Parent Class	Field	Ontology Identifier	Definition	Guidance	Examples	Deprecated Label Deprecated ID	Version Tracking		
	Colour Code Legend					deprecated, always with replacement version	Label	ID	Description/Gui dance
	field name in yellow = required					provided. If a term changes in its meaning, a new term will be created.			
	field name in purple = recommended field name in white = optional								
	Database Identifiers	GENEPIO:0001122							
		January Communication of the C		Store the collector sample ID. If this number is considered identifiable information, provide an alternative ID. Be sure to store the key that maps between the original and alternative IDs for traceability and follow up if necessary. Every collector sample ID from a single submitter must be unique. It can have any format, but we	5				
Database identifiers	specimen_collector_sample_ID	GENEPIO:0001123	The user-defined name for the sample.	suggest that you make it concise, unique and consistent within your lab.	ASDFG123		1.0.0	1.0.0	1.0.0
Database identifiers	specimen_collector_subsample_ID	GENEPIO:0100752	The user-defined identifier assigned to a portion of the original sample.	Store the ID for the subsample/aliquot.	ASDFG123 12	,	1.0.0	1.0.0	1.0.0
Database Identifiers	opeoinon_concoto_custampleic	SENEI ISISTISTISE	The dear defined advisiner designed to a portion or the original earlipse.	If the sample being analyzed is the result of	7.001 0120_12		1.0.0	1.0.0	1.0.0
			The user-defined identifier assigned to a combined (pooled) set of	pooling individual samples, rename the pooled sample with a new identifier. Store the pooled					
Database identifiers	pooled_sample_ID	GENEPIO:0100996	samples.	sample ID.  Store the ID for the site from which a sample	12345AYZ		1.0.0	1.0.0	1.0.0
			The user-defined identifier assigned to a specific location from which	was taken. The "site" is user defined (e.g. it may be a building and its environs, a specific entity within an environment). Please use the same site ID for all samples from a given site, regardless of when these samples were taken. Any important changes in site location, should					
Database identifiers	sampling_site_ID	GENEPIO:0100760	samples are taken.	be represented with a new site ID.  Store the ID for the event during which a sample	Site 12A		1.0.0	1.0.0	1.0.0
Database identifiers	sampling_event_ID	GENEPIO:0100761	The user-defined identifier assigned to a specific event during which one or more samples are taken, from one or more sites.	or samples were taken. For example, an event could be one person taking samples from multiple sites, or multiple people taking samples from one site.	Event 120522.	1	1.0.0	1.0.0	1.0.0
Database identifiers	BioProject_accession	GENEPIO:0001136	The INSDC (i.e., ENA, NCBI, or DDBJ) accession number of the BioProject(s) to which the BioSample belongs.	Store the BioProject accession number BioProjects are an organizing tool that links together raw sequence data, assemblies, and their associated metadata. Each province will be assigned a different bioproject accession number by the National Microbiology Lab. A valid NCBI BioProject accession has prefix PRJN e.g., PRJNA12345, and is created once at the beginning of a new sequencing project.			1.0.0	1.0.0	1.0.0
Database identifiers	BioSample_accession	GENEPIO:0001139	The identifier assigned to a BioSample in INSDC (i.e., ENA, NCBI, or DDBJ) archives.	Store the accession returned from the BioSample submission. NCBI BioSamples will have the prefix SAMN, ENA have the prefix SAMEA. DDBJ have SAMD	SAMN1418020 2, SAMD00000000		1.0.0	1.0.0	1.0.0
Database Identifiers	Diocumpio_diocession	SENEI ISISSO IIIS	DDDO) drawroc.	o uner, esse have orang			1.0.0	1.0.0	1.0.0
			The versioned identifier assigned to an assembly or consensus	Store the versioned GenBank accession					
Database identifiers	GenBank_accession_(versioned)	GENEPIO:0100754	sequence in GenBank archives.	assigned to the submitted sequence.	LZ986655.1		1.0.0	1.0.0	1.0.0
Database identifiers	SRA_accession	GENEPIO:0001142	The Sequence Read Archive (SRA) identifier linking raw read data, methodological metadata and quality control metrics submitted to the INSDC.	Store the accession assigned to the submitted sequence. NCBI-SRA accessions start with SRR.	SRR11177792		1.0.0	1.0.0	1.0.0
Database identifiers	ENA_accession	GENEPIO:0100755	The identifier assigned to a sequence in the European Nucleotide Archive (ENA).	Store the accession assigned to the submitted sequence. ENA sequence accessions start with ERR.	ERR123456		1.0.0	1.0.0	1.0.0
Database identifiers	DRA_accession	GENEPIO:0100757	The identifier assigned to a sequence in DNA Data Bank of Japan (DDBJ) sequence read archives.	Store the accession assigned to the submitted sequence. DRA accessions start with DRR.	DRR123456		1.0.0	1.0.0	1.0.0
	Sample collection and processing	GENEPIO:0001150							
			The name of the individual responsible for the data governance,	Provide the name of the sample collection data					
Sample collection and processing	sample_collection_data_steward_name	GENEPIO:0100762	(meta)data usage and distribution of the sample.	steward.  Provide the email address of the sample	Joe Bloggs		1.0.0	1.0.0	1.0.0
			The email address of the individual responsible for the data	collection data steward. This may or may not be the same individual/organization that collected the sample. If the contact is the same, provide the same address as the "sample collector	bloggsj@aglab				
Sample collection and processing	sample_collection_data_steward_contact_email	GENEPIO:0101107	governance, (meta)data usage and distribution of the sample.	contact email".	ca		1.0.0	1.0.0	1.0.0

Parent Class	Field	Ontology Identifier	Definition	Guidance	Examples	Deprecated Label Deprecated ID	Version Tracking		
Sample collection and processing	sample_collected_by	GENEPIO:0001153	The name of the organization with which the sample collector is affiliated.	The name of the agency should be written out in full, (with minor exceptions) and be consistent across multiple submissions.			1.0.0	1.0.0	1.0.0
Sample collection and processing	sample_collector_contact_email	GENEPIO:0001156	The email address of the contact responsible for follow-up regarding the sample.	The email address can represent a specific individual or lab e.g. johnnyblogs@lab.ca, or Respl.ab@lab.ca	WaterTester@f acility.ca		1.0.0	1.0.0	1.0.0
