should not be confused with, nor replaced by a Best Before date or other food quality date. If the date is known, leave blank or provide a null value.   | 2020-05-25  |                    |              | 6.4.2                        | 6.4.2  | 6.4.2                    |
| Sample collection and processing                                     | collection_device                                   | GENEPIO:0001234                    | The instrument or container used to collect the sample e.g. swab.  | This field includes animal feed. If applicable, select the standardized term and ontology ID for the anatomical material from the picklist provided. Multiple values can be provided, separated by a semi-colon.  | Drag swab [OBI:0002822]   |                    |              | 3.0.0                        | 3.0.0  | 3.0.0                    |
| Sample collection and processing                                     | collection_method                                   | GENEPIO:0001241                    | The process used to collect the sample e.g. phlebotomy, necropsy.  | If applicable, provide the standardized term and ontology ID for the anatomical material from the picklist provided. Multiple values can be provided, separated by a semi-colon.  | Rinsing for specimen collection<br>[GENEPIO_0002116]                            |                    |              | 3.0.0                        | 3.0.0  | 3.0.0                    |
| Sample collection and processing                                     | sample_volume_measurement_value                     | GENEPIO:0100768                    | The numerical value of the volume measurement of the sample collected.   | Provide the numerical value of volume.  | 5   |                    |              | 11.1.1                       | 11.1.1 | 11.1.1                   |
| 2  | sample_volume_measurement_unit                      | GENEPIO:0100769                    | The units of the volume measurement of the sample collected  | Provide the units from the pick list.   | milliliter (mL) [UO:0000098]  |                    |              | 11.1.1                       | 11.1.1 | 11.1.1                   |
| Sample collection and processing<br>Sample collection and processing | residual_sample_status                              | GENEPIO:0101090                    | collected.   |   |   |                    |              | 11.1.1                       | 11.1.1 | 11.1.1                   |
|  |   |                                    | A specification of the way that a  | Provide a description of how the same was   | Frozen at -20C  |                    |              |                              |        |                          |
| Sample collection and processing                                     | sample_storage_method                               | GENEPIO:0100448                    |  | stored.  Provide a description of the material or matrix  |   |                    |              | 3.0.0                        | 3.0.0  | 3.0.0                    |
| Sample collection and processing                                     | sample_storage_medium sample_storage_duration_value | GENEPIO:0100449<br>GENEPIO:0101014 | sample is stored.  The numerical value of the time   | used to store the sample.  Provide the numerical value of time.   | PBS + 20% glycerol 5  |                    |              | 3.0.0                        | 3.0.0  | 3.0.0                    |
| Sample collection and processing                                     |   |                                    | measurement during which a sample is in storage.   |   |   |                    |              | 11.1.1                       | 11.1.1 | 11.1.1                   |
| Sample collection and processing                                     | sample_storage_duration_unit                        | GENEPIO:0101015                    | storage duration.  | Provide the units from the pick list.   | Day [UO:0000033]  |                    |              | 11.1.1                       | 11.1.1 | 11.1.1                   |
| Sample collection and processing                                     | nucleic_acid_storage_duration_value                 | GENEPIO:0101085                    | The numerical value of the time measurement during which the extracted nucleic acid is in storage.   | Provide the numerical value of time.  | 5   |                    |              | 11.1.1                       | 11.1.1 | 11.1.1                   |
| Sample collection and processing                                     | nucleic_acid_storage_duration_unit                  | GENEPIO:0101086                    | The units of a measured extracted nucleic acid storage duration.   | Provide the units from the pick list.   | Year [UO:0000036]   |                    |              | 11.1.1                       | 11.1.1 | 11.1.1                   |
| Sample collection and processing                                     | available_data_types                                | GENEPIO:0100690                    | The type of data that is available, that may or may not require permission to access.  | This field provides information about additional data types that are available that may provide context for interpretation of the sequence data. Provide a term from the picklist for additional data types that are available. Additional data types may require special permission to access. Contact the data provider for more information. | Total coliform count data   |                    |              | 3.0.0                        | 3.0.0  | 3.0.0                    |
| · •  |   |                                    | Detailed information regarding   | provide context for interpreting genomic  | Pooled metagenomes<br>containing extended spectrum<br>beta-lactamase (ESBL)     |                    |              |                              |        |                          |
| Sample collection and processing                                     | available_data_type_details                         | GENEPIO:0101023                    | other available data types.  | sequence data.  | bacteria  |                    |              | 9.0.0                        | 9.0.0  | 9.0.0                    |
| Environmental conditions and<br>measurements                         |   |                                    |  |   |   |                    |              |                              |        |                          |
| Sample collection and processing                                     | water_depth   | GENEPIO:0100440                    | The depth of some water.   | only (without units).   | 5   |                    |              | 3.0.0                        | 3.0.0  | 9.0.0                    |
|  | water_depth_units                                   | GENEPIO:0101025                    | The units of measurement for<br>water depth.   | Provide the units of measurement for which<br>the depth was recorded.   | m   |                    |              |                              |        | 9.0.0                    |

| Parent Class   | Field                                   | Ontology Identifier                | Definition   | Guidance   | Examples   | Deprecated Label | Deprecated ID | Version<br>Tracking<br>Label | ID             | Description/Gui |
|--|---|------------------------------------|--|--|--|------------------|---------------|------------------------------|----------------|-----------------|
|  | Colour Code Legend                      |                                    |  |  |  |                  |               | Label                        | טו             | dance           |
| Sample collection and processing                                   | sediment_depth                          | GENEPIO:0100697                    | The depth of some sediment.  | Provide the numerical depth only of the sediment (without units).  | 2  |                  |               | 7.6.4                        | 7.6.4          | 9.0.0           |
| Sample collection and processing                                   | sediment_depth_units                    | GENEPIO:0101026                    | The units of measurement for sec   | Provide the units of measurement for which the depth was recorded.   | m  |                  |               | 9.0.0                        | 9.0.0          | 9.0.0           |
| Sample collection and processing                                   | air_temperature                         | GENEPIO:0100441                    | The temperature of some air.   | Provide the numerical value for the temperature of the air (without units).  | 25   |                  |               | 3.0.0                        | 3.0.0          | 9.0.0           |
| OIIIti   | -1- 4                                   | OENEDIO 0404007                    |  | Provide the units of measurement for which   | celsius  |                  |               | 0.00                         | 0.00           | 000             |
| Sample collection and processing  Sample collection and processing | air_temperature_units water_temperature | GENEPIO:0101027<br>GENEPIO:0100698 | temperature.  The temperature of some water.   | the temperature was recorded.  Provide the numerical value for the temperature of the water (without units).   | 4  |                  |               | 9.0.0<br>7.6.4               | 9.0.0<br>7.6.4 | 9.0.0           |
| Sample collection and processing                                   | water_terriperature                     | GENEFIO.0100090                    | The units of measurement for   | Provide the units of measurement for which   | celsius  |                  |               | 7.0.4                        | 7.0.4          | 9.0.0           |
| Sample collection and processing                                   | water_temperature_units                 | GENEPIO:0101028                    | water temperature.  The state of the atmosphere at a place and time as regards heat,   | the temperature was recorded.  | Rain [ENVO:01001564]   |                  |               | 9.0.0                        | 9.0.0          | 9.0.0           |
| Sample collection and processing                                   | weather_type                            | GENEPIO:0100442                    | dryness, sunshine, wind, rain, etc.  | Provide the weather conditions at the time of sample collection.   |  |                  |               | 3.0.0                        | 3.0.0          | 3.0.0           |
| lost information   |   | GENEPIO:0001268                    |  |  |  |                  |               |                              |                |                 |
