

STOMICS ANALYSIS WORKFLOW SOFTWARE SUITE USER MANUAL

Revision History

Manual Version: A0
Software Version: V1.0.0
Date: Nov. 2021
Description: Initial release

Manual Version: A1
Software Version: V2.1.0
Date: Dec. 2021

Description:

- New feature: addition of manual registration function, some fine-tuning will be automatically performed after manually registering;
- Improvement: performance improvement for mapping; the order of sequencing saturation calculation has switched, in v2.1.0 we only compute the saturation of tissue-covered region;
- Bug fix: fixed the bug of indexing at mapping step; fix the bug of long waiting time at register step.

Manual Version: A1.1
Software Version: V2.1.0
Date: Jan. 2022

Description:

- Add error handling;
- Update demo output.

Manual Version: A2
Software Version: V4.1.0
Date: Apr. 2022

Description:

- New feature: support to process fused micrographs; employ Stereopy in clustering; new gene expression matrix file format; including a file format convertor and a mapping memory estimator;
- Improvement: performance improvements for mapping; updated gene annotation approach in count; updated image stitching and tissue segmentation model for better performance on image stitching and segmentation for tissue with voids.

Note: Please download the latest version of the manual and use it with the corresponding software version.

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WORKFLOW

Minimum requirements:

Higher requirements:

64 bit CentOS/RedHat 7.8

1 G reads

64 bit Ubuntu 20.04 12 cores = 256 G 🖺 1 TB Reference **FASTQ** Mask Main Results mapping ~60 G **(1)**barcode mapping sequencing report (.summaryReport.html) 2 adapter filter bam index (.bam.bai) **3RNA** alignment statistical report (.stat &.Log.final.out) merge ○ ~5 min == ~5 G merge barcode reads count reads count (.barcodeReadsCount.txt) count **1**annotation annotated BAM (.bam) 2deduplication raw matrix (.raw.gef) 3get gef statistical report (.stat) image register ~20 min == ~20 G image registration image pyramid (.rpi) tissueCut (1) tissue cut statistical report (.stat) 2statistics matrix of tissue covered region (.tissue.gef) spatial cluster (.h5ad) spatialCluster saturation saturation plot (.png) ~15 min == ~30 G statistical report (.json) report web report (.report.html) ↑ ~1 min = ~1 G

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CHAPTER 1 STOMICS ANALYSIS WORKFLOW SOFTWARE SUITE

1.1. Software Introduction

STOmics Analysis Workflow¹ (SAW) software suite is a set of pipelines bundled to position sequenced reads to their spatial location on the tissue section, quantify spatial gene expression and visually present spatial expression distribution. SAW processes the sequencing data of Stereo-Seq² to generate spatial gene expression matrices, and then users could take these files as the starting point to perform downstream analysis. SAW includes eight essential pipelines:

- **mapping:** Corresponds *in situ* captured sequenced reads recorded in FASTQ^{3,4} files by Stereo-Seq with their spatial information. It also aligns reads to the reference genome and generates coordination sorted BAM files.
- **merge (optional):** Combines mapping of CID (same as barcodes) listed files with reads count from multiple runs of mapping. Only applicable for an analysis that requires to combine multiple pairs of FASTQ.
- **count:** Reads BAM files generated from mapping to perform gene annotation, de-duplication, and gene expression analysis on the aligned reads.
- **register (optional):** Align microscopic tissue staining image with gene expression matrix file (GEF) generated from count.
- **tissueCut:** Identify tissue coverage area on the chip and extract gene expression matrix of the corresponding spatial location by taking inputs from both count and register or count pipeline alone.
- **saturation:** Calculate sequencing saturation of tissue coverage area based on the file that used for sampling data generated from count.
- **spatialCluster:** Perform clustering analysis for spots (bin200) according to the gene expression matrix of the tissue covered area generated from tissueCut.
- **report:** Generate a JSON format statistical summary report that integrate the analysis result from each step, as well as an HTML web analysis report, showing spatial expression distribution of genes, key statistical metrices, sequencing saturation plots, and clustering analysis results.