Sample collection and processing	geo_loc_name_(country)	GENEPIO:0001181	The country of origin of the sample.	If known, select a value from the pick list.  Provide the state/province/territory name from the GAZ geography ontology. Search for	Canada		1.0.0	1.0.0	1.0.0
Sample collection and processing	geo_loc_name_(state/province/territory)	GENEPIO:0001185	The state/province/territory of origin of the sample.	geography terms here: https://www.ebi.ac.uk/ols/ontologies/ga	Western Cape		1.0.0	1.0.0	1.0.0
Sample collection and processing	geo_loc_name_(county/region)	GENEPIO:0100280	The county/region of origin of the sample.	Provide the county/region name from the GAZ geography ontology. Search for geography terms here: https://www.ebi.ac.uk/ols/ontologies/gaz	Derbyshire		1.0.0	1.0.0	1.0.0
Sample collection and processing	geo_loc_name_(city)	GENEPIO:0001189	The city of origin of the sample.	Provide the city name from the GAZ geography ontology. Search for geography terms here: https://www.ebi.ac.uk/ols/ontologies/gaz	Vancouver		1.0.0	1.0.0	1.0.0
Sample collection and processing	geo_loc_name_(site)	GENEPIO:0100436	The name of a specific geographical location e.g. Credit River (rather than river).	Provide the name of the specific geographical site using a specific noun (a word that names a certain place, thing).	Credit River		1.0.0	1.0.0	1.0.0
Sample collection and processing	geo loc latitude	GENEPIO:0100309	The latitude coordinates of the geographical location of sample collection.	Provide latitude coordinates if available. Do not use the centre of the city/region/province/state/country or the location of your agency as a proxy, as this implicates a real location and is misleading. Specify as degrees latitude in format "fd[d.dddd] NJS".	38.98 N		1.0.0	100	100
Sample collection and processing	geo_io_autule	GENEFIO.0100309	The longitude coordinates of the geographical location of sample	Provide longitude coordinates if available. Do not use the centre of the city/region/province/state/country or the location of your agency as a proxy, as this implicates a real location and is misleading. Specify as			1.0.0	1.0.0	1.0.0
Sample collection and processing	geo_loc_longitude	GENEPIO:0100310	collection.	Provide the official nomenclature for the organism(s) present in the sample. Multiple organisms can be entered, separated by semicolons. Avoid abbreviations. Search for taxonomic names here:	77.11 W		1.0.0	1.0.0	1.0.0
Sample collection and processing	organism	GENEPIO:0001191	Taxonomic name of the organism.	ncbi.nlm.nih.gov/taxonomy.	Vibrio cholerae		1.0.0	1.0.0	1.0.0
Sample collection and processing	influenza_subtype	GENEPIO:0101108					1.0.0	1.0.0	1.0.0
Sample collection and processing	influenza_subtyping_scheme_name	GENEPIO:0101109					1.0.0	1.0.0	1.0.0
Sample collection and processing	taxonomic_identification_process	GENEPIO:0100583					1.0.0	1.0.0	1.0.0
Sample collection and processing	virus_identifier	GENEPIO:0101110					1.0.0	1.0.0	1.0.0
Sample collection and processing	WHO/OIE/FAO_H5_clade	GENEPIO:0101111		If your cample is a continuous cample places			1.0.0	1.0.0	1.0.0
			The date on which the sample was collected, or sampling began for a	If your sample is a continuous sample please use this field to capture your start date. Sample collection date is critical for surveillance and many types of analyses. Required granularity includes year, month and day. The date should be provided in ISO 8601 standard format					
Sample collection and processing	sample_collection_date	GENEPIO:0001174	continuous sample.	"YYYY-MM-DD".  Provide the sample received date in ISO 8601	2020-03-16		1.0.0	1.0.0	1.0.0
Sample collection and processing	sample received date	GENEPIO:0001179	The date on which the sample was received.	format, i.e. "YYYY-MM-DD".  Provide the sample processed date in ISO 8601	2020-03-28		1.0.0	1.0.0	1.0.0
Sample collection and processing	sample processing date	GENEPIO:0100763	The date on which the sample was processed.	format, i.e. "YYYY-MM-DD". The sample may be collected and processed (e.g. filtered, extraction) on the same day, or on different dates.			1.0.0	1.0.0	1.0.0

Parent Class	Field	Ontology Identifier	Definition	Guidance	Examples	Deprecated Label	Deprecated ID	Version Tracking		
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 manufactured 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.	·			1.0.0	1.0.0	100
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 [FOODON:000 03927]; Bone meal [ENVO:020000 54]; Chicken breast [FOODON:000 02703]			1.0.0	1.0.0	1.0.0
			Any characteristic of the food product pertaining to its state, processing,	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 ontainment (e.g. canned), and any information about a label claim	Food (chopped) [FOODON:000 02777]; Ready-to-eat (RTE) I [FOODON:033					
Sample collection and processing	food_product_properties	GENEPIO:0100445	a label claim, or implications for consumers.		16636] Plastic tray or pan [FOODON:034			1.0.0	1.0.0	1.0.0
Sample collection and processing  Sample collection and processing	food_packaging  food_quality_date	GENEPIO:0100447  GENEPIO:0100615	The type of packaging used to contain a food product.  A date recommended for the use of a product while at peak quality, this date is not a reflection of safety unless used on infant formula.	food product was packaged.  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.	90126]			1.0.0	1.0.0	1.0.0
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.				1.0.0	1.0.0	1.0.0
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.	Poultry hatchery [ENVO:010018 74]			1.0.0	1.0.0	1.0.0
Sample collection and processing	environmental_material	GENEPIO:0001223	A substance obtained from the natural or man-made environment e.g. soil, water, sewage, door handle, bed handrall, face mask.	If applicable, select the standardized term and ontology ID for the environmental material from the picklist provided. Separated by a semi-colon.	Soil [ENVO:000019 98]; Water [CHEBI:15377]; Wastewater [ENVO:000020 01]; Broom [ENVO:035013 77]			1.0.0	1.0.0	1.0.0
Sample collection and processing	collection device	GENEPIO:0001223	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.				1.0.0	1.0.0	1.0.0
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 [GENEPIO_00 02116]			1.0.0	1.0.0	1.0.0