| Host information   | host (common name)                      | GENEPIO:0001386                    | The commonly used name of the host.  | If the sample is directly from a host, either a common or scientific name must be provided (although both can be included, if known). If known, provide the common name.   | Cow [NCBITaxon:9913];<br>Chicken [NCBITaxon:9913],<br>Human [NCBITaxon:9606] |                  |               | 3.0.0                        | 3.0.0          | 3.0.0           |
| Host information   | host (scientific name)                  | GENEPIO:0001387                    | The taxonomic, or scientific name of the host.   | If the sample is directly from a host, either a common or scientific name must be provided (although both can be included, if known). If known, select the scientific name from the picklist provided.   | Bos taurus [NCBITaxon:9913];<br>Homo sapiens<br>[NCBITaxon:9103]             |                  |               | 3.0.0                        | 3.0.0          | 3.0.0           |
| Host information   | host (ecotype)                          | GENEPIO:0100450                    | The biotype resulting from selection in a particular habitat, e.g. the A. thaliana Ecotype Ler.  | Provide the name of the ecotype of the host organism.  | Sea ecotype  |                  |               | 3.0.0                        | 3.0.0          | 3.0.0           |
|  |   |                                    | A breed is a specific group of domestic animals or plants having homogeneous appearance, homogeneous behavior, and other characteristics that distinguish it from other animals or plants of the same species and that were arrived at through selective | Provide the name of the breed of the host  | Holstein   |                  |               |                              |                |                 |
| Host information   | host (breed)                            | GENEPIO:0100451                    | breeding.  | organism.  |  |                  |               | 3.0.0                        | 3.0.0          | 3.0.0           |
|  |   |                                    | The name of the host at a certain stage of food production, which may depend on its age or stage   | Select the host's food production name from  | Calf [FOODON:03411349]   |                  |               |                              |                |                 |
| Host information   | host (food production name)             | GENEPIO:0100452                    | of sexual maturity.  Age of host at the time of  | the pick list.  Select the corresponding host age bin from the   | First summer   |                  |               | 3.0.0                        | 3.0.0          | 3.0.0           |
| Host information   | host_age_bin                            | GENEPIO:0001394                    | sampling, expressed as an age group.   | pick list provided in the template. If not available, provide a null value or leave blank.   | [GENEPIO:0100685]  |                  |               | 7.6.4                        | 7.6.4          | 7.6.4           |
|  |   |                                    | The name of the disease  | This field is only required if the Pathogen.cl package was selected. If the host was sick, provide the name of the disease. The standardized term can be sourced from this look-up service: https://www.ebi.ac.uk/ols/ontologies/doid if the   | mastitis, gastroenteritis  |                  |               |                              |                |                 |
| Host information   | host_disease                            | GENEPIO:0001391                    | experienced by the host.   | disease is not known, put "missing".   |  |                  |               | 3.0.0                        | 3.0.0          | 3.0.0           |
| 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   |                  |               | 3.0.0                        | 3.0.0          | 3.0.0           |
| Strain and isolation information                                   | strain                                  | GENEPIO:0100455                    | The strain identifier.   | A population or type of organisms that is genetically different from others of the same species and possessing a set of defined characteristics. If the isolate represents or is derived from, a lab reference strain or strain from a type culture collection, provide the strain identifier.   | K12  |                  |               | 3.0.0                        | 3.0.0          | 3.0.0           |
|  |   |                                    |  | 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. In 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 asure |  |                  |               |                              |                |                 |
| Strain and isolation information                                   | isolate_ID                              | GENEPIO:0100456                    | Identifier of the specific isolate.  | to include it in the alternative_isolate_ID field.   |  |                  |               | 3.0.0                        | 3.0.0          | 3.0.0           |

|                                  |  |                     |   |  |                                    | Deprecated Label | Deprecated ID | Version           |       |                 |
|----------------------------------|--|---------------------|---|--|------------------------------------|------------------|---------------|-------------------|-------|-----------------|
| Parent Class                     | Field                                    | Ontology Identifier | Definition  | Guidance   | Examples                           |                  |               | Tracking<br>Label | ID    | Description/Gui |
|                                  | Colour Code Legend                       | _                   |   |  |                                    |                  |               | Label             | טו    | dance           |
|                                  |  |                     | An alternative isolate_ID assigned to the isolate by  | 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]. Agency short forms include the following: Public Health Agency of Canadia: PHAC Canadian Food Inspection Agency: CFIA Agriculture and Agri-Food Canada: AFC Fisheries and Oceans Canada: DFO Environment and Climate Change Canada: ECC Health Canada: HC An example of a properly formatted alternative isolate identifier would be e.g. XY24567(CFIA) Multiple alternative isolate identifier would be | GHIF3456[PHAC];<br>QWICK222[CFIA]  |                  |               |                   |       |                 |
| Strain and isolation information | alternative_isolate_ID                   | GENEPIO:0100457     | another organization.   | provided, separated by semi-colons.  | OUD ON 4500                        |                  |               | 3.0.0             | 3.0.0 | 3.0.0           |
|                                  |  |                     | The identifier assigned to a<br>progenitor isolate derived from an<br>isolate that was directly obtained  | If your sequence data pertains to progeny of   | SUB_ON_1526                        |                  |               |                   |       |                 |
| Strain and isolation information | progeny_isolate_ID                       | GENEPIO:0100458     | from a sample.  | progeny_isolate_ID.  |                                    |                  |               | 3.0.0             | 3.0.0 | 3.0.0           |
| Strain and isolation information | IRIDA_isolate_ID                         | GENEPIO:0100459     | The identifier of the isolate in the IRIDA platform.  | Provide the "sample ID" used to track information linked to the isolate in IRIDA. IRIDA sample IDs should be unqiue to avoid ID olash. This is very important in large Projects, especially when samples are shared from different organizations. Download the IRIDA sample ID and add it to the sample data in your spreadsheet as part of good data management practices.  |                                    |                  |               | 3.0.0             | 3.0.0 | 3.0.0           |
| Strain and isolation information | IRIDA_project_ID                         | GENEPIO:0100460     | The identifier of the Project in the iRIDA platform.  | Provide the IRIDA "project ID".  | 666                                |                  |               | 3.0.0             | 3.0.0 | 3.0.0           |
|                                  | IKIDA_project_iD                         |                     | The name of the agency, organization or institution with which the individual who performed the isolation | Provide the INDA project to  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   | Canada (PHAC)<br>[GENEPIO:0100551] |                  |               |                   |       | 3.0.0           |
| Strain and isolation information | isolated_by                              | GENEPIO:0100461     | procedure is affiliated.  | provide a null value.  |                                    |                  |               | 3.0.0             | 3.0.0 | 3.0.0           |
| 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                           |                  |               | 3.0.0             | 3.0.0 | 3.0.0           |
|                                  |  |                     | The name or title of the contact responsible for follow-up  | 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,   |                                    |                  |               |                   |       |                 |
| Strain and isolation information | isolated_by_contact_name                 | GENEPIO:0100463     | regarding the isolate.  | leave blank or provide a null value.  Provide the email associated with the listed   | enterics@lab.ca                    |                  |               | 3.0.0             | 3.0.0 | 3.0.0           |
| Strain and isolation information | isolated_by_contact_email                | GENEPIO:0100464     | The email address of the contact responsible for follow-up regarding the isolate.                         | contact. As personnel furmover 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.  |                                    |                  |               | 3.0.0             | 3.0.0 | 3.0.0           |
| 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".   | 2020-10-30                         |                  |               | 3.0.0             | 3.0.0 | 3.0.0           |
|                                  |  |                     | The date on which the isolate   | Provide the date according to the ISO 8601 standard "YYYY-MM-DD", "YYYY-MM" or   | 2020-11-15                         |                  |               |                   |       |                 |
