

SMaHT Sample & File Nomenclature

Resource for the SMaHT Network Members Version 1.1 Last modified: 2024-June-20

Description	
Overview	
Naming schema:	
Tissue	
Cell line	6
Additional sample/assay metadata in the name	7
Additional file information in the name	
Benchmarking Cell Line Names	11
File Name Examples	

Description

The SMaHT sample and file names are the primary identifiers of biosamples and files generated by the Tissue Procurement Center (TPC) and Data Analysis Center (DAC) of the SMaHT Network ("Network").

Overview

The SMaHT sample and file names contain identifiers that are unique and immovable, as well as semi-human readable codes that correspond to metadata. This document describes the naming schema and tables of codes for each metadata type that are included in sample and file names.

Naming schema:

[Project][Kit/Donor ID] - [Protocol ID] - [Tissue aliquot & Tissue core IDs] - [Sex & Age] - [Sequencing platform & Sequencing assay code] - [Center code] - [Unique Accession ID] - [Analysis info] . [File format]

Tissue

The donor, tissue, and sample-related metadata and identifiers are generated at TPC/NDRI and are used to name samples that are sent to GCCs and TTDs. Donor/tissue metadata fields are delimited by a hyphen ("-"). "#" indicates a number, and "A" indicates an alphabet letter.

For benchmarking tissues, the project is "ST".

For production tissues, the project is "SMHT".

Benchmarking tissues: ST### - #A - ###A#Production tissues: SMHT### - #A - ###A#

Note:

- The majority of benchmarking tissues will be homogenized. At specific requests by certain GCCs, non-homogenized benchmarking tissues have been made available for sequencing.
- 2. All production tissues will *not* be homogenized.
- 3. For non-homogenized tissues (for both benchmarking or production), there will be tissue aliquots from each tissue, which then get sectioned into "tissue cores" or "tissue samples", which are larger than cores and smaller than tissue aliquots.
- 4. GCCs do not need to rename their samples received from TPC when submitting the metadata to DAC. They need to provide DAC with the metadata they received from TPC and any information related to sample processing performed at GCC, e.g. aliquot numbers for benchmark homogenate tissues; which aliquot(s) were used to create a library.
- Project: ST for a benchmarking tissue sample; SMHT for a production tissue sample.
- Kit ID: Provided by TPC/NDRI. It is a three-digit pre-generated kit identifier that
 references the donor. 001–150 for production tissues; 001–004 for benchmarking
 tissues.
- Protocol ID: Automatically generated by NDRI's Rhythm system. A single internal
 protocol group number is followed by a letter designation (A-ZZ) corresponding to a
 specific combination of tissue type and tissue preservation method (Table 1).
- Tissue aliquot & core IDs: Provided by TPC/NDRI. Aliquot ID refers to spatially numbered tissue aliquots from longitudinally bisected or non-bisected tissue samples. Serial tissue cores subsampled from within an aliquot are designated by a letter followed by a number (Figures 1 and 2). No aliquots or cores will be available for buccal swab samples. No cores will be available for isolated fibroblasts and benchmarking tissue homogenate samples. Null values are represented as "x".
 - o For the homogenate benchmarking tissue (no core or tissue sample):
 - [Metadata from GCC] Tissue aliquot number will be provided by TPC, e.g., ST### - #A - ###X

- [File names at DAC] Tissue aliquot and Core ID are both "x", e.g.,
 ST### #A XX
 - To avoid having multiple homogenate aliquots associated with a file name.
- For the non-homogenate benchmarking tissue (in either core or tissue sample form):
 - [All] ST### #A ###A#
- Core ID for the core: ID is comprised of a letter between A-F, followed by a digit between 1-6 (ID is associated with the spatial information of the grid, representing a series of cores taken from a tissue aliquot See Figure 1).
- Core ID for the tissue specimen: ID is comprised of a letter between S-W followed by 1-9 (no spatial information associated with the ID).
- Core ID for the homogenate benchmarking tissues: Null value for the ID, i.e., "x".

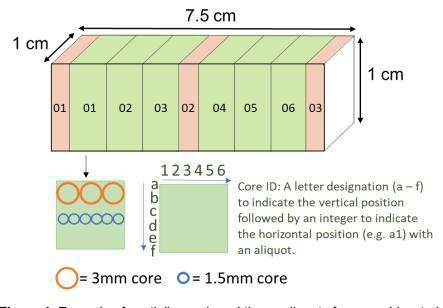


Figure 1. Example of spatially numbered tissue aliquots from non-bisected tissue samples. Fixed (pink) or frozen aliquots (green) are numbered separately. "X" represents null values.