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			1.0.0	1.0.0	1.0.0
Sample collection and processing	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)			1.0.0	1.0.0	1.0.0
Sample collection and processing	residual_sample_status	GENEPIO:0101090	The status of the residual sample (whether any sample remains after its original use).	Residual samples are samples that remain after the sample material was used for its original purpose. Select a residual sample status from the picklist. If sample still exists, select "Residua sample remaining (some sample left)".	(sample all used)			1.0.0	1.0.0	1.0.0
				The reason a sample was collected may provide information about potential biases in sampling strategy. Provide the purpose of sampling from the picklist in the template. Most likely, the sample was collected for Public health surveillance. The reason why a sample was originally collected may differ from the reason why it was selected for sequencing, which should be indicated in the "purpose of	Public health					
Sample collection and processing	purpose_of_sampling	GENEPIO:0001198	The reason that the sample was collected.	sequencing" field.	surveillance			1.0.0	1.0.0	1.0.0

		Ontology			Deprecated Label	Deprecated ID	Version		
Parent Class	Field	Identifier	Definition	Guidance	Examples	- Spresence II	Tracking		
Sample collection and processing	presampling_activity	GENEPIO:0100433	The activities or variables upstream of sample collection that may affect the sample.	If there was an activity that would affect the sample prior to collection (this is different than sample processing), provide the activities by selecting one or more values from the template pick list. If the Information is unknown or cannot be provided, leave blank or provide a null value.			1.0.0	1.0.0	1.0.0
Sample collection and processing	presampling_activity_details	GENEPIO:0100434	The details of the activities or variables that affected the sample collected.	Briefly describe the presampling activities using free text.	Agricultural waste from large farm contributes waste to the site sampled.		1.0.0	1.0.0	1.0.0
	p			Provide details of how the sample was stored from time of collection until time of processing. If there were issues with the cold chain storage,	The sample was placed in a tube in a cooler bag during transportation (~3 hours) to the lab site. At this point the sample was placed in storage medium and put in a -10C freezer until it was processed and extracted 5				
Sample collection and processing  Sample collection and processing	sample_storage_method sample_storage_medium	GENEPIO:0100448  GENEPIO:0100449	The process used to store the sample.  The medium in which a sample is stored.	note those here.  Provide the name of the transport medium or storage medium used for this sample. If none was used, leave blank or write "None".	days later.  Cary-Blair transport medium		1.0.0	1.0.0	1.0.0
Sample collection and processing	sample_storage_duration_value	GENEPIO:0101014	The numerical value of the time measurement during which a sample is in storage.	Provide the numerical value of time.	5		1.0.0	1.0.0	1.0.0
Sample collection and processing	sample_storage_duration_unit	GENEPIO:0101015	The units of a measured sample storage duration.	Provide the units from the pick list.	Day		1.0.0	1.0.0	1.0.0
Sample collection and processing	specimen_processing	GENEPIO:0001253	Any processing applied to the sample during or after receiving the sample.	Select processes from the picklist that were applied to this sample.	Centrifugation		1.0.0	1.0.0	1.0.0
			The details of the processing applied to the sample during or after		25 samples were pooled and further prepared as a single sample during library				
Sample collection and processing  Sample collection and processing	specimen_processing_details  experimental_protocol	GENEPIO:0100311  GENEPIO:0101029	receiving the sample.  The name of the overarching experimental methodology that was used to process the biomaterial.	sample.  Provide the name of the methodology used in your study. If available, provide a link to the protocol.	prep.		1.0.0	1.0.0	1.0.0
		2.12.13.10.1023		Samples can play different types of roles in experiments. A sample under study in one experiment may act as a control or be a replicate of another sample in another experiment. This field is used to distinguish samples under study from controls, replicates, etc. If the sample acted as an experimental control or a replicate, select a role type from the pictilist. If the sample	Positive				10.0
Sample collection and processing	experimental_specimen_role_type	GENEPIO:0100921	The type of role that the sample represents in the experiment.	was not a control, leave blank or select "Not Applicable".	experimental control		1.0.0	1.0.0	1.0.0
Sample collection and processing	experimental_specimen_details	GENEPIO:0101112					1.0.0	1.0.0	1.0.0
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 provided for more information.	Total coliform count [GENEPIO:010 0729]		1.0.0	1.0.0	1.0.0
			F	and data provider for more information.				1.0.0	

Parent Class	Field	Ontology Identifier	Definition	Guidance	Examples	Deprecated Label	Deprecated ID	Version Tracking		
Falelit Class	rietu	identiner	Definition	Use this field to provide free text details	Pooled metagenomes containing extended spectrum			3		
				describing other available data types that may provide context for interpreting genomic	beta-lactamase (ESBL)					
Sample collection and processing	available_data_type_details	GENEPIO:0101023	Detailed information regarding other available data types.	sequence data.	bacteria			1.0.0	1.0.0	1.0.0
Sequence information	Sequence information library_ID	GENEPIO:0001441  GENEPIO:0001448	The user-specified identifier for the library prepared for sequencing.	Every "library ID" from a single submitter must be unique. It can have any format, but we suggest that you make it concise, unique and consistent within your lab, and as informative as possible.	123446			1.0.0	1.0.0	1.0.0
Sequence information	sequencing_assay_type	GENEPIO:0100997	The overarching sequencing methodology that was used to determine the sequence of a biomaterial.	Example Guidance: Provide the name of the DNA or RNA sequencing technology used in your study. If unsure refer to the protocol documentation, or provide a null value.	whole genome sequencing assay [OBI:0002117]			1.0.0	1.0.0	1.0.0