| Strain and isolation information | isolate_received_date                    | GENEPIO:0100466     | was received by the laboratory.   | "YYYY".  | Salmonella enterica subsp.         |                  |               | 3.0.0             | 3.0.0 | 3.0.0           |
| Strain and isolation information | organism                                 | GENEPIO:0001191     | Taxonomic name of the organism.   | Put the genus and species (and subspecies if applicable) of the bacteria, if known. The standardized term can be sourced from this look-up service: https://www.ebi.ac.uk/ols/ontologies/ncbitaxon.  | enterica [NCBITaxon:59201]         |                  |               | 3.0.0             | 3.0.0 | 3.0.0           |
| Strain and isolation information | taxonomic_identification_process         | GENEPIO:0100583     | The type of planned process by which an organismal entity is associated with a taxon or taxa.             | Provide the type of method used to determine the taxonomic identity of the organism by selecting a value from the pick list. If the information is unknown or cannot be provided, leave blank or provide a null value.   | PCR assay [OBI:0002740]            |                  |               | 4.2.1             | 4.2.1 | 4.2.1           |
| Strain and isolation information | taxonomic_identification_process_details | GENEPIO:0100584     | The details of the process used to determine the taxonomic identification of an organism.                 | Briefly describe the taxonomic identififcation method details using free text.   | Biolog instrument                  |                  |               | 4.2.1             | 4.2.1 | 4.2.1           |

| Parent Class  | Field                        | Ontology Identifier                | Definition   | Guidance  | Examples                            | Deprecated Label De | precated ID | Version<br>Tracking |        |                |
|---|------------------------------|------------------------------------|--|---|-------------------------------------|---------------------|-------------|---------------------|--------|----------------|
|   | Colour Codo Logand           | 0,                                 |  |   |                                     |                     |             | Label               | ID     | Description/Gu |
|   | Colour Code Legend           |                                    |  | Only include this information if it has been  | Heidelberg                          |                     |             |                     |        | dance          |
|   |                              |                                    |  | determined by traditional serological methods   |                                     |                     |             |                     |        |                |
| Strain and isolation information                                  | serovar                      | GENEPIO:0100467                    | The serovar of the organism.   | or a validated in silico prediction tool e.g. SISTR.  |                                     |                     |             | 3.0.0               | 3.0.0  | 3.0.0          |
|   |                              |                                    |  | If the serovar was determined via traditional   | SISTR 1.0.1                         |                     |             |                     |        |                |
|   |                              |                                    |  | serotyping methods, put "Traditional<br>serotyping". If the serovar was determined via            |                                     |                     |             |                     |        |                |
| Strain and isolation information                                  |                              | OENEDIO 0400400                    | The method used to determine   | in silico methods, provide the name and   |                                     |                     |             | 0.00                | 200    | 200            |
| Strain and isolation information Strain and isolation information | serotyping_method phagetype  | GENEPIO:0100468<br>GENEPIO:0100469 | the serovar.  The phagetype of the organism.                           | version number of the software.  Provide if known. If unknown, put "missing".                     | 4'                                  | 7                   |             | 3.0.0               | 3.0.0  | 3.0.0          |
| Sequence information  | phagotype                    | GENEPIO:0001441                    | The phagetype of the organism.   | Torido II Milonii II dindromi, pat miloniig .   |                                     |                     |             | 0.0.0               | 0.0.0  | 0.0.0          |
| ·   |                              |                                    |  | Every "library ID" from a single submitter must   | LS_2010_NP_123446                   |                     |             |                     |        |                |
|   |                              |                                    | The user-specified identifier for                                      | be unique. It can have any format, but we<br>suggest that you make it concise, unique and         |                                     |                     |             |                     |        |                |
| Sequence information  | library_ID                   | GENEPIO:0001448                    | the library prepared for sequencing.                                   | consistent within your lab, and as informative as possible.                                       |                                     |                     |             | 3.0.0               | 3.0.0  | 3.0.0          |
| sequence information  | library_iD                   | GENEPIO.0001446                    | sequencing.  | Provide the name of the agency, organization  | Public Health Agency of             |                     |             | 3.0.0               | 3.0.0  | 3.0.0          |
|   |                              |                                    | The name of the agency,  | or institution that performed the sequencing in   | Canada (PHAC)                       |                     |             |                     |        |                |
|   |                              |                                    | organization or institution responsible for sequencing the             | full (avoid abbreviations). If the information is<br>unknown or cannot be provided, leave blank o |                                     |                     |             |                     |        |                |
| Sequence information  | sequenced_by                 | GENEPIO:0100416                    | isolate's genome.  | provide a null value.   |                                     |                     |             | 3.0.0               | 3.0.0  | 3.0.0          |
|   |                              |                                    |  | Provide the name of the specific laboratory<br>that that performed the sequencing in full         | Topp Lab                            |                     |             |                     |        |                |
|   |                              |                                    |  | (avoid abbreviations). If the information is  |                                     |                     |             |                     |        |                |
| Sequence information  | sequenced_by_laboratory_name | GENEPIO:0100470                    | of the responsible for sequencing<br>the isolate's genome.             | unknown or cannot be provided, leave blank o<br>provide a null value.                             | r                                   |                     |             | 3.0.0               | 3.0.0  | 3.0.0          |
| '   |                              |                                    | Ŭ  | Provide the name of an individual or their job  | Enterics Lab Manager                |                     |             |                     |        |                |
|   |                              |                                    |  | title. As personnel turnover may render the<br>contact's name obsolete, it is more prefereable    | _                                   |                     |             |                     |        |                |
|   |                              |                                    |  | to provide a job title for ensuring accuracy of   |                                     |                     |             |                     |        |                |
|   |                              |                                    | The name or title of the contact<br>responsible for follow-up          | information and institutional memory. If the<br>information is unknown or cannot be provided,     |                                     |                     |             |                     |        |                |
| Sequence information  | sequenced_by_contact_name    | GENEPIO:0100471                    | regarding the sequence.  | leave blank or provide a null value.  |                                     |                     |             | 3.0.0               | 3.0.0  | 3.0.0          |
|   |                              |                                    |  | Provide the email associated with the listed<br>contact. As personnel turnover may render an      | enterics@lab.ca                     |                     |             |                     |        |                |
|   |                              |                                    |  | individual's email obsolete, it is more   |                                     |                     |             |                     |        |                |
|   |                              |                                    |  | prefereable to provide an address for a<br>position or lab, to ensure accuracy of                 |                                     |                     |             |                     |        |                |
|   |                              |                                    | The email address of the contact<br>responsible for follow-up          | information and institutional memory. If the information is unknown or cannot be provided,        |                                     |                     |             |                     |        |                |
| Sequence information  | sequenced_by_contact_email   | GENEPIO:0100422                    | regarding the sequence.  | leave blank or provide a null value.  |                                     |                     |             | 3.0.0               | 3.0.0  | 3.0.0          |