1.2. System Requirements

SAW runs on Linux systems that meet the following minimum requirements:

8-core Intel or AMD processor (24 cores recommended)
128GB RAM (256GB recommended)
1TB free disk space
64-bit CentOS/RedHat 7.8 or Ubuntu 20.04

To install and run SAW, please install one kind of the following software:

docker⁵: version 20.10.8 or higher singularity⁶: version 3.8 or higher



1.3. SAW Docker Image Installation

SAW is delivered as a docker image that bundles all of its required software dependencies. You can pull the SAW docker image from Docker Hub to your local system and run analyses offline.

Please download the latest version of the software and use it with the corresponding **STOmics Analysis Workflow Softwate Suite User Manual** version.

Docker Hub link: https://hub.docker.com/r/stomics/saw/tags

We support using Singularity and Docker to install and run the SAW.

Here we take SAW version 4.1.0 as an example.



1.3.1 SAW Installation via Singularity

• CRITICAL STEPS: For red highlighted inputs, please replace with your own path or data.

```
Step 1: Pull the SAW docker image (2 options):
```

```
$ singularity build SAW_v4.1.0.sif docker://stomics/saw:04.1.0 ## option 1
$ singularity build --sandbox SAW_v4.1.0/ docker://stomics/saw:04.1.0 ## option 2
```

For non-root users, especially who don't have enough space in the /home/ directory, please try:

```
$ export SINGULARITY_CACHEDIR=/path/to/build
$ singularity build --sandbox SAW_v4.1.0/ docker://stomics/saw:04.1.0
```

Step 2: Run pipelines (3 options):

Note! All the requested paths need to be mounted before input. For example, it is necessary to bind directories that store input files (/path/to/data), reference genome (/path/to/genomeDir), and outputs (/path/to/output).

```
$ export SINGULARITY_BIND=" /path/to/data,/path/to/genomeDir,/path/to/output"
```

Option 1: Run pipelines within the container from the host system.

```
$ /path/to/SAW_v4.1.0.sif <application> ## option 1.1
$ singularity exec /path/to/SAW_v4.1.0.sif <application> ## option 1.2
```

Option 2: Shell into the SAW container and interactively run bash commands. Run exit to exit the environment.

```
$ /path/to/SAW_v4.1.0.sif /bin/bash ## option 2.1
Singularity>
Singularity> exit
$
$ singularity shell /path/to/SAW_v4.1.0.sif ## option 2.2
Singularity>
Singularity> <shell-command>
Singularity> exit
$
$ singularity shell SAW_v4.1.0 ## option 2.3 for sandbox
Singularity>
Singularity> singularity> exit
$
$ singularity> command>
Singularity> command>
Singularity> command>
Singularity> exit
$
```

Option 3: Use "-B directory on the host-machine:directory in the container" to mount a host directory into the container and execute the command in the container.

```
$ singularity shell -B /path/to/directory/on/the/host-machine:/path/to/directory/mounted/in/the/container/path/to/SAW_v4.1.0.sif
Singularity>
Singularity> <shell-command>
Singularity> exit
$
```

1.3.2 SAW Installation via Docker

```
Step 1: Pull the SAW docker image

to the state of the st
```

1.4 SAW Github

SAW GitHub: https://github.com/BGIResearch/SAW

Please visit GitHub for instruction regarding the **installation of singularity** and **indexing reference genome**. The page also provides **SAW shell script** examples for users.

• Note! Please build your reference before running SAW analysis.

1.5 SAW Test Data

SAW test data can be downloaded from SAW GitHub page. Key outputs are shown in <u>Chapter 3 Test Data</u> Demonstration. The reference genome version for SAW testing is:

genome-build: GRCm38.p6

genome-version: GRCm38

genome-date: 2012-01

genome-build-accession: NCBI:GCA_000001635.8

1.6 SAW Output File Format

Please check <u>STOmics File Format Manual</u> to get more information on SAW output files format.



^{*} Please be noted that these two lines belong to the same line of command.