Sequence information	sequencing_date	GENEPIO:0001447	The date the sample was sequenced.	ISO 8601 standard "YYYY-MM-DD".	2020-06-22					
Sequence information	purpose of sequencing	GENEPIO:0001445	The reason that the sample was sequenced.	The reason why a sample was originally collected may differ from the reason why it was selected for sequencing. The reason a sample was sequenced may provide information about potential biases in sequencing strategy. Provide the purpose of sequencing from the picklist in the template. The reason for sample collection should be indicated in the "purpose of sampling" field.				1.0.0	1.0.0	1.0.0
Sequence information	purpose_of_sequencing_details	GENEPIO:0001446	The description of why the sample was sequenced providing specific details.	Provide an expanded description of why the sample was sequenced using free text. The description may include the importance of the sequences for a particular public health investigation/surveillance activity/research question. Suggested standardized descriptions include: Assessing public health control measures, Determining early introductions and spread. Investigating aritine-related exposures, investigating remote regions, investigating health care workers, investigating schools/universities.	Investigating schools/universities			1.0.0	1.0.0	1.0.0
			The name of the agency, organization or institution responsible for	Provide the name of the agency, organization or institution that performed the sequencing in full (avoid abbreviations). If the information is unknown or cannot be provided, leave blank or	Canada (PHAC) [GENEPIO:010					
Sequence information	sequenced_by	GENEPIO:0100416	sequencing the isolate's genome.	provide a null value.	0551]			1.0.0	1.0.0	1.0.0
Sequence information	sequenced by laboratory name	GENEPIO:0100470	The specific laboratory affiliation of the responsible for sequencing the isolate's genome.	Provide the name of the specific laboratory that that performed the sequencing in full (avoid abbreviations). If the information is unknown or cannot be provided, leave blank or provide a nul value.	I Topp Lab			1.0.0	1.0.0	1.0.0
Sequence information	sequenced by contact name	GENEPIO:0100471	The name or title of the contact responsible for follow-up regarding the sequence.	Provide the name of an individual or their job title. As personnel turnover may render the contact's name obsolete, it is more 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			1.0.0	1.0.0	1.0.0
Sequence information	sequenced by contact email	GENEPIO:0100422	The email address of the contact responsible for follow-up regarding the sequence.	Provide the email associated with the listed contact. As personnel turnover may render an individual's email obsolete, it is more prefereable to provide an address for a position or lab, to ensure accuracy of information and institutional	enterics@lab.c			1.0.0	1.0.0	1.0.0
·				The name of the agency should be written out in full, (with minor exceptions) and be consistent across multiple submissions. For Canadian institutions submitting specimens rather than sequencing data, please put the "National	Public Health					
Sequence information	sequence_submitted_by	GENEPIO:0001159	The name of the agency that submitted the sequence to a database.  The email address of the contact responsible for follow-up regarding the	Microbiology Laboratory (NML)".  The email address can represent a specific	Ontario (PHO)  RespLab@lab.			1.0.0	1.0.0	1.0.0
Sequence information	sequence_submitter_contact_email	GENEPIO:0001165	sequence.	individual or laboratory.	ca			1.0.0	1.0.0	1.0.0

		Ontology Identifier			Deprecated Label	Deprecated ID	Version		
Parent Class	Field	Identifier	Definition	Guidance	Examples Direct		Tracking		
					wastewater RNA capture and purification via the "Sewage, Salt, Silica and SARS-CoV-2 (4S)" method v4 found at				
Sample collection and processing	nucleic_acid_extraction_method	GENEPIO:0100939	The process used to extract genomic material from a sample.	Briefly describe the extraction method used.	v4 round at https://www.pro tocols.io/view/v 4-direct-waste water-ma-captu re-and-purificati on-38wgq581y gk5/v4		1.0.0	1.0.0	1.0.0
Sample collection and processing	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.	QIAamp PowerFecal Pro DNA Kit		1.0.0	1.0.0	1.0.0
Sample collection and processing	endogenous control details	GENEPIO:0100923	The description of the endogenous controls included when extracting a sample.	Provide the names of endogenous controls that were used as a reference during extraction. If relevant, include titers of these controls, as well as whether any controls were expected but not identified in the sample.			1.0.0	1.0.0	1.0.0
Sequence information	sequencing_project_name	GENEPIO:0100472	The name of the project/initiative/program for which sequencing was performed.	Provide the name of the project and/or the project ID here. If the information is unknown or cannot be provided, leave blank or provide a null value.	AMR-GRDI (PA-1356)		1.0.0	1.0.0	1.0.0
Sequence information	sequencing_platform	GENEPIO:0100473	The platform technology used to perform the sequencing.	Provide the name of the company that created the sequencing instrument by selecting a value from the template pick list. If the information is unknown or cannot be provided, leave blank or provide a null value.	Illumina [GENEPIO:000 1923]		1.0.0	1.0.0	1.0.0
				Provide the model sequencing instrument by selecting a value from the template pick list. If the information is unknown or cannot be	Illumina HiSeq 2500 [GENEPIO:010				
Sequence information	sequencing_instrument	GENEPIO:0001452	The model of the sequencing instrument used.	provided, leave blank or provide a null value.	0117]		1.0.0	1.0.0	1.0.0
Sequence information							1.0.0	1.0.0	1.0.0
Sequence information	library_preparation_kit	GENEPIO:0001450	The name of the DNA library preparation kit used to generate the library being sequenced.	Provide the name of the library preparation kit used.	Nextera XT		1.0.0	1.0.0	1.0.0
Sequence information	DNA_fragment_length	GENEPIO:0100843	The length of the DNA fragment generated by mechanical shearing or enzymatic digestion for the purposes of library preparation.	Provide the fragment length in base pairs (do not include the units).	400		1.0.0	1.0.0	1.0.0