|   |                              |                                    |  | Provide the reason for sequencing by  | Research                            |                     |             |                     |        |                |
|   |                              |                                    |  | selecting a value from the following pick list:<br>Diagnostic testing, Surveillance, Monitoring,  | [GENEPIO:0100003]                   |                     |             |                     |        |                |
|   |                              |                                    |  | Clinical trial, Field experiment, Environmental testing. If the information is unknown or canno   |                                     |                     |             |                     |        |                |
|   |                              |                                    | The reason that the sample was   |   | ·                                   |                     |             |                     |        |                |
| Sequence information  | purpose_of_sequencing        | GENEPIO:0001445                    | sequenced.   | value.  | 111D ODD! (D1 1050)                 |                     |             | 3.0.0               | 3.0.0  | 3.0.0          |
|   |                              |                                    | The name of the<br>project/initiative/program for                      | Provide the name of the project and/or the<br>project ID here. If the information is unknown      | AMR-GRDI (PA-1356)                  |                     |             |                     |        |                |
| Sequence information  | sequencing_project_name      | GENEPIO:0100472                    | which sequencing was<br>performed.                                     | or cannot be provided, leave blank or provide a null value.                                       |                                     |                     |             | 3.0.0               | 3.0.0  | 3.0.0          |
| Sequence information  | sequenting_project_name      | GENEFIO.0100472                    | periornieu.  | Provide the name of the company that created  | Illumina [GENEPIO:0001923]          |                     |             | 3.0.0               | 3.0.0  | 3.0.0          |
|   |                              |                                    |  | the sequencing instrument by selecting a value from the template pick list. If the                |                                     |                     |             |                     |        |                |
|   |                              |                                    | The platform technology used to  | information is unknown or cannot be provided,   |                                     |                     |             |                     |        |                |
| Sequence information  | sequencing_platform          | GENEPIO:0100473                    | perform the sequencing.  | leave blank or provide a null value.  |                                     |                     |             | 3.0.0               | 3.0.0  | 3.0.0          |
|   |                              |                                    |  | Provide the model sequencing instrument by<br>selecting a value from the template pick list. If   | [GENEPIO:0100117]                   |                     |             |                     |        |                |
| Sequence information  | sequencing instrument        | GENEPIO:0001452                    | The model of the sequencing instrument used.                           | the information is unknown or cannot be provided, leave blank or provide a null value.            |                                     |                     |             | 3.0.0               | 3.0.0  | 3.0.0          |
| ocquence inionnation  | sequencing_instrument        | GENEPIO:0001452<br>GENEPIO:0100997 | The overarching sequencing   | Example Guidance: Provide the name of the   | whole genome sequencing             |                     |             | 3.0.0               | 3.0.0  | 3.0.0          |
|   |                              | -                                  | methodology that was used to   | DNA or RNA sequencing technology used in  | assay [OBI:0002117]                 |                     |             |                     |        |                |
| Sequence information  | sequencing_assay_type        |                                    | determine the sequence of a biomaterial.                               | your study. If unsure refer to the protocol documentation, or provide a null value.               |                                     |                     |             | 10.0.0              | 10.0.0 | 10.0.0         |
|   |                              |                                    | The name of the DNA library  | Devide the second of the liberary of the  |                                     |                     |             |                     |        |                |
| Sequence information  | library_preparation_kit      | GENEPIO:0001450                    | preparation kit used to generate the library being sequenced.          | Provide the name of the library preparation kit<br>used.  | Nextera XT                          |                     |             | 3.0.0               | 3.0.0  | 3.0.0          |
|   | _                            | GENEPIO:0100843                    | The length of the DNA fragment   | Provide the fragment length in base pairs (do   | 400                                 | )                   |             |                     |        |                |
|   |                              |                                    | generated by mechanical<br>shearing or enzymatic digestion             | not include the units).   |                                     |                     |             |                     |        |                |
| Sequence information  | DNA fragment length          |                                    | for the purposes of library  |   |                                     |                     |             |                     |        |                |
| Sequence information  | DNA fragment length          | GENEPIO:0100966                    | preparation. The molecular technique used to                           | Provide the name of the enrichment method   | Hybrid selection method             |                     |             |                     |        |                |
|   |                              | 32.120.0100300                     | selectively capture and amplify<br>specific regions of interest from a |   | (bait-capture)<br>[GENEPIO:0001950] |                     |             |                     |        |                |
|   |                              |                                    |  |   |                                     |                     |             |                     |        |                |

| Parent Class                                       | Field  | Ontology Identifier                | Definition  | Guidance  | Examples   | Deprecated Label | Deprecated ID | Version<br>Tracking |       |                |
|--|--|------------------------------------|---|---|--|------------------|---------------|---------------------|-------|----------------|
|  |  | Sitto og jaldentille               |   |   | - Allipioo   |                  |               | Label               | ID    | Description/Gu |
|  | Colour Code Legend   | GENEPIO:0100967                    | Details that provide additional                                   | Provide details that are applicable to the  | enrichment was done using  |                  |               |                     |       | dance          |
|  |  | GENEPIO:0100967                    | context to the molecular  | method you used. Note: If bait-capture  | Twist's respiratory virus  |                  |               |                     |       |                |
|  |  |                                    | technique used to selectively                                     | methods were used for enrichment, provide   | research panel:  |                  |               |                     |       |                |
|  |  |                                    | capture and amplify specific                                      | the panel name and version number (or a URL   |  |                  |               |                     |       |                |
| equence information                                | genomic target enrichment method details   |                                    | regions of interest from a genome.                                | providing that information).  | m/products/ngs/fixed-panels/re<br>spiratory-virus-research-panel |                  |               |                     |       |                |
|  | g  | GENEPIO:0001456                    | 9   | Provide the name and version of the primer  | artic v3   |                  |               |                     |       |                |
|  |  |                                    | (primer sequences, binding  | scheme used to generate the amplicons for   |  |                  |               |                     |       |                |
|  |  |                                    | positions, fragment size  | sequencing.   |  |                  |               |                     |       |                |
| Sequence information                               | amplicon pcr primer scheme   |                                    | generated etc) used to generate<br>the amplicons to be sequenced. |   |  |                  |               |                     |       |                |
|  |  | GENEPIO:0001449                    | The length of the amplicon  | Provide the amplicon size expressed in base   | 300  |                  |               |                     |       |                |
| Sequence information                               | amplicon size  |                                    | generated by PCR amplification.                                   | pairs.  | B 0 4 4  |                  |               |                     |       |                |
|  |  |                                    | The version number of the flow<br>cell used for generating        | Flow cells can vary in terms of design,<br>chemistry, capacity, etc. The version of the         | R.9.4.1  |                  |               |                     |       |                |
