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© Populating NCBI template for submissions using BioNumerics v7.6 V.3

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GenomeTrakr

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PURPOSE: to define the standard operating procedure for collecting isolate metadata using BioNumerics for submission of food/environmental isolates to NCBI.

SCOPE: to provide a standardized procedure to collect isolate metadata using BioNumerics for submission of food/environmental isolates to NCBI.

RESPONSIBILITIES- SOP Responsible Officials: Ruth Timme, Maria Balkey

The GenomeTrakr Network Management will be responsible to monitor GenomeTrakr submissions processed through Bionumerics and ensure that all GT labs are familiar with the mandatory metadata fields required for submission of GenomeTrakr sequencing records to NCBI.

V3: Dropdown menus from controlled vocabulary added to the ncbi_update submission sheet

DOI

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NCBI submission, BioNumerics, biosample, SRA, metadata, bioproject

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1 Metadata SampleSheet preparation

Before uploading your sequencing run or linking NCBI sequencing records at the BioNumerics platform make sure to fill out the metadata spreadsheet form.

Please download the template and guidelines included in the file 'GT_BioNumerics_spreadsheet_v2.xlsx'.

Create the fields NCBI_bioproject, Attribute_package, Organism_name, NCBI_LabID, SourceCountryState, Latitude_longitude, Reference_material, Culture_collection or Description if **they are not** in the BioNumerics interface and are needed to process the metadata for your isolates.

Once you have filled out the template information, save the **template sheet** as **.csv** and import the metadata to BioNumerics.

@ GT_BioNumerics_spreadsheet_v2.xlsx

2 NCBI Submission Settings (Manage submission template)

Create the NCBI metadata template in BioNumerics following PulseNet instructions making sure fields are populated according to GT requirements which are described in the following steps.

2.1 BioProject and Organization: GenomeTrakr labs by submitting independently become owners of their data and are responsible for managing individual bioprojects for each sequenced organism. The term 'field content' denotes that the template value e.g. BioProject accession is mapping to the field in BioNumerics e.g. NCBI_bioproject.

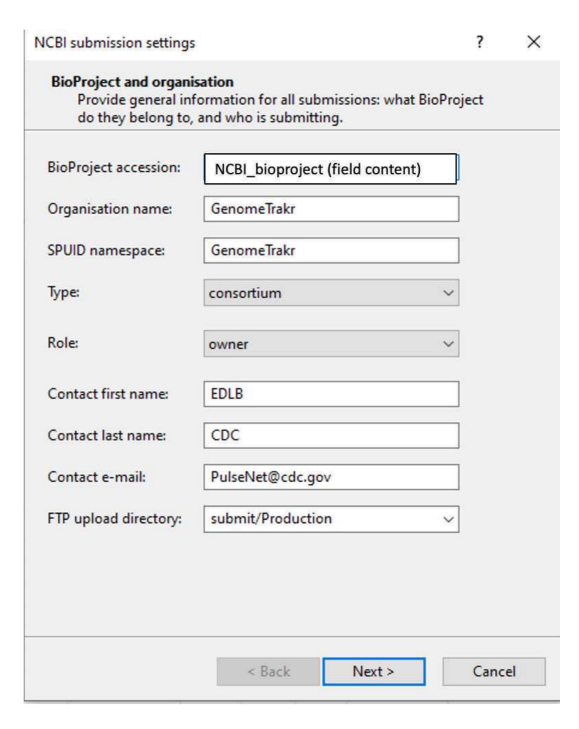


Fig 1. NCBI Submission Template: BioProject and Organization

Α	В	С
Name of Field in	Description	Example
BioNumerics Template		
BioProject accession	Identifier for NCBI data	PRJNA514285
	collection that contatins data	
	associated to	
	GenomeTrakr. Specific for	
	organism and lab submitter	
Organization name	Surveillance Program	GenomeTrakr
	(example is the default value	
	for GenomeTrakr	
	submissions)	
SPUID namespace	Surveillance Program	GenomeTrakr
	(example is the default value	
	for GenomeTrakr	
	submissions)	
Туре	organization type (example is	consortium
	the default value for	
	GenomeTrakr submissions)	
Role	laboratory responsibility	owner
	(example is the default value	
	for GenomeTrakr	
	submissions)	
Contact first name	First name for Lab POC for	First Name
	NCBI submissions. Lab might	
	choose to create alias name	
	for WGS team	
Contact last name	Last name Lab POC for NCBI	Last Name
	submissions. Lab might	
	choose to create alias name	
	for WGS team	
Contact e-mail	email for Lab POC for NCBI	first.last@lab.gov
	submissions. Lab might	
	choose to create alias name	
	for WGS team	
FTP upload directory	Name of directory at NCBI	submit/Production
	FTP site (example is the	
	default value for	
	GenomeTrakr submissions)	