Sequence information	genomic_target_enrichment_method	GENEPIO:0100966	The molecular technique used to selectively capture and amplify specific regions of interest from a genome.	Provide the name of the enrichment method	Hybrid selection method (bait-capture) [GENEPI0::000 1950]		1.0.0	1.0.0	1.0.0
Sequence information	genomic_target_enrichment_method_details	GENEPIO:0100967	Details that provide additional context to the molecular technique used to selectively capture and amplify specific regions of interest from a genome.	Provide details that are applicable to the method you used. Note: If balt-capture methods were	enrichment was done using Twist's respiratory virus research panel: https://www.twi		1.0.0	1.0.0	1.0.0
			The specifications of the primers (primer sequences, binding positions, fragment size generated etc) used to generate the amplicons to be	Provide the name and version of the primer scheme used to generate the amplicons for					
Sequence information	amplicon_pcr_primer_scheme	GENEPIO:0001456	sequenced.	sequencing.	artic v3		1.0.0	1.0.0	1.0.0
Sequence information	amplicon_size	GENEPIO:0001449	The length of the amplicon generated by PCR amplification.	Provide the amplicon size expressed in base pairs.	300		1.0.0	1.0.0	1.0.0

		Ontology Identifier				Deprecated Label	Deprecated ID	Version		
Parent Class	Field	Identifier	Definition	Flow cells can vary in terms of design, chemistry, capacity, etc. The version of the 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				Tracking		
Sequence information	sequencing_flow_cell_version	GENEPIO:0101102	The version number of the flow cell used for generating sequence data.	not include "version" or "v" in the version number.	R.9.4.1			1.0.0	1.0.0	1.0.0
				Provide the name and version of the procedure or protocol used for sequencing. You can also	https://www.pro tocols.io/view/n cov-2019-sequ encing-protocol -bbmuik6w?ver sion warning=n					
Sequence information	sequencing_protocol	GENEPIO:0001454	The protocol or method used for sequencing.	provide a link to a protocol online.	Ω			1.0.0	1.0.0	1.0.0
Sequence information	r1_fastq_filename	GENEPIO:0001476	The user-specified filename of the r1 FASTQ file.	Provide the r1 FASTQ filename.	ABC123_S1_L 001_R1_001.fa stq.gz			1.0.0	1.0.0	1.0.0
Sequence information	r2_fastq_filename	GENEPIO:0001477	The user-specified filename of the r2 FASTQ file.	Provide the r2 FASTQ filename.	ABC123_S1_L 001_R2_001.fa stq.gz			1.0.0	1.0.0	1.0.0
Sequence information	fast5_filename	GENEPIO:0001480	The user-specified filename of the FAST5 file.	Provide the FAST5 filename. Provide the name and version number, with the	batch1a_seque nces.fast5			1.0.0	1.0.0	1.0.0
Sequence information	genome sequence file name	GENEPIO:0101715	The name of the sequence file.	Frovide the inaine and version furnine, with the file extension, of the processed genome sequence file e.g. a consensus sequence FASTA file or a genome assembly file.	mpxvassembly. fasta			1.0.0	1.0.0	1.0.0
Sequence information	assembly_filename	GENEPIO:0001461	The user-defined filename of the FASTA file.	Provide the FASTA filename.	pathogenasse mbly123.fasta			1.0.0	1.0.0	1.0.0
	Bioinformatics and QC metrics	GENEPIO:0001457						1.0.0	1.0.0	1.0.0
Bioinformatics and QC metrics	quality control method name	GENEPIO:0100557	The name of the method used to assess whether a sequence passed a predetermined quality control threshold.	Providing the name of the method used for quality control is very important for interpreting the rest of the QC information. Method names can be provided as the name of a pipeline or a link to a Gilt-lub repository. Multiple methods should be listed and separated by a semi-colon. Do not include QC tags in other fields if no method name is provided.	ncov-tools			1.0.0	1.0.0	1.0.0
Bioinformatics and QC metrics	quality control method version	GENEPIO:0100558	The version number of the method used to assess whether a sequence passed a predetermined quality control threshold.	Methods updates can make big differences to their outputs. Provide the version of the method used for quality control. The version can be expressed using whatever convention the developer implements (e.g. date, semantic versioning). If multiple methods were used, record the version numbers in the same order as the method names. Separate the version numbers using a semi-colon numbers using a semi-colon the province of the version numbers using a semi-colon.	1.2.3			1.0.0	1.0.0	1.0.0
Bioinformatics and QC metrics	quality control determination	GENEPIO:0100559	The determination of a quality control assessment.	Select a value from the pick list provided. If a desired value is missing, submit a new term request to the PHA4GE QC Tag GitHub	sequence failed quality control					
Bioinformatics and QC metrics	quality control issues	GENEPIO:0100560	The reason contributing to, or causing, a low quality determination in a quality control assessment.	Select a value from the pick list provided. If a desired value is missing, submit a new term request to the PHA4GE QC Tag GitHub	low average genome coverage			1.0.0	1.0.0	1.0.0
			The details surrounding a low quality determination in a quality control	Provide notes or details regarding QC results	CT value of 39. Low viral load. Low DNA concentration after					
Bioinformatics and QC metrics  Bioinformatics and QC metrics	quality control details  raw sequence data processing method	GENEPIO:0100561  GENEPIO:0001458	assessment.  The method used for raw data processing such as removing barcodes, adapter trimming, filtering etc.	using free text.  Raw data processing can have a significant impact on data quality and how it can be used. Provide the names and version numbers of software used for trimming adaptors, quality filtering, etc (e.g. Trimmomatto v. 0.38, Porechop v. 0.2.3), or a link to a Gilf-tub protocol.	amplification.  Porechop 0.2.3			1.0.0	1.0.0	1.0.0
Bioinformatics and QC metrics	dehosting method		The method used to remove host reads from the pathogen sequence.	Provide the name and version number of the software used to remove host reads.	Nanostripper			1.0.0	1.0.0	1.0.0

ld	Identifier	Definition	Guidance	Examples		Tracking		
			Provide the name of the software used to	SPAdes Genome Assembler, Canu, wtdbg2,				