|  |  |                                    | sequence data.  | flow cell used to generate sequence data can  |  |                  |               |                     |       |                |
|  |  |                                    |   | affect sequence quantity and quality. Record  |  |                  |               |                     |       |                |
|  |  |                                    |   | the version of the flow cell used to generate sequence data. Do not include "version" or "v"    |  |                  |               |                     |       |                |
| Sequence information                               | sequencing flow cell version   | GENEPIO:0101084                    |   | in the version number.  |  |                  |               |                     |       |                |
|  |  |                                    |   | Provide the name and version of the   | https://www.protocols.io/view/n                                  |                  |               |                     |       |                |
|  |  |                                    | The protocol or method used for                                   | procedure or protocol used for sequencing.<br>You can also provide a link to a protocol         | cov-2019-sequencing-protocol-<br>bbmuik6w?version_warning=n      |                  |               |                     |       |                |
| Sequence information                               | sequencing_protocol  | GENEPIO:0001454                    | sequencing.   | online.   | Q  |                  |               | 3.0.0               | 3.0.0 | 3.0.0          |
|  |  |                                    | The user-specified filename of                                    |   | ABC123_S1_L001_R1_001.fa   |                  |               |                     |       |                |
| Sequence information                               | r1_fastq_filename  | GENEPIO:0001476                    | the r1 FASTQ file.  | Provide the r1 FASTQ filename.  | stq.gz   |                  |               | 3.0.0               | 3.0.0 | 3.0.0          |
| Sequence information                               | r2_fastq_filename  | GENEPIO:0001477                    | The user-specified filename of<br>the r2 FASTQ file.              | Provide the r2 FASTQ filename.  | ABC123_S1_L001_R2_001.fa<br>stq.gz                               |                  |               | 3.0.0               | 3.0.0 | 3.0.0          |
| eequenee internation                               | 12_tastq_monamo  | CENER IO.SSOTTIT                   | The user-specified filename of                                    | Trestad die 12 17te Fig. Illestante.  | batch1a_sequences.fast5  |                  |               | 0.0.0               | 0.0.0 | 0.0.0          |
| Sequence information                               | fast5_filename   | GENEPIO:0001480                    | the FAST5 file.   | Provide the FAST5 filename.   |  |                  |               | 3.0.0               | 3.0.0 | 3.0.0          |
|  |  |                                    | The user-defined filename of the                                  |   | pathogenassembly123.fasta  |                  |               |                     |       |                |
| Sequence information Bioinformatics and QC metrics | assembly_filename  | GENEPIO:0001461<br>GENEPIO:0001457 | FASTA file.   | Provide the FASTA filename.   |  |                  |               | 3.0.0               | 3.0.0 | 3.0.0          |
| Bioinformatics and QC metrics                      | quality control method name  | GENEPIO:0100557                    | The name of the method used to                                    | Providing the name of the method used for   | ncov-tools   |                  |               |                     |       |                |
|  | quality control metrod frame   | GENERIO DI 1000001                 | assess whether a sequence   | quality control is very important for interpreting  |  |                  |               |                     |       |                |
|  |  |                                    | passed a predetermined quality                                    | the rest of the QC information. Method names  |  |                  |               |                     |       |                |
|  |  |                                    | control threshold.  | can be provided as the name of a pipeline or a<br>link to a GitHub repository. Multiple methods | 1  |                  |               |                     |       |                |
|  |  |                                    |   | should be listed and separated by a   |  |                  |               |                     |       |                |
| Bioinformatics and QC metrics                      |  |                                    |   | semi-colon. Do not include QC tags in other   |  |                  |               |                     |       |                |
| Bioinformatics and QC metrics                      | quality control method version   | GENEPIO:0100558                    | The version number of the   | fields if no method name is provided.  Methods updates can make big differences to              | 123  |                  |               |                     |       |                |
|  | quality contact monion version   | 02112110.010000                    | method used to assess whether                                     | their outputs. Provide the version of the   | 1.2.0  |                  |               |                     |       |                |
|  |  |                                    | a sequence passed a   | method used for quality control. The version  |  |                  |               |                     |       |                |
|  |  |                                    | predetermined quality control<br>threshold                        | can be expressed using whatever convention<br>the developer implements (e.g. date, semantic     |  |                  |               |                     |       |                |
|  |  |                                    | an ostioia.   | versioning). If multiple methods were used,   |  |                  |               |                     |       |                |
|  |  |                                    |   | record the version numbers in the same order as the method names. Separate the version          |  |                  |               |                     |       |                |
| Bioinformatics and QC metrics                      |  |                                    |   | numbers using a semi-colon.   |  |                  |               |                     |       |                |
|  | quality control determination  | GENEPIO:0100559                    | The determination of a quality                                    | Select a value from the pick list provided. If a  | sequence failed quality control                                  |                  |               |                     |       |                |
|  |  |                                    | control assessment.   | desired value is missing, submit a new term   |  |                  |               |                     |       |                |
|  |  |                                    |   | request to the PHA4GE QC Tag GitHub<br>issuetracker using the New Term Request                  |  |                  |               |                     |       |                |
| Bioinformatics and QC metrics                      |  |                                    |   | form.   |  |                  |               |                     |       |                |
|  | quality control issues   | GENEPIO:0100560                    | The reason contributing to, or                                    | Select a value from the pick list provided. If a  | low average genome coverage                                      |                  |               |                     |       |                |
|  |  |                                    | causing, a low quality<br>determination in a quality control      | desired value is missing, submit a new term request to the PHA4GE QC Tag GitHub                 |  |                  |               |                     |       |                |
|  |  |                                    | assessment.   | issuetracker using the New Term Request   |  |                  |               |                     |       |                |
| Bioinformatics and QC metrics                      |  | 051500 04005                       |   | form.   | 07 1 100 1 111   |                  |               |                     |       |                |
|  | quality control details  | GENEPIO:0100561                    | The details surrounding a low quality determination in a quality  | Provide notes or details regarding QC results using free text.                                  | CT value of 39. Low viral load.<br>Low DNA concentration after   |                  |               |                     |       |                |
| Bioinformatics and QC metrics                      |  |                                    | control assessment.   |   | amplification.   |                  |               |                     |       |                |
|  | raw sequence data processing method  | GENEPIO:0001458                    | The method used for raw data                                      | Raw data processing can have a significant  | Porechop 0.2.3   |                  |               |                     |       |                |
|  |  |                                    | processing such as removing<br>barcodes, adapter trimming,        | impact on data quality and how it can be used.<br>Provide the names and version numbers of      |  |                  |               |                     |       |                |
|  |  |                                    | filtering etc.  | software used for trimming adaptors, quality  |  |                  |               |                     |       |                |
|  |  |                                    |   | filtering, etc (e.g. Trimmomatic v. 0.38,   |  |                  |               |                     |       |                |
| Bioinformatics and QC metrics                      |  |                                    |   | Porechop v. 0.2.3), or a link to a GitHub protocol.   |  |                  |               |                     |       |                |
|  | dehosting method   | GENEPIO:0001459                    | The method used to remove host                                    | Provide the name and version number of the  | Nanostripper   |                  |               |                     |       |                |