Table 1. Guidelines for Bioproject and Organization metadata

Laboratories will be submitting to specific bioprojects for lab/organisms. Find the



2.2 organism/lab specific bioproject under each of the GenomeTrakr umbrella bioprojects included at https://www.ncbi.nlm.nih.gov/bioproject/593772

Make sure to submit to your lab bioproject. **Please don't submit to umbrella bioprojects.**

2.3 BioSample: Metadata associate to the isolate might require the creation of new fields in BioNumerics. The term 'field content' denotes that the template value e.g. Organism name is mapping to the field in BioNumerics e.g. Organism_name. The template values might map to default values e.g. Pathogen: environmental/food/other; version 1.0. Make sure to include the metadata associated to the isolates in the mandatory fields such as: Submitter Provided Unique ID, BioSample accession (output), Organism name, Title, Attribute package, Strain name and Isolate name alias. Isolate name alias is a mandatory field for GenomeTrakr submissions. Provide serovar when available.

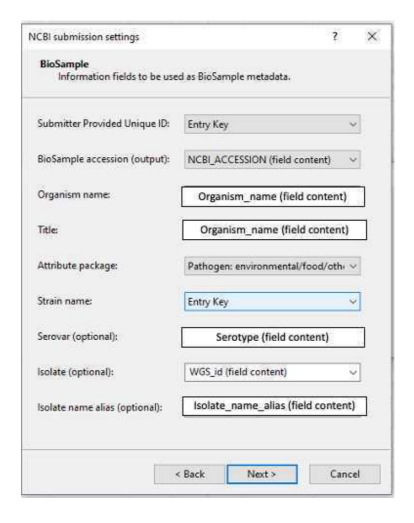


Fig 2. NCBI Submission Template:BioSample

Α	В	С	D
Name of	Description	Name of Field in	Example of metadata
Field in		BioNumerics DataBase	value
BioNumerics			
Template			
Submitter	Local lab strain	Entry Key	21B00181-5
Provided	ID		
Unique ID			
BioSample	NCBI accession	NCBI_ACCESSION (field	SAMN17385051
accession	will get	content)	
(output)	populated		
	upon		
	submission to		
	NCBI		
Organism	Genus -	Organism_name (field	Listeria monocytogenes
name	species for	content)	
	organism		
Title	Organism	Organism_name (field	Listeria monocytogenes
	name	content)	
Attribute	Sample	Pathogen:	Pathogen:
package	category	environmental/food/other;	environmental/food/other;
		version 1.0	version 1.0
Strain name	PNUSA	WGS_id (field content)	PNUSAL008933
	identifier		
	(automatically		
	populates at		
	the time of		
	registration)		
Serovar	Serotyping	Serovar (field content)	missing
(optional)	information for		
	Escherichia coli		
	and		
	Salmonella		
	enterica		
Isolate	Field is not	<missing></missing>	missing
(optional)	required for		
	GenomeTrakr		
Isolate namea	Optional	Isolate_name_alias (field	21B00181-5; RS_21290
alias (optional)	identifier for	content)	
	collaboration		
	projects		

Table 2. Guidelines for BioSample metadata

2.4 BioSample: Make sure to include the metadata associated to the isolates in the mandatory fields such as: Collected by, Collection / Isolate date, Collection / Isolate date format, Title, Geographical origin and Isolate source. Isolate name alias is a mandatory field for GenomeTrakr submissions. Provide Geographical coordinates when available. Host or host disease are provided only for isolates obtained from human, indicate "missing" for isolates from food or environmental sources.