uence assembly software name	GENEPIO:0100825	The name of the software used to assemble a sequence.	assemble the sequence.	velvet		1.0.0	1.0.0	1.0.0
			Provide the version of the software used to					
uence assembly software version	GENEPIO:0100826	The version of the software used to assemble a sequence.	assemble the sequence.	3.15.5		1.0.0	1.0.0	1.0.0
sensus sequence software name	GENEPIO:0001463	The name of the software used to generate the consensus sequence.	Provide the name of the software used to generate the consensus sequence.	iVar		1.0.0	1.0.0	1.0.0
sensus sequence software version	GENEPIO:0001469	The version of the software used to generate the consensus sequence.	Provide the version of the software used to generate the consensus sequence.	1.3	3	1.0.0	1.0.0	1.0.0
adth of coverage value	GENEPIO:0001472	The percentage of the reference genome covered by the sequenced data, to a prescribed depth.	Provide value as a percent.	95	5	1.0.0	1.0.0	1.0.0
			·					
th of coverage value	GENEPIO:0001474	The average number of reads representing a given nucleotide in the reconstructed sequence.	Provide value as a fold of coverage.	400		1.0.0	1.0.0	1.0.0
th of coverage threshold	GENEPIO:0001475	The threshold used as a cut-off for the depth of coverage.	Provide the threshold fold coverage.	100		1.0.0	1.0.0	1.0.0
ome completenese	GENEDIO:0100844	The percentage of expected genes identified in the genome being sequenced. Missing genes indicate missing genomic regions (incompleteness) in the detail.	Provide the genome completeness as a percent	98		100	100	1.0.0
one completeness	GENEFIO.0100044	(incompleteness) in the data.	(no need to include units).	00	,	1.0.0	1.0.0	1.0.0
nber of base pairs sequenced	GENEPIO:0001482	The number of total base pairs generated by the sequencing process.	Provide a numerical value (no need to include units).	387566	3	1.0.0	1.0.0	1.0.0
nber of total reads	GENEPIO:0100827	The total number of non-unique reads generated by the sequencing process.	Provide a numerical value (no need to include units).	423867		1.0.0	1.0.0	1.0.0
nber of unique reads	GENEPIO:0100828	The number of unique reads generated by the sequencing process.	Provide a numerical value (no need to include units).	248236	3	1.0.0	1.0.0	1.0.0
imum nost-trimming read length	GENEPIO:0100829	The threshold used as a cut-off for the minimum length of a read after	Provide a numerical value (no need to include units)	150		100	100	1.0.0
intum post-timining read length	OENET 10.0100023		unia).	100		1.0.0	1.0.0	1.0.0
nber of contigs	GENEPIO:0100937	The number of contigs (contiguous sequences) in a sequence assembly.	Provide a numerical value.	10		1.0.0	1.0.0	1.0.0
cent Ns across total genome length	GENEPIO:0100830	The percentage of the assembly that consists of ambiguous bases (Ns).	Provide a numerical value (no need to include units).	2	2	1.0.0	1.0.0	1.0.0
per 100 kbp	GENEPIO:0001484	The number of ambiguous bases (Ns) normalized per 100 kilobasepairs (kbp).	Provide a numerical value (no need to include units).	342		1.0.0	1.0.0	1.0.0
	GENEPIO:0100938	The length of the shortest read that, together with other reads, represents at least 50% of the nucleotides in a set of sequences.	Provide the N50 value in Mb.	150		1.0.0	1.0.0	1.0.0
cent read contamination	GENEPIO:0100845	The percent of the total number of reads identified as contamination (not belonging to the target organism) in a sequence dataset	Provide the percent contamination value (no need to include units)	,		1.0.0	100	1.0.0
, , , , , , , , , , , , , , ,								
uence assembly length	GENEPIO:0100846	scaffold or by reference-based mapping.	units).	34272		1.0.0	1.0.0	1.0.0
sensus genome length	GENEPIO:0001483	The length of the genome defined by the most common nucleotides at each position.	Provide a numerical value (no need to include units).	38677	,	1.0.0	1.0.0	1.0.0
rence genome accession	GENEPIO:0001485	A persistent, unique identifier of a genome database entry	Provide the accession number of the reference genome.	NC 045512.2		1.0.0	1.0.0	1.0.0
s s a t t c n n n ii	th of coverage value  h of coverage value  h of coverage threshold  me completeness  ber of base pairs sequenced  ber of total reads  ber of unique reads  mum post-trimming read length  ber of contigs  ent Ns across total genome length  ent read contamination  sence assembly length	tensus sequence software name  GENEPIO.0001463  GENEPIO.0001469  GENEPIO.0001472  GENEPIO.0001474  GENEPIO.0001474  GENEPIO.0001475  GENEPIO.0001475  GENEPIO.0001475  GENEPIO.0001475  GENEPIO.0001482  GENEPIO.0001482  GENEPIO.0100827  GENEPIO.0100828  GENEPIO.0100829  GENEPIO.0100829  GENEPIO.0100830  GENEPIO.0100830  GENEPIO.0100830  GENEPIO.0100836  GENEPIO.0100836  GENEPIO.0100836  GENEPIO.0100846  GENEPIO.0100846  GENEPIO.0100846	GENEPIC-0001483 The name of the software used to generate the consensus sequence.  GENEPIC-0001499 The version of the software used to generate the consensus sequence.  The percentage of the reference genome covered by the sequenced data, is a prescribed depth.  The average number of reads representing a given nucleotide in the reconstructed sequence.  GENEPIC-0001472 The the percentage of expected genes identified in the genome being sequenced. Missing genes indicate missing genomic regions (excomplateness) in the data.  GENEPIC-0001482 The percentage of expected genes identified in the genome being sequenced. Missing genes indicate missing genomic regions (excomplateness) in the data.  GENEPIC-0100844 The number of total base pairs generated by the sequencing process.  The total number of non-unique reads generated by the sequencing process.  The threshold used as a cut-off for the minimum length of a read after transmit.  The number of unique reads generated by the sequencing process.  The threshold used as a cut-off for the minimum length of a read after transmit.  GENEPIC-0100829 The number of unique reads generated by the sequencing process.  The threshold used as a cut-off for the minimum length of a read after transmit.  The number of unique reads generated by the sequencing process.  The number of unique reads generated by the sequencing process.  The number of antique reads generated by the sequencing process.  The number of antique reads generated by the sequencing process.  