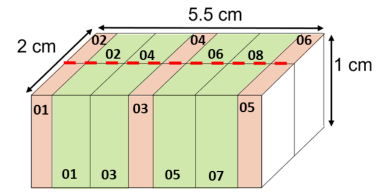


Figure 2. Example of spatially numbered tissue aliquots from longitudinally bisected tissue samples. Fixed (pink) or frozen (green) aliquots are numbered separately, in sequential order starting from the superior end of the sample.

Table 1. Protocol IDs for SMaHT benchmarking tissues

Sample Protocol ID	Tissue Name	Preservation	Notes
1A	Liver	Snap Frozen	Homogenates and non-homogenate samples
1C	Liver	Fixed	
1D	Lung	Snap Frozen	Homogenates and non-homogenate samples
1F	Lung	Fixed	
1G	Colon	Snap Frozen	Homogenates and non-homogenate samples
11	Colon	Fixed	
1J	Skin	Snap Frozen	Collected but not utilized for study; homogenization attempted but not successful.
1K	Skin	Snap Frozen	Cores made (intact tissue)
1L	Skin	Fixed	
1Q	Brain, Frontal Lobe	Snap Frozen	Homogenates and non-homogenate samples

Table 2. Protocol IDs for SMaHT production tissues.

Protocol ID	Tissue Name for Container	Preservation
3A	Blood, Whole	Frozen
3B	Buccal Swab	Fresh
3C	Esophagus	Frozen
3D	Esophagus	Fixed
3E	Colon, Ascending	Frozen

3F	Colon, Ascending	Fixed
3G	Colon, Descending	Frozen
ЗН	Colon, Descending	Fixed
31	Liver Sample	Frozen
3J	Liver Sample	Fixed
3K	Adrenal Gland, Left	Frozen
3L	Adrenal Gland, Left	Fixed
ЗМ	Adrenal Gland, Right	Frozen
3N	Adrenal Gland, Right	Fixed
30	Aorta, Abdominal	Frozen
3P	Aorta, Abdominal	Fixed
3Q	Lung	Frozen
3R	Lung	Fixed
3S	Heart, LV	Frozen
3T	Heart, LV	Fixed
3U	Testis, Left	Frozen
3V	Testis, Left	Fixed
3W	Testis, Right	Frozen
3X	Testis, Right	Fixed
3Y	Ovary, Left	Frozen
3Z	Ovary, Left	Fixed
3AA	Ovary, Right	Frozen
3AB	Ovary, Right	Fixed
3AC*	Skin, Calf	Fresh
3AD	Skin, Calf	Frozen
3AE	Skin, Calf	Fixed
3AF	Skin, Abdomen	Frozen
3AG	Skin, Abdomen	Fixed
ЗАН	Muscle	Frozen
		-

3AI	Muscle	Fixed
3AJ**	Brain	Fresh

^{*} Will be used to isolate fibroblasts. Could potentially still use "3AC" protocol for fibroblasts, but will need discussion.

Cell line

For benchmarking samples that are not distributed by TPC (e.g. UW, Yale), Kit/Sample ID is designated by DAC as shown in Table 2. As the TPC-based protocol and tissue aliquots do not apply to cell line cultures, the schema for the benchmarking cell lines is the following: ("A" indicates an alphabet letter, and "x" indicates a null value.

For benchmarking cell line (Table 2): SMHTAAAA - X - X

Table 2. Benchmarking cell line codes.

Kit/Sample ID	Cell line description	
COLO829T	COLO829 tumor cell line	
COLO829BL	COLO829BL normal lymphoblast cell line	
COLO829BLT50	COLO829 1:50 admixture	
НАРМАР6	Cell admixture of six HapMap cell lines	
HG002	HG002 HapMap cell line	
HG005	HG005 HapMap cell line	
HG02257	HG02257 HapMap cell line	
HG02486	HG02486 HapMap cell line	
HG02622	HG02622 HapMap cell line	
HG00438	HG00438 HapMap cell line	
LBLA2	LB-LA2 fibroblast cell line	
LBIPSC1	iPSC line from clone #1 derived from the LB-LA2 fibroblast cell line	
LBIPSC2	iPSC line from clone #2 derived from the LB-LA2 fibroblast cell line	
LBIPSC4	iPSC line from clone #4 derived from the LB-LA2 fibroblast cell line	
LBIPSC52	iPSC line from clone #52 derived from the LB-LA2 fibroblast cell line	

^{**} Will be frozen and subsectioned by NBB-Maryland. Will need subsection IDs from NBB-Maryland.

PSC line from clone #60 derived from the LB-LA2 fibroblast cell line
--

Additional sample/assay metadata in the name

Additional sample- and assay-related metadata are delimited by a hyphen ("-") as shown below.