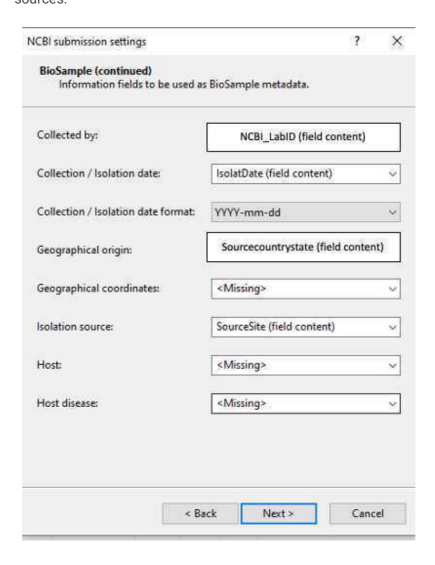


Fig 2. NCBI Submission Template: BioSample_2

Α	В	С	D
Name of	Description	Name of Field in	Example of
Field in		BioNumerics	metadata value
BioNumerics		DataBase	
Template			
Collected by	Full name of	NCBI_LabID (field	NY Department of
	laboratory that	content)	Agriculture and
	collected the sample		Markets
	or has taken over		
	curation of the		
	isolate.		
Collection date	Date on which the	IsolateDate (field	2020
	sample was	content)	
	collected.		
Geographical	Country and State	SourceCountryState	USA:NY
location	for sample	(field	
	collection	content)	
Geographical	latitude and	<missing></missing>	missing
coordinates	longitude for site		
	of collection.		
	Missing if it is not		
	provided		
Isolation source	Detailed description	SourceSite (field	cheese
	for sample	content)	
	product or		
	environmental		
	source		
Host	Only provided for	<missing></missing>	missing
	human		
	isolates		
Host disease	Only provided for	<missing></missing>	missing
	human		
	isolates		

Table 3. Guidelines for BioSample metadata (2)

2.5 NCBI submission settings – SRA Experiment and Run

Populate fields for SRA Experiment and Run according to PulseNet instructions.

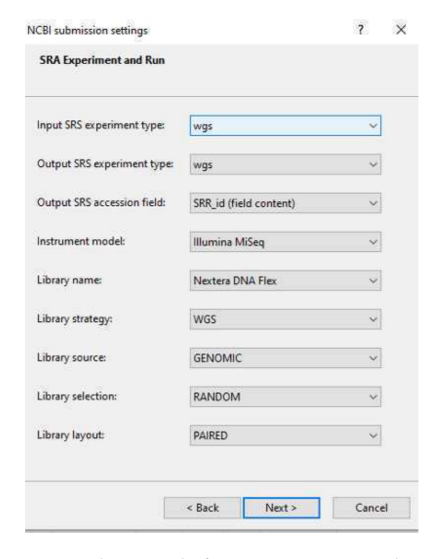


Fig 4. NCBI Submission Template forBioNumerics, SRA Experiment and run: Make sure to map collection attributes to the corresponding fields.

3 NCBI submission settings - Submission Template

Save submission template according to PulseNet Instructions as -GenomeTrakr-Template-.

4 Import data

4.1 Import the *GenomeTrakr Metadata form for BioNumerics*(GT_BioNumerics_spreadsheet_v2.csv) according to PulseNet Instructions.

- 4.2 When importing rules, the field source should match destination fields.
- 4.3 In the importing links section, choose the -key- for linking records to database entries.
- **4.4** Proceed with sequencing data import according to PulseNet Instructions.
- 4.5 Submit data to NCBI according to PulseNet Instructions. If NCBI accessions are not available at BioNumerics in 1 business day, please contact NCBI and PulseNet to troubleshoot issues with submissions.
- 4.6 Contact GenomeTrakr by email genomeTrakr if issues with submissions are delayed for more than 3 days. GenomeTrakr can support urgent submissions if needed.
- 5 NCBI submission for fields not included in the BioNumerics Template.

Laboratories need to include the name of the laboratory sequencing the isolates and the surveillance effort name in the **sequenced_by** and **project_name** fields, respectively. After receiving biosample accessions, fill out the <code>@ BioNumerics_update.xlsx</code> spreadsheet and submit the update for these fields to NCBI by contacting biosamplehelp@ncbi.nlm.nih.gov.