The number of antique reads generated by the sequencing process.  The number of antique reads generated by the sequencing process.  The number of antique reads generated by the sequencing process.  The number of antique reads generated by the sequence after transmit.  The number of configs (configuous bases (No)) normalized per 100 kilobasepairs (Rep).  GENEPIC-0100839 The length of the genome generated by assembling reads using a scaffold or by reference-based mapping.  The length of the genome defined by the most common nucleot	GENEPIO 0101433 The name of the software used to generate the consensus sequence.  GENEPIO 0001443 The version of the software used to generate the consensus sequence.  Provide the name of the software used to generate the consensus sequence.  Provide the version of the software used to generate the consensus sequence.  Provide the version of the software used to generate the consensus sequence.  Provide the version of the software used to generate the consensus sequence.  Provide the version of the software used to generate the consensus sequence.  Provide the version of the software used to generate the consensus sequence.  Provide the version of the software used to generate the consensus sequence.  The version of the software used to generate the consensus sequence.  Provide the version of the software used to generate the consensus sequence.  Provide the version of the software used to generate the consensus sequence.  Provide the version of the software used to generate the consensus sequence.  Provide the version of the software used to generate the consensus sequence.  Provide the version of the software used to generate the consensus sequence.  Provide the version of the software used to generate the consensus sequence.  Provide the version of the software used to generate the consensus sequence.  Provide the version of the software used to generate the consensus sequence.  Provide the version of the software used to generate the consensus sequence.  Provide the version of the software used to generate the consensus sequence.  Provide the version of the software used to generate the consensus sequence.  Provide the version of the software used to generate the consensus sequence.  Provide the version of the software used to generate the consensus sequence.  Provide the version of the software used to generate the consensus sequence.  Provide the version of the software used to generate the consensus sequence.  Provide the version of the software used to generate the provide in the generate the provide t	annum sequence software name  GENEPIO.0001469. The name of the software used to generate the consensus sequence.  GENEPIO.0001469 The version of the software used to generate the consensus sequence.  GENEPIO.0001460 The version of the software used to generate the consensus sequence.  The personal part of an influence personal coverad by the sequenced of the software used to generate the consensus sequence.  The personal part of the influence used to generate the consensus sequence.  The personal part of the influence personal coverad by the sequenced of the software used to generate the consensus sequence.  The personal part of the influence personal coverad by the sequenced of the influence used to generate the consensus sequence.  The personal part of the influence personal coverad by the sequenced of the influence of the influence used to generate the consensus sequence.  The personal part of the influence personal coverage personal coverage personal coverage personal coverage threshold  GENEPIO.0001172 The threshold used as a cut-off for the depth of coverage.  The threshold used as a cut-off for the depth of coverage.  The threshold used as a cut-off for the depth of coverage.  The threshold used as a cut-off for the depth of coverage.  Provide the threshold fold coverage.  Provide the threshold scale as a percent personal coverage threshold.  GENEPIO.0001482 The number of from unique reads generated by the sequencing process.  The bold number of from unique reads generated by the sequencing process.  The bold number of from unique reads generated by the sequencing process.  The threshold used as a cut-off for the minimum length of a read after threshold scale as a cut-off for the minimum length of a read after threshold.  The threshold used as a cut-off for the minimum length of a read after threshold and the personal personal coverage.  Provide a numeric	SCHEPTO-000143 The name of this software under the	actives sequence coffware version  CELEPTOLOGISTAL  The reservor of the orthware used to generate the configuration of the coffware used to generate the construction sequence.  CELEPTOLOGISTAL  The reservor of the coffware used to generate the construction sequence.  CELEPTOLOGISTAL  The reservor of the coffware used to generate the construction sequence.  CELEPTOLOGISTAL  The reservor of the coffware used to generate the construction sequence.  CELEPTOLOGISTAL  The reservor of the coffware used to generate the construction sequence.  The reservor of the coffware used to generate the construction sequence.  CELEPTOLOGISTAL  The reservor of the coffware used to generate the construction sequence.  The reservor of the coffware used to generate the construction sequence.  The security of the subservor and used to the first sequence.  The security of the subservor and used to the first sequence.  The security of the subservor and used to the first sequence.  CELEPTOLOGISTAL  The security of the security growth reservor generate construction from the security growth in sequence of the security growth in security growth in sequence of the security growth in s	entions sequence software name  GENEPO0001497  The service of the entioners used to generate the consensus sequence software name  GENEPO0001497  The service of the entioners used to generate the consensus sequence software viriation  GENEPO0001497  The service of the entioners used to generate the consensus sequence software used to generate the consensus sequences of the security of the sequence of the sequences of the consensus sequences and the consensus sequences software used to generate the consensus sequences of the sequences o

		Ontology			D	precated Label	Donge control ID	Version		
Parent Class	Field	Ontology Identifier	Definition	Guidance	Examples	eprecated Laber	Deprecated ID	Tracking		