[Project][Kit/Donor ID] - [Protocol ID] - [Tissue aliquot & Tissue core IDs] - [Sex & Age] - [Sequencing platform & Sequencing assay code] - [Center code] - [Unique Accession ID] - [Analysis info] . [File format]

- Sex & Age: Sex and age of the donor "M" (for male) or "F" (for female), followed by the donor's age in years. "N" and "N" will be used when sex or age is unknown or not applicable (e.g. cell line mixture).
- Sequencing platform & assay code: A code consisting of a single alphabet corresponding to a sequencing platform, followed by a 3-digit number designated for a specific experimental assay type (Tables 3A and 3B).
- Center code: Letters corresponding to the centers, i.e. GCCs and TTDs that generate original/raw sequencing data and submit to DAC (Table 4).

Table 3A. Sequencing platform codes.

SMaHT code	Sequencing platform
А	Illumina NovaSeq X
В	PacBio Revio HiFi
С	Illumina NovaSeq 6000
D	ONT PromethION 24
E	ONT PromethION 2 Solo
F	ONT MinION Mk1B
G	Illumina HiSeq X
Н	Illumina NovaSeq X Plus
(set the codes as data are generated on different sequencing platforms and submitted to DAC)	ONT MinION, Ultima Genomics, PacBio Onso, Element Aviti, PacBio Sequel

Table 3B. Experimental assay codes.

Code	DNA/RNA/Others	Assay Level	Assay Type

000	Null or N/A			
	DNA below			
001	DNA, PCR-free	Bulk	Whole genome sequencing (WGS)	
002	DNA	Bulk	WGS	
003	DNA	Bulk	Ultra-Long HMW WGS	
004	DNA	Bulk	Fiber-seq	
005	DNA	Bulk	Hi-C	
006	DNA	Bulk	NTSeq	
007	DNA	Single-molecule	CODEC	
800	DNA	Single-molecule	Duplex sequencing	
009	DNA	Single-molecule	NanoSeq	
010	DNA	Single-molecule, single-cell	scNanoSeq	
011	DNA	Single-cell	DLP+	
012	DNA	Single-cell	MALBAC-amplified WGS	
013	DNA	Single-cell	PTA-amplified WGS	
014	DNA	Single-cell	scDip-C	
015	DNA	Single-molecule	CompDuplex-seq	
016	DNA	Single-molecule, single-cell	scCompDuplex-seq	
			(Include additional DNA assays)	
	RNA below			
101	RNA	Bulk	RNA-seq	
102	RNA	Bulk	MAS-ISO-Seq	
103	RNA	Single-cell/nucleus	snRNA-Seq	
104	RNA	Single-cell/nucleus	STORM-Seq	
105	RNA	Single-cell/nucleus	Tranquil-Seq	
			(Include additional RNA assays)	
	Epigenetics below			
201			(Include additional epigenetic assays)	

	Others below		
301			(Include additional assays)

Table 4. SMaHT data generation center codes.

Code	Category	Institute	Contact PI
bcm	GCC	Baylor College of Medicine	Richard Gibbs
broad	GCC	Broad Institute	Kristin Ardlie
nygc	GCC	New York Genome Center	Soren Germer
uwsc	GCC	University of Washington & Seattle Children's Hospital	Jimmy Bennett
washu	GCC	Washington University in St. Louis	Ting Wang

Code	Category	Institute	Contact PI
bcm1	TTD	Baylor College of Medicine	Chuck Zong
bcm2	TTD	Baylor College of Medicine	Fritz Sedlazeck
bch1	TTD	Boston Children's Hospital	Christopher Walsh
bch2	TTD	Boston Children's Hospital	Sangita Choudhury
broad1	TTD	Broad Institute	Fei Chen
cwru	TTD	Case Western Reserve University	Fulai Jin
dfci	TTD	Dana-Farber Cancer Institute	Kathleen Burns
mayo	TTD	Mayo Clinic	Alexej Arbyzov
nyu	TTD	New York University	Gilad Evrony
stfd	TTD	Stanford University	Alexander Urban
umass	TTD	University of Massachusetts	Thomas Fazzio
umich	TTD	University of Michigan	Ryan Mills
uutah	TTD	University of Utah	Gabor Marth
wcnygc	TTD	Weill Cornell Medicine & New York Genome Center	Dan Landau

Code	Category	Institute	Contact PI
------	----------	-----------	------------

dac	DAC	Harvard Medical School	Peter Park
Code	Category	Institute	Contact PI
tpc	TPC	National Disease Research Interchange (NDRI)	Thomas Bell

Additional file information in the name

The file and analysis-related metadata and identifiers are generated at DAC.

