





© Guidance for populating GenomeTrakr metadata templates (BioSample and SRA)

V.7



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PURPOSE: Guidance on how to populate NCBI's metadata packages, maximizing interoperability for foodborne pathogen surveillance.

SCOPE: This protocol provides detailed instructions for populating the following two templates:

- 1. **BioSample metadata**: guidelines to populate the GenomeTrakr-extended pathogen package.
- 2. **SRA metadata:** NCBI's generic sequence metadata template for SRA submissions.

Versions:

v6: Added the One Health Enteric package presented at IAFP 2021 meeting. v7: Updated the picklists in the GenomeTrakr-extended pathogen package, "GT-pathogen package-OHE v0.2.2.xlsx" and added an incremental update file for the DRAFT One Health Enteric Package that includes extensive edits compared to v6.

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protocols.io

https://dx.doi.org/10.17504/protocols.io.by2mpyc6 Ruth Timme

protocol

Timme, R.E., Wolfgang, W.J., Balkey, M. et al. Optimizing open data to support one health: best practices to ensure interoperability of genomic data from bacterial pathogens. One Health Outlook 2, 20 (2020). https://doi.org/10.1186/s42522-020-00026-3

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Updated the picklists in the GenomeTrakr-extended pathogen package, "GT-pathogen package-OHE v0.2.2.xlsx" and added an incremental update file for the DRAFT One Health Enteric Package that includes extensive edits compared to v6.

GenomeTrakr, metadata, Pathogen package, NCBI Pathogen Detection, INSDC

_____ protocol,

Oct 13, 2021

Nov 10, 2021

54061

Part of collection

Utilizing the Public GenomeTrakr Database for Foodborne Pathogen Traceback Utilizing the Public GenomeTrakr Database for Foodborne Pathogen Traceback

Gather the following contextual information for each pure culture isolate:

- 1. organism name
- 2. lab name that collected the sample
- 3. collection date
- 4. collection source
- 5. Geographic location of sample collection

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Before collecting sequence data for your isolates, ensure that you can provide the minimum metadata recommended by your coordinating surveilliance body. The INSDC, in collaboration with the Global Microbial Identifer (GMI)

(<u>https://www.globalmicrobialidentifier.org</u>), recommends using the Pathogen metadata template for pathogen surveilliance submissions: (NCBI:

https://www.ncbi.nlm.nih.gov/pathogens/submit-data/and EMBL-EBI: https://www.ebi.ac.uk/ena/submit/pathogen-data).

Overview

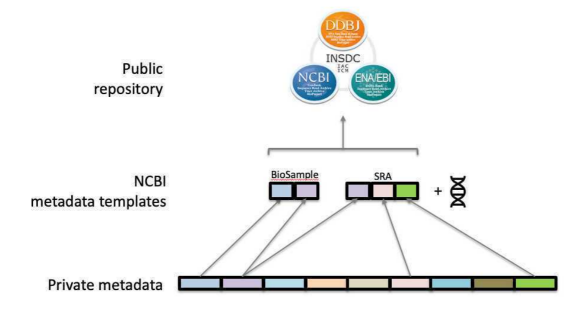
1 Guidance for organizing and populating the metadata templates required for direct submission to NCBI. This guidance is applicable for most enterics and/or microbial pathogens.

****If your laboratory uses the BioNumerics platform for submission, please follow this <u>protocol</u>.****

Two metadata templates are required:

- 1. BioSample metadata (metadata describing the sample source and submitter)
- 2. SRA metadata (metadata describing the sequence data collection)





BioSample metadata template

2 Template for BioSample submission:

Download the **GenomeTrakr-extended pathogen package** and follow the guidance included in this template.

■ GT-pathogen package-OHE v0.2.2.xlsx

- 2.1 DRAFT One Health Enteric Package, announced at IAFP 2021 (and the fall GenomeTrakr meeting) is ready for review and comment. Please download the file, review the attributes, and provide feedback here or email directly to ruth.timme@fda.hhs.gov.
 - One Health Enteric Package-DRAFT_v0.5.1-Nov1.xlsx



NOTE: DO NOT USE THIS VERSION FOR NCBI SUBMISSION.

We expect the v1.0 release to be ready for use in Fall 2021.

SRA sequence metadata template

3 Template for SRA metadata submission:



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Citation: Ruth Timme, Maria Balkey, William Wolfgang, Errol Strain Guidance for populating GenomeTrakr metadata templates (BioSample and SRA) https://dx.doi.org/10.17504/protocols.io.by2mpyc6

Download the "<u>Metadata spreadsheet with sample names</u>" file from the NCBI Submission Templates page:

https://submit.ncbi.nlm.nih.gov/templates/

And follow the guidance in the following table:

PRO TIPS:

- 1. If you have sequences to submit that belong to more than one BioProject, create a separate submission + metadata table for each of your BioProjects.
- 2. Entering fastq filenames in the spreadsheet. On a Mac, you can directly copy the file names from the folder into a spreadsheet. This is not possible on a PC using copy and paste but can be done with some command-line operation.
- 3. Finally, it is important to develop a QA/QC step to make sure the files are associated with the correct sample name. For example, use a left function in excel to strip of the appended text in the file name and then use the exact match to make sure the name matches the sample name.

3.1

Α	В	С
Field	Description	Example
sample_name	Include the same ID here as you entered for "sample_name" in the	UT-12345
	BioSample submission template.	
	Populate this field using the values	
	in the PHA4GE specification for	
	"specimen collector sample ID".	
library_ID	The library name should be a	UT-12345.6
	unique ID relevant to your	
	workflow. It can be an	
	autogenerated ID from your LIMS	
	system or a modification of your	
	sample_name.	
	Populate this field using the values	
	in the PHA4GE specification for	
	"library_id".	
Title	Short, free text description that	WGS of Salmonella enterica:
	identifies the data on public pages.	UT-12345
	For Example:	
	{methodology} of {organism}:	
	{sample_name}	



library_strategy	Overall sequencing strategy or approach.	See NCBI SRA pick list. (e.g. WGS)
	Choose from NCBI pick list	
library_source	molecule type used to make the	See NCBI SRA pick list. (e.g.
	library	Genomic)
library_selection	Library capture method	See NCBI SRA pick list. (e.g.
		random, PCR)
Library_layout	Choose from NCBI pick list	See NCBI SRA pick list,
		choose "paired"
platform	Sequencing platform	See NCBI SRA pick list. (e.g.,
		Illumina).
instrument_model	Name of the sequencing	See NCBI SRA pick list. (e.g.
	instrument.	Illumina MiSeq, iSeq 100)
Design_description	optional field for free text	
	description of methods	
Filetype	File format name for the raw	See NCBI SRA pick list. (e.g.
	sequence data	Fastq)
	Choose from NCBI pick list	
Filename	include ALL of the files resulting	genome_r1.fastq (*must be
	from this library. **Add additional	exact)
	fields if there are more than two	
	files (e.g. Filename3).	
	Populate this field using the values	
	in the PHA4GE specification for "r1	
	fastq filename".	
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Filename2	genome_r2.fastq (*must be exact)	genome_r2.fastq (*must be exact)
	Populate this field using the values	
	in the PHA4GE specification for "r2	
	fastq filename".	
Filename3-8	list other fastq file names (e.g. for	
	NextSeq data)	

Save the second sheet (SRA_data) as a TSV (tab-delimited file) for upload in the "SRA metadata" tab within the submission portal.

^{*}NCBI should also accept the original excel formatted file.