ioinformatics and QC metrics	deduplication method	GENEPIO:0100831	The method used to remove duplicated reads in a sequence read dataset.	Provide the deduplication software name followed by the version, or a link to a tool or method.	DeDup 0.12.8			1.0.0	1.0.0	1.0.0
				Further details regarding the methods used to process raw data, and/or generate assemblies, 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	https://github.c					
oinformatics and QC metrics	bioinformatics protocol	GENEPIO:0001489	A description of the overall bioinformatics strategy used.	link to a pipeline or workflow.	f			1.0.0	1.0.0	1.0.0
	Taxonomic identification information	GENEPIO:0101082						1.0.0	1.0.0	1.0.0
			The name of the software used to map sequence reads to a reference		Bowtie2, BWA-MEM,					
xonomic identification information	read mapping software name	GENEPIO:0100832	genome or set of reference genes.	Provide the name of the read mapping software.	TopHat		,	1.0.0	1.0.0	1.0.0
xonomic identification information	read mapping software version	GENEPIO:0100833	The version of the software used to map sequence reads to a reference genome or set of reference genes.	Provide the version number of the read mapping software.	2.5.1			1.0.0	1.0.0	1.0.0
			The name of the taxonomic reference database used to identify the	Provide the name of the taxonomic reference						
xonomic identification information	taxonomic reference database name	GENEPIO:0100834	organism.	database.	NCBITaxon					
avanamia idantification information	tavanania usfavana adatabasa varsian	CENEDIO 0400925	The version of the taxonomic reference database used to identify the	Provide the version number of the taxonomic reference database.	1.3			100	100	100
axonomic identification information	taxonomic reference database version	GENEPIO:0100835	organism.	reference database.	1.3			1.0.0	1.0.0	1.0.0
xonomic 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.doc			1.0.0	1.0.0	1.0.0
				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						
xonomic identification information	taxonomic analysis date	GENEPIO:0101075	The date a taxonomic analysis was performed.	format, i.e. "YYYY-MM-DD".	2024-02-01			1.0.0	1.0.0	1.0.0
xonomic identification information	read mapping criteria	GENEPIO:0100836	A description of the criteria used to map reads to a reference sequence.	Provide a description of the read mapping criteria.	Phred score >20			1.0.0	1.0.0	1.0.0
	Pathogen diagnostic testing	GENEPIO:0001506						1.0.0	1.0.0	1.0.0
				Provide the full name of the gene used in the test. Standardized gene names can be found in						
- 44		GENEPIO:0101116	The control of the control of the Australia	the Gene Ontology using this look-up service:				1.0.0	1.0.0	1.0.0
athogen diagnostic testing	genetic target name	GENEPIO:0101116	The name of the genetic marker used for testing.	https://bit.ly/2Sq1Lbl	gyrase A			1.0.0	1.0.0	1.0.0
thogen diagnostic testing	genetic target region	GENEPIO:0101117						1.0.0	1.0.0	1.0.0
thogen diagnostic testing	genetic target region reference genome	GENEPIO:0101118								
athogen diagnostic testing	diagnostic target presence	GENEPIO:0100962	The binary value of the result from a diagnostic test.	Select a value from the pick list provided, to describe whether a target was determined to be present or absent within a sample.	diagnostic target present			1.0.0	1.0.0	1.0.0
				Provide the numerical result of a diagnostic test						
athogen diagnostic testing	diagnostic measurement value	GENEPIO:0100963	The value of the result from a diagnostic test.	(no need to include units).	1000			1.0.0	1.0.0	1.0.0
				Select a value from the pick list provided, to	cycle threshold					
thogen diagnostic testing	diagnostic measurement unit	GENEPIO:0100964	The unit of the result from a diagnostic test.	describe the units of the given diagnostic test.	(Ct)			1.0.0	1.0.0	1.0.0
		051/50/00/00/00		Select a value from the pick list provided to describe the method used for a given diagnostic	202					
thogen diagnostic testing	diagnostic measurement method	GENEPIO:0100965	The method by which a diagnostic result was determined.	test.	qPCR			1.0.0	1.0.0	1.0.0
athogen diagnostic testing	diagnostic testing threshold value	GENEPIO:0101104						1.0.0	1.0.0	1.0.0
0g		22.12.12.101101								
thogen disappetic testing	diagnostic testing threshold units	GENEPIO:0101105					.	1.0.0	1.0.0	1.0.0
athogen diagnostic testing	uragnostic testing threshold units	GENEPIO.0101105						1.0.0	1.0.0	1.0.0

Parent Class	Field	Ontology Identifier	Definition	Guidance	Examples	Deprecated Label	Deprecated ID	Version Tracking		
Pathogen diagnostic testing	diagnostic testing details	GENEPIO:0101106						1.0.0	1.0.0	1.0.0
	Risk assessment information	GENEPIO:0100478						1.0.0	1.0.0	1.0.0
Risk assessment information	prevalence_metrics	GENEPIO:0100480	Metrics regarding the prevalence of the pathogen of interest obtained from a surveillance project.	Risk assessment requires detailed information regarding the quantities of a pathogen in a specified location, commodity, or environment. As such, it is useful for risk assessors to know what types of information are available through documented methods and results. Provide the metric types that are available in the surveillance project sample plan by selecting them from the pick list. The metrics of interest 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.				1.0.0	1.0.0	1.0.0
Risk 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.	Hazard organism counts (i.e. Salmonella) do not distinguish between serovars.			1.0.0	1.0.0	1.0.0
Risk assessment information	stage_of_production	GENEPIO:0100482	The stage of food production.	Provide the stage of food production as free text.	Abattoir [ENVO:010009 25]					
Risk assessment information	experimental_intervention	GENEPIO:0100483	The category of the experimental intervention applied in the food production system.	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 list provided.	Vaccination [NCIT:C15346]			1.0.0	1.0.0	1.0.0
Risk assessment information	experiment_intervention_details	GENEPIO:0100484	The details of the experimental intervention applied in the food production system.	If an experimental intervention was applied in the survey, provide details in this field as free text.				1.0.0	1.0.0	1.0.0