[Project][Kit/Donor ID] - [Protocol ID] - [Tissue aliquot & Tissue core IDs] - [Sex & Age] -[Sequencing platform & Sequencing assay code] - [Center code] - [Unique Accession ID] -[Analysis info] . [File format]

- Unique Accession ID: Automatically generated unique accession ID with the "SM" prefix (for SMaHT) for a file stored at DAC. This ID can also be used as an identifier to search the file via the SMaHT data portal.
- Analysis info and File format: Information about the analysis used to generate the file can be added here using the following format, delimited by an underscore (" ").
 - For unaligned raw sequence files, a null value, "x", is set for the analysis info field.
 - [Tool name and version; separated by " "] [Genome version] [Variant type / expression] _ [Other info]

```
e.g. Unaligned FASTQ
```

...-SM192K2-X.fastq.qz

e.g. Unaligned BAM

 \dots -SM192K2-X.bam

e.g. BAM aligned against GRCh38 using Sentieon BWA-MEM

...-SMRO2AR-sentieon bwamem 202308.01 GRCh38.aligned.sorted. bam

E.g. VCF of somatic calls using Strelka2 called against the GRCh38 reference genome

```
...-SM092E4-strelka2 4.14 GRCh38 snv.vcf
```

- Tool name and version (mandatory): Software/Computational tool name and version used to generate the file, e.g. bwamem 0.7.17.
- Genome version (optional): Reference genome version used for aligned sequence files, or variant type for VCF files (Table 4A).
- Variant type or gene expression (optional; Table 4B).
- Additional key info (optional) can be added, delimited by an underscore ("_").

Table 4. Genome version (A) and variant type (B) tables. **(A)**

Reference Genome	Code
GRCh38 without ALT contigs	GRCh38
T2T CHM13	CHM13
Donor-specific assembly	DSA

(B)

Variant Type	Code
Substitutions, short insertions and deletions	snv
Copy number variants	cnv
Structural variants including large insertions and deletions, duplications, inversions, translocations	sv
Mobile element insertions	mei

Benchmarking Cell Line Names

```
1. COLO829
```

```
SMHTCOLO829T - X - X - M45
```

2. COLO829BL

```
SMHTCOLO829BL - X - X - M45
```

3. COLO829 1:50 mixture

```
SMHTCOL0829BLT50 - X - X - M45
```

4. HapMap mixture

```
SMHTHAPMAP6 - X - X - NN
```

5. iPSC / fibroblast, e.g. LB-IPSC 1

SMHTLBIPSC1 - X - X - [metadata from the Vaccarino lab]

File Name Examples

(space was added to make the delimiters (i.e. "-" and "_") more visible)

[Project][Kit/Donor ID] - [Protocol ID] - [Tissue aliquot & Tissue core IDs] - [Sex & Age] - [Sequencing platform & Sequencing assay code] - [Center code] - [Unique Accession ID] - [Analysis info] . [File format]

Bulk WGS on Illumina NovaSeqX - FASTQ file of the COLO829T-COLO829BL 1:50 admixture from BCM-GCC:

```
SMHTCOLO829BLT50 - X - X - M45 - A001 - bcm - SM123JK0 - X. fastq.gz
```

FiberSeq on PacBio HiFi - unaligned BAM file of the COLO829 tumor cell line from UW-GCC:

```
SMHTCOLO829T - X - X - M45 - B003 - uwsc - SM89QQ3LA - X . bam
```

FiberSeq on PacBio HiFi - BAM file of the COLO829 tumor cell line from UW-GCC, aligned to GRCh38, sorted by read name, and phased:

```
SMHTCOL0829T - X - X - M45 - B003 - uwsc - SM57S7SLA - pbmm2.6 GRCh38 . aligned.sorted.phased.bam
```

Bulk WGS on Ultima Genomics - SNVs called using Strelka2 from COLO829 1:50 admixture against COLO829BL:

Bulk WGS on Illumina NovaSeqX - FASTQ file of the HapMap mixture from Broad:

```
SMHTHAPMAP6 - X - X - NN - A001 - broad - SM89XAY38 - X. fastq.qz
```

• BAM file from the FASTQ file above:

```
SMHTHAPMAP6 - X - X - NN - A001 - broad - SM33A9K6R - sentieon bwamem 202308.01 GRCh38 . aligned.sorted.bam
```

 VCF that contains HapMap mixture-specific variants identified by running Mutect 2 on the BAM file above:

```
SMHTHAPMAP6 - X - X - NN - A001 - broad - SM3R123U9 - mutect4.1.4.1_GRCh38_snv . vcf
```

 In the current plan, for VCF files that are generated by GCCs and TTDs, and submitted to DAC, DAC will not rename the file names according to the SMaHT nomenclature.
 DAC will keep the original file names provided by the submitters, and the associated metadata will indicate who the submitting group is.