

# Quick-Start Protocol for GMI Proficiency Test, 2017

---

HISTORY OF CHANGES; version 3

Under 3.1 corrected to: *there should be only 14 st, 11 ec, and 11 sa samples in each file – no more no less*

---

This document describes the basic instructions for both the dry and wet lab components of the PT. See the full Protocol document including appendices for more information:

<http://www.globalmicrobialidentifier.org/workgroups/about-the-gmi-proficiency-test-2017>.

## 1 OVERVIEW

The proficiency test, 2017, consists of three parts:

- 1a. DNA extraction, purification, library-preparation, and whole-genome-sequencing (WGS) from **live cultures**
- 1b. Whole-genome-sequencing of **pre-prepared DNA**
2. Phylogenetic/clustering analysis of three **fastq datasets**

## 2 SHIPPING, RECEIPT AND STORAGE OF BACTERIAL STRAINS

All bacterial strains and DNA are shipped as UN3373, Biological substance category B. **Please confirm receipt of the parcel through the confirmation form enclosed in the shipment.**

## 3 PROCEDURE AND ANALYSIS OF TEST MATERIAL

### 3.1 Bacterial cultures and DNA

Subculture the bacterial strains on a relevant growth medium of the laboratory's own choice and incubate. Following incubation and assessment of purity of the bacterial cultures, perform DNA extraction and whole-genome-sequencing according to the laboratory's standard procedure.

For the purified PT-DNA received, perform whole-genome-sequencing according to the laboratory's standard procedure.

The metadata file and WGS data must be submitted as a batch-upload. The web-interface of the (<https://cge.cbs.dtu.dk/services/ringtrials/>) provides a possibility to upload several isolates in a single submission.

Step 1; Login to the server using provided username (gmi\_xx) and password (pink area).

Step 2; Download the Excel Metadata template to your computer (green area).

Step 3; Fill in the required fields with all the relevant information (metadata) about the isolates. Note that **sample name** should be the **same as label-name of the sample**, e.g. **GMI17-003-BACT** or **GMI17-003-DNA**.

Step 4; Upload the metadata file as well as the individual WGS files to the web-interface by dropping the files to the 'Drop metadata and sequence files here' (grey area).



**Step 5;** Click on the Submit button to upload the files. It is important to keep the window opened until the upload is completed.

Via the Internet-based survey

([https://www.surveymonkey.com/r/PT\\_2017\\_bacterial\\_cultures\\_and\\_DNA](https://www.surveymonkey.com/r/PT_2017_bacterial_cultures_and_DNA); see also Appendix 2 in the full protocol document), answers should be submitted to the questions related to the analysed bacterial cultures and DNA.

### 3.1 Fastq dataset

The three fastq datasets should be downloaded from the ftp-site. They are organized into three different .zip archives appropriately labeled with the taxon they represent (st, ec, or sa). Within each archive the participant will find the paired-end reads.

The participant should perform variant detection and clustering (phylogenetics) of all files within each .zip archive according to participant's standard procedure.

For each .zip dataset the following should be uploaded to the ftp-site:

1. The DNA sequence matrix used for clustering should be in fasta format (.fasta file) and the clusters should be in newick format (.tre file)
  - The matrix and tree file should only contain **only** those samples provided through the ftp site (i.e., there should be only 14 st, 11 ec, and 11 sa samples in each file).
  - Syntax for the names of samples in each file should be **only** the prefix preceding the first underscore in the file name. For example, **st1\_1.fastq** should be named **st1** in the matrix and tree files.
  - The file should be named as follows **GMI LabID\_Taxon.fasta** (e.g., **GMI01\_st.FASTA**, **GMI01\_ec.FASTA**, **GMI01\_sa.FASTA**, **GMI01\_st.TRE**, **GMI01\_ec.TRE**, or **GMI01\_sa.TRE**).
2. The vcf (variant call format) files for each sample if a reference based approach was used and such files were produced.
  - The number of vcf files should match the number of samples found in the zipped archive from the ftp site.
  - Syntax for the names of the files should be **only** the prefix preceding the first underscore in the file name. For example, **st1\_1.fastq** should be named **st1** in the matrix.
  - The file should be named as follows **GMI LabID\_Taxon.tre** (e.g., **GMI01\_st1.vcf**, **GMI01\_ec1.vcf**, or **GMI01\_sa1.vcf**).

If performing a reference based approach for variant detection, the reference applied for the analysis **must be** st\_reference.fasta (*Salmonella*), ec\_reference.fasta (*E. coli*) and sa\_reference.fasta (*S. aureus*).

Via the Internet-based survey ( [https://www.surveymonkey.com/r/PT\\_2017\\_FASTQ\\_dataset](https://www.surveymonkey.com/r/PT_2017_FASTQ_dataset); see also Appendix 2 in the full protocol document), answers should be submitted to the questions related to the analysed of the fastq dataset.



#### 4 DISCUSSION FORUM

A web-based discussion forum (<https://foros.isciii.es/viewforum.php?f=7>) is available for participants in the GMI PT 2017. Appendix 4 in the full protocol document presents detailed information on the PT discussion forum.

#### 5 CONTACT INFO

**If you have any questions or concerns, please do not hesitate to contact us**, preferably by using the web-based discussion forum (<https://foros.isciii.es/viewforum.php?f=7>).

##### **PT organizer related to the dry-lab fastq datasets:**

James Pettengill  
U.S. Food and Drug Administration  
Center for Food Safety and Applied Nutrition  
CPK1 RM2D0195100 Paint Branch Parkway  
College Park, MD 20740, US  
Tel: +1 240-402-1992  
E-mail: [James.Pettengill@fda.hhs.gov](mailto:James.Pettengill@fda.hhs.gov)

##### **PT organizer in relation to other issues, e.g. organizational issues, please contact the GMI PT Coordinator:**

Susanne Karlsmosen Pedersen  
National Food Institute, Technical University of Denmark  
Kemitorvet, Building 204,  
DK-2800 Kgs. Lyngby, DENMARK  
Tel: +45 3588 6601  
E-mail: [suska@food.dtu.dk](mailto:suska@food.dtu.dk)

--- --- ---

##### **Note:**

This document describes the basic instructions for both the dry and wet lab components of the PT. See the full protocol document including appendices for more information:  
<http://www.globalmicrobialidentifier.org/workgroups/about-the-gmi-proficiency-test-2017>.