|  | -  |                                    | reads from the pathogen   | software used to remove host reads.   |  |                  |               |                     |       |                |
|  |  |                                    |   |   |  |                  |               |                     |       |                |
| Bioinformatics and QC metrics                      | and the second s | CENEDIO 0100807                    | sequence.   | Dravide the name of the coffusion   | CDA des Conomo Asso  |                  |               |                     |       |                |
| Bioinformatics and QC metrics                      | sequence assembly software name  | GENEPIO:0100825                    | sequence.   | Provide the name of the software used to assemble the sequence.                                 | SPAdes Genome Assembler,<br>Canu, wtdbg2, velvet                 |                  |               |                     |       |                |
|  | sequence assembly software name sequence assembly software version   | GENEPIO:0100825<br>GENEPIO:0100826 | sequence.  The name of the software used to assemble a sequence.  |   | SPAdes Genome Assembler,<br>Canu, wtdbg2, velvet<br>3.15.5       |                  |               |                     |       |                |

| Parent Class  | Field                                 | Ontology Identifier                | Definition  | Guidance   | Examples  | Deprecated Label | Deprecated ID | Version<br>Tracking |    |                |
|---|---------------------------------------|------------------------------------|---|--|---|------------------|---------------|---------------------|----|----------------|
| - arone-olass   |                                       | Ontology Identifier                | Definition  | - ourdance   | Examples  |                  |               | Label               | ID | Description/Gu |
|   | Colour Code Legend                    |                                    |   |  |   |                  |               |                     |    | dance          |
| Bioinformatics and QC metrics                               | consensus sequence software name      | GENEPIO:0001463                    | The name of the software used to<br>generate the consensus<br>sequence.                       | Provide the name of the software used to generate the consensus sequence.  | iVar  |                  |               |                     |    |                |
| Bioinformatics and QC metrics                               | consensus sequence software version   | GENEPIO:0001469                    |   | Provide the version of the software used to generate the consensus sequence.   | 1.3   |                  |               |                     |    |                |
|   | breadth of coverage value             | GENEPIO:0001472                    | The percentage of the reference genome covered by the sequenced data, to a prescribed         | Provide value as a percent.  | 95  |                  |               |                     |    |                |
| Bioinformatics and QC metrics                               | depth of coverage value               | GENEPIO:0001474                    | representing a given nucleotide   | Provide value as a fold of coverage.   | 400   |                  |               |                     |    |                |
| Bioinformatics and QC metrics Bioinformatics and QC metrics | depth of coverage threshold           | GENEPIO:0001475                    |   | Provide the threshold fold coverage.   | 100   |                  |               |                     |    |                |
| Bioinformatics and QC metrics                               | genome completeness                   | GENEPIO:0100844                    |   | Provide the genome completeness as a percent (no need to include units).   | 85  |                  |               |                     |    |                |
| Bioinformatics and QC metrics                               | number of base pairs sequenced        | GENEPIO:0001482                    | The number of total base pairs  | Provide a numerical value (no need to include units).  | 387566  |                  |               |                     |    |                |
| Bioinformatics and QC metrics                               | number of total reads                 | GENEPIO:0100827                    | The total number of non-unique  | Provide a numerical value (no need to include units).  | 423867  |                  |               |                     |    |                |
| Bioinformatics and QC metrics                               | number of unique reads                | GENEPIO:0100828                    | The number of unique reads  | Provide a numerical value (no need to include units).  | 248236  |                  |               |                     |    |                |
| 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).  | 150   |                  |               |                     |    |                |
| Bioinformatics and QC metrics                               | number of contigs                     | GENEPIO:0100937                    | The number of contigs (contiguous sequences) in a sequence assembly.                          | Provide a numerical value.   | 10  |                  |               |                     |    |                |
| Bioinformatics and QC metrics                               | percent Ns across total genome length | GENEPIO:0100830                    |   | Provide a numerical value (no need to include units).  | 2   |                  |               |                     |    |                |
| Bioinformatics and QC metrics                               | Ns per 100 kbp                        | GENEPIO:0001484                    |   | Provide a numerical value (no need to include units).  | 342   |                  |               |                     |    |                |
| Bioinformatics and OC metrics                               | N50                                   | GENEPIO:0100938                    | that, together with other reads,<br>represents at least 50% of the<br>nucleotides in a set of | Provide the N50 value in Mb.   | 150   |                  |               |                     |    |                |
|   | percent read contamination            | GENEPIO:0100845                    | of reads identified as<br>contamination (not belonging to<br>the target organism) in a        | Provide the percent contamination value (no need to include units).  | 2   |                  |               |                     |    |                |
| Bioinformatics and QC metrics                               | sequence assembly length              |                                    |   | Provide a numerical value (no need to include units).  | 34272   |                  |               |                     |    |                |
| Bioinformatics and QC metrics                               | consensus genome length               | GENEPIO:0100846<br>GENEPIO:0001483 | reference-based mapping. The length of the genome   | Provide a numerical value (no need to include units).  | 38677   |                  |               |                     |    |                |
| Bioinformatics and QC metrics                               | reference genome accession            | GENEPIO:0001485                    |   | Provide the accession number of the reference  | NC_045512.2                                       |                  |               |                     |    |                |
| Bioinformatics and QC metrics                               | deduplication method                  | GENEPIO:0100831                    | The method used to remove   | genome.  Provide the deduplication software name followed by the version, or a link to a tool or   | DeDup 0.12.8                                      |                  |               |                     |    |                |
| Bioinformatics and QC metrics                               |                                       |                                    | read dataset.   | method.  |   |                  |               |                     |    |                |
| Bioinformatics and QC metrics                               | bioinformatics protocol               | GENEPIO:0001489                    | bioinformatics strategy used.   | Further details regarding the methods used to process raw data, and/or generate assemblies, and/or generate consensus sequences can. This information can be provided in an SOP or protocol or pipeline/workflow. Provide the name and version number of the protocol, or a GitHub link to a pipeline or workflow. | https://github.com/phac-nml/nc<br>ov2019-artic-nf |                  |               |                     |    |                |
| Taxonomic identification                                    |                                       | CENEDIO 0404000                    |   |  |   |                  |               |                     |    |                |
| information   | read mapping software name            | GENEPIO:0101082<br>GENEPIO:0100832 |   | Provide the name of the read mapping software.   | Bowtie2, BWA-MEM, TopHat                          |                  |               |                     |    |                |

| Parent Class                         | Field                                | Ontology Identifier              | Definition  | Guidance  | Examples   | Deprecated Label | Deprecated ID | Version<br>Tracking |             |                |
|--------------------------------------|--------------------------------------|----------------------------------|---|---|--|------------------|---------------|---------------------|-------------|----------------|
| aront oldso                          |                                      | Ontology Identifier              | Johnson   | Caracinoc   |  |                  |               | Label               | ID          | Description/Gu |
|                                      | Colour Code Legend                   |                                  |   |   |  |                  |               |                     |             | dance          |
| axonomic identification information  | read mapping software version        | GENEPIO:0100833                  | The version of the software used<br>to map sequence reads to a<br>reference genome or set of<br>reference genes.  | Provide the version number of the read mapping software.  | 2.5.1  |                  |               |                     |             |                |
|                                      | taxonomic reference database name    | GENEPIO:0100834                  | The name of the taxonomic reference database used to  | Provide the name of the taxonomic reference database.   | NCBITaxon  |                  |               |                     |             |                |
| axonomic identification information  | taxonomic reference database version | GENEPIO:0100835                  | identify the organism.  The version of the taxonomic  | Provide the version number of the taxonomic   | 1.3  |                  |               |                     |             |                |
| Taxonomic identification information |                                      | GENEPIO:0100835                  | reference database used to identify the organism.   | reference database.   |  |                  |               |                     |             |                |
| axonomic identification information  | taxonomic analysis report filename   | GENEPIO:0101074                  | 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.do  |                  |               |                     |             |                |
| Faxonomic identification information | taxonomic analysis date              | GENEPIO:0101075                  | 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 in ISO 8601 format, i.e. "YYYY-MM-DD".   | 2024-02-01   |                  |               |                     |             |                |
|                                      | read mapping criteria                | GENEPIO:0100836                  | A description of the criteria used to map reads to a reference  | Provide a description of the read mapping criteria.   | Phred score >20  |                  |               |                     |             |                |
| axonomic identification information  |                                      | GENEPIO:0100477                  | sequence.   |   |  |                  |               |                     |             |                |
| Public repository information        |                                      | GENEPIO:0100477                  | The name of the agency that   | The name of the agency should be written out in full, (with minor exceptions) and be consistent across multiple submissions. If submitting specimens rather than sequencing   | Environment and Climate<br>Change Canada (EC)<br>[GENEPIO:0100555] |                  |               |                     |             |                |
|                                      |                                      |                                  | submitted the sequence to a   | data, please put the "National Microbiology   |  |                  |               |                     |             |                |
| Public repository information        | sequence_submitted_by                | GENEPIO:0001159                  | database.   | Laboratory (NML)".  Provide the name of an individual or their job  |  |                  |               | 3.0.0               | 3.0.0       | 3.0.0          |
| Public repository information        | sequence_submitted_by_contact_name   | GENEPIO:0100474                  | The name or title of the contact responsible for follow-up regarding the submission of the sequence to a repository or database.  | 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   |                  |               | 3.0.0               | 3.0.0       | 3.0.0          |
| rubiic repository information        | sequence_submitted_by_contact_name   | GENEPIO.0100474                  |   | The email address can represent a specific  | RespLab@lab.ca   |                  |               | 3.0.0               | 3.0.0       | 3.0.0          |
|                                      |                                      |                                  | responsible for submission of the   | individual or lab e.g. johnnyblogs@lab.ca, or   | 1100p2ab@iab.sa  |                  |               |                     |             |                |
| Public repository information        | sequence_submitted_by_contact_email  | GENEPIO:0001165                  | sequence.   | RespLab@lab.ca  If the isolate is associated with a published   | PMID: 33205991   |                  |               | 3.0.0               | 3.0.0       | 3.0.0          |
|                                      |                                      |                                  |   | work which can provide additional information,<br>provide the PubMed identifier of the<br>publication. Other types of identifiers (e.g.   | PMID: 33205991   |                  |               |                     |             |                |
| Public repository information        | publication_ID                       | GENEPIO:0100475                  | The identifier for a publication.   | DOI) are also acceptable.   |  |                  |               | 3.0.0               | 3.0.0       | 3.0.0          |
|                                      |                                      |                                  | The attribute package used to structure metadata in an INSDC  | If the sample is from a specific human or<br>animal, put "Pathogen.cl". If the sample is from<br>an environmental sample including food, feed,<br>production facility, farm, water source, manure   |  |                  |               |                     |             |                |
| Public repository information        | attribute_package                    | GENEPIO:0100476                  | BioSample.  | etc, put "Pathogen.env".  |  |                  |               | 3.0.0               | 3.0.0       | 3.0.0          |
| Public repository information        | bioproject accession                 | GENEPIO:0001136                  | The INSDC accession number of the BioProject(s) to which the BioSample belongs.   | Required if submission is linked to a BioProject. BioProjects are an organizing tool that links together raw sequence data, assemblies, and their associated metadata. A valid BioProject accession has prefix PRJN, PRJE or PRJD, e.g., PRJNA12345 and is created once at the beginning of a new sequencing project. Your laboratory can have one or many BioProjects. | PRJNA12345   |                  |               | 3.0.0               | 3.0.0       | 3.0.0          |
| · ·                                  |                                      |                                  | The identifier assigned to a  | Store the accession returned from the BioSample submission. NCBI BioSamples will have the prefix SAMN, whileEMBL- EBI   | SAMN14180202   |                  |               |                     |             |                |
| Public repository information        | biosample_accession                  | GENEPIO:0001139  GENEPIO:0001142 | BioSample in INSDC archives. The Sequence Read Archive (SRA), European Nucleotide Archive (ENA) or DDBJ Sequence Read Archive (DRA) identifier linking raw read data, methodological metadata and quality control metrics submitted to the INSDC. | BioSamples will have the prefix SAMEA.  Store the accession assigned to the submitted "run". NCBI-SRA accessions start with SRR, EBI-ENA runs start with ERR and DRA accessions start with DRA  | SRR11177792  |                  |               | 3.0.0               | 3.0.0       | 3.0.0          |
| Public repository information        | GenBank_accession                    | GENEPIO:0001142                  | The GenBank/ENA/DDBJ identifier assigned to the   | Store the accession returned from a GenBank/ENA/DDBJ submission.  | MN908947.3   |                  |               | 3.0.0               | 3.0.0       | 3.0.0          |
| Risk assessment information          | 1                                    | GENEPIO:0100478                  | ,   |   |  |                  |               | ,,,,,,,             | , , , , , , |                |

| Parent Class                | Field  | Ontology Identifier | Definition   | Guidance   | Examples   | Deprecated Label | Deprecated ID | Version<br>Tracking | ID    | Description             |
|-----------------------------|--|---------------------|--|--|--|------------------|---------------|---------------------|-------|-------------------------|
|                             | Colour Code Legend   |                     |  |  |  |                  |               | Label               | טו    | Description/Gu<br>dance |
|                             |  |                     |  | Risk assessment requires detailed information<br>regarding the quantities of a pathogen in a<br>specified location, commodity, or environment.<br>As such, it is useful for risk assessors to know<br>what types of information are available through<br>documented methods and results. Provide the<br>metric types that are available in the<br>surveillance project sample plan by selecting<br>them from the roick list. The metrics of interest | collected, Number of positive samples                          |                  |               |                     |       |                         |
| isk assessment information  | prevalence metrics   | GENEPIO:0100480     | Metrics regarding the prevalence of the pathogen of interest obtained from a surveillance project.                             | are "Number of total samples collected", "Number of positive samples", "Average count of hazard organism", "Average count of indicator organism". You do not need to provide the actual values, just indicate that the information is available.   |  |                  |               | 3.0.0               | 3.0.0 | 3.0.0                   |
| isk assessment information  | prevalence_metrics_details                                 | GENEPIO:0100481     | The details pertaining to the prevalence metrics from a surveillance project.  | If there are details pertaining to samples or organism counts in the sample plan that might be informative, provide details using free text.   |  |                  |               | 3.0.0               | 3.0.0 | 3.0.0                   |
| tisk assessment information | stage_of_production  | GENEPIO:0100482     | The stage of food production.  | Provide the stage of food production as free text.   | Abattoir   |                  |               | 3.0.0               | 3.0.0 | 3.0.0                   |
| Risk assessment information |  | GENEPIO:0100483     | The category of the experimental intervention applied in the food  | In some surveys, a particular intervention in the food supply chain in studied. If there was an intervention specified in the sample plan, select the intervention category from the pick  | Vaccination [NCIT:C15346]                                      |                  |               |                     |       | 3.0.0                   |
| Risk assessment information | experimental_intervention  experiment_intervention_details | GENEPIO:0100483     | production system.  The details of the experimental intervention applied in the food production system.                        | list provided.  If an experimental intervention was applied in the survey, provide details in this field as free text.   | 2% cranberry solution mixed in feed                            |                  |               | 3.0.0               | 3.0.0 | 3.0.0                   |
| Antimicrobial resistance    | experiment_intervention_details                            | GENEPIO:0100484     | production system.   | text.  |  |                  |               | 3.0.0               | 3.0.0 | 3.0.0                   |
| Antimicrobial resistance    | antimicrobial_agent_name                                   | GENEPIO:0100521     | The name of the agent that kills or slows the growth of microorganisms, including bacteria, viruses, fungi and protozoans.     | The names of the drug have already been matched with measurement, breakpoint, and phenotype fields in the template. No need to add these unless the drug of interest is not present. Use the Term Request System to request the addition of other agents.  | Amoxicillin-clavulanic<br>[ARO:3003997]                        |                  |               | 3.0.0               | 3.0.0 | 3.0.0                   |
| Antimicrobial resistance    | AMR_testing_by   | GENEPIO:0100511     | The name of the organization that performed the antimicrobial resistance testing.  | Provide the name of the agency, organization or institution that performed the AMR testing, in full (avoid abbreviations). If the information is unknown or cannot be provided, leave blank or provide a null value.   | Canadian Food Inspection<br>Agency (CFIA)<br>[GENEPIO:0100552] |                  |               | 3.0.0               | 3.0.0 | 3.0.0                   |
| Antimicrobial resistance    | AMR_testing_by_laboratory_name                             | GENEPIO:0100512     | The name of the lab within the organization that performed the antimicrobial resistance testing.                               | Provide the name of the specific laboratory that performed the AMR testing (avoid abbreviations). If the information is unknown or cannot be provided, leave blank or provide a null value.  | Topp Lab   |                  |               | 3.0.0               | 3.0.0 | 3.0.0                   |
|                             |  | GENEPIO:0100513     | The name of the individual or the individual's role in the organization that performed the                                     | Provide the name of an individual or their job title. As personnel tumover 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,  | Enterics Lab Manager   |                  |               | 3.0.0               | 3.0.0 | 3.0.0                   |
| Antimicrobial resistance    | AMR_testing_by_contact_name                                | GENEFIC.UIUS13      | antimicrobial resistance testing.  The email of the individual or the individual's role in the organization that performed the | leave blank or provide a null value.  Provide the email associated with the listed contact. As personnel turnover may render an individual's email obsolete, it is more preferable 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  |  |                  |               | 3.0.0               | 3.0.0 | 3.0.0                   |
| Antimicrobial resistance    | AMR_testing_by_contact_email                               | GENEPIO:0100514     | antimicrobial resistance testing.  The date the antimicrobial resistance testing was   | provide a null value.  Provide the date according to the ISO 8601 standard "YYYY-MM-DD", "YYYY-MM" or  | 2022-04-03   | 3                |               | 3.0.0               | 3.0.0 | 3.0.0                   |
| ntimicrobial resistance     | AMR_testing_date   | GENEPIO:0100515     | performed.  The antimicrobial resistance phenotype, as determined by the antibiotic susceptibility                             | "YYYY".  | Susceptible antimicrobial phenotype [ARO:3004302]              |                  |               | 3.0.0               | 3.0.0 | 3.0.0                   |
| Antimicrobial resistance    | antimicrobial_resistance_phenotype                         |                     | measurement and testing standard for this antibiotic   | Select a phenotype from the pick list provided.  |  |                  |               | 7.7.5               |       | 3.0.0                   |
| Antimicrobial resistance    | antimicrobial_measurement                                  |                     | The measured value of antimicrobial resistance.  | This field should only contain a number (either an integer or a number with decimals).   |  | •                |               | 7.7.5               |       | 3.0.0                   |
| Antimicrobial resistance    | antimicrobial_measurement_units                            |                     | The units of the antimicrobial resistance measurement.   | Select the units from the pick list provided.<br>Use the Term Request System to request the<br>addition of other units if necessary.   | ug/mL [UO:0000274]   |                  |               | 7.7.5               |       | 3.0.0                   |
| Antimicrobial resistance    | antimicrobial measurement sign                             |                     | The qualifier associated with the antibiotic susceptibility measurement  | Select the comparator sign from the pick list provided. Use the Term Request System to request the addition of other signs if necessary.   | greater than (>)<br>[GENEPIO:0001006]                          |                  |               | 7.7.5               |       | 3.0.0                   |

| Parent Class             | Field  | Ontology Identifier | Definition   | Guidance  | Examples   | Deprecated Label | Deprecated ID | Version<br>Tracking<br>Label | ID | Description/Gui |
|--------------------------|--|---------------------|--|---|--|------------------|---------------|------------------------------|----|-----------------|
|                          | Colour Code Legend                               |                     |  |   |  |                  |               | Label                        | טו | dance           |
| Antimicrobial resistance | antimicrobial laboratory typing method           |                     | The general method used for antibiotic susceptibility testing.   | Select a typing method from the pick list<br>provided. Use the Term Request System to<br>request the addition of other methods if<br>necessary. | Broth dilution [ARO:3004397]   |                  |               | 7.7.5                        |    | 3.0.0           |
| Antimicrobial resistance | antimicrobial laboratory typing platform         |                     | The brand/platform used for antibiotic susceptibility testing  | Select a typing platform from the pick list provided. Use the Term Request System to request the addition of other platforms if necessary.      | Sensitire [ARO:3004402]  |                  |               | 7.7.5                        |    | 3.0.0           |
| Antimicrobial resistance | antimicrobial_laboratory_typing_platform_version |                     | The specific name and version of<br>the plate, panel, or other platform<br>used for antibiotic susceptibility<br>testing.  | Include any additional information about the<br>antimicrobial susceptibility test such as the<br>drug panel details.                            | CMV3AGNF   |                  |               | 7.7.5                        |    | 3.0.0           |
| Antimicrobial resistance | antimicrobial_vendor_name                        |                     | The name of the vendor of the testing platform used.   | Provide the full name of the company (avoid abbreviations).   | Sensititre [ARO:3004402]   |                  |               | 7.7.5                        |    | 3.0.0           |
| Antimicrobial resistance | antimicrobial_testing_standard                   |                     | The testing standard used for determination of resistance phenotype  | Select a testing standard from the pick list provided.  | Clinical Laboratory and<br>Standards Institute (CLSI)<br>[ARO:3004366] |                  |               | 7.7.5                        |    | 3.0.0           |
| Antimicrobial resistance | antimicrobial testing standard version           |                     | The version number associated with the testing standard used for determination of resistance phenotype   | If applicable, include a version number for the testing standard used.  | M100   |                  |               | 7.7.5                        |    | 3.0.0           |
| Antimicrobial resistance | antimicrobial testing standard details           |                     | The additional details associated  | This information may include the year or location where the testing standard was published. If not applicable, leave blank.                     | 27th ed. Wayne, PA: Clinical and Laboratory Standards Institute; 2017. |                  |               | 77.5                         |    | 3.0.0           |
| Antimicrobial resistance | antimicrobial susceptible breakpoint             |                     | The maximum measurement, in the units specified in the "AMR_measurement_units" field, for a sample to be considered "sensitive" to this antibiotic                                       | This field should only contain a number (either an integer or a number with decimals), since the "<=" qualifier is implied.                     | 8  |                  |               | 7.7.5                        |    | 3.0.0           |
| Antimicrobial resistance | antimicrobial_intermediate_breakpoint            |                     | The intermediate measurement(s), in the units specified in the "AMR measurement_units" field, where a sample would be considered to have an "intermediate" phenotype for this antibiotic | ,   | 16   |                  |               | 7.7.5                        |    | 3.0.0           |
| Antimicrobial resistance | antimicrobial_resistant_breakpoint               |                     | The minimum measurement, in the units specified in the "AMR_measurement_units" field, for a sample to be considered "resistant" to this antibiotic                                       | This field should only contain a number (either an integer or a number with decimals), since the ">=" qualifier is implied.                     | 32   |                  |               | 7.7.5                        |    | 3.0.0           |