CHAPTER 2 SAW PIPELINES & ARGUMENTS

2.1 mapping

Each Stereo-Seq sequenced read contains a CID sequence which is used as a key to spatially map the read back to its original location on the tissue slice. **mapping** pipeline matches CID of the original sequencing reads stored in the FASTQ file with the records of CID-coordinates key-value pairs saved in the STOmics Chip Mask file. Coordinate information for reads that CID could be paired with will be added based on the records of the Mask file. Reads that get the coordinate annotations are Valid CID mRNA Reads (Valid CID Reads). After filtering, the filtered Valid CID Reads are the Clean Reads. **mapping** pipeline maps Clean Reads to the reference genome, and output sorted BAM⁷ format alignments and summary report.

Run mapping requires the following files:

- Stereo-Seq sequenced reads FASTQ files (.fq.gz)
- STOmics Chip Mask file (.h5)
- Indexed reference genome
- bcPara file (.bcPara), please check the content of Table 2-2
- Expected running time for ~1G reads: ~3.5 h, Memory: ~60G
- NOTE! Before proceeding, if users need to estimate the number of CIDs and the memory needed for running mapping, please refer to 2.9.1 CIDcount for more information.

2.1.1 Arguments and Options

As mapping encapsulate STAR⁸ function, it accepts additional options beyond those shown in the table below.

Table 2-1 mapping Arguments and Options

Parameter	Function
outSAMattributes spatial	Set to turn on spatial BAM file format mode.
outSAMtype BAM SortedByCoordinate	(STAR option) Set output BAM file sorted by coordinate.
genomeDir	(STAR option) Path to the directory where the genome indices are stored.
runThreadN	(STAR option; defaults to 1) Set the number of threads to be used.
outFileNamePrefix	(STAR option) Custom output file prefix.
sysShell /bin/bash	(STAR option) Path to the shell binary.
stParaFile	(Required) Name of a parameters file defines CID mapping options. Options are specified in Table 2-2.
readNameSeparator \" \"	(STAR option) Character(s) separating the part of the read names that will be trimmed in output.
limitBAMsortRAM	(STAR option) Maximum available RAM (bytes) for sorting BAM.
limitOutSJcollapsed	(STAR option) Max number of collapsed junctions.
limitIObufferSize	(STAR option) Max available buffers size (bytes) for input/output, per thread.
outBAMsortingBinsN	(STAR option; defaults to 50) number of genome bins for coordinate-sorting.If the read2 FASTQ file size is greater than 200, it is better to set this value to 100.



Table 2-2 mapping --stParaFile Arguments and Options

Parameter	Function
in	(Required) Path to the STOmics Chip Mask file.
in1	(Required) Path to the FASTQ file. If PE sequencing, then specify the path to the FASTQ file of read1 here.
in2	(Optional) Path to the FASTQ file of read2. Only valid for PE sequencing.
encodeRule	(Required) Encoding rule for the four bases. ACTG stands for A->0, C->1, T->2, G->3.
out	(Optional) Set output file prefix.
action	(Required; defaults to 1) Action number. Valid options: 1=CID stat, 2=CID overlap, 3=get CID position map, 4=map CID to slide, or 5=merge CID list.
barcodeReadsCount	(Required) Mapped CID list file with reads counts for each CID.
platform	(Optional) Sequencing platform. Valid options: SEQ500, T1, or T10.
barcodeStart	(Required; defaults to 0) CID start position.
barcodeLen	(Required; defaults to 25) CID length.
umiStart	(Required; defaults to 25) MID start position.
umiLen	(Required; defaults to 10) MID length.
umiRead	(Required; defaults to 1) Declare the read contains MID.
mismatch	(Required; defaults to 0) Max mismatch tolerant.
bcNum	(Optional) CID count. Please check 2.9.1 CIDcount for more information.

2.1.2 Usage Example

Prepare mapping --stParaFile input file {lane}.bcPara as shown below:

- **○** The same applies to all examples.

```
$ mkdir /path/to/output/00.mapping
$ vim /path/to/output/00.mapping/{lane}.bcPara
in=/path/to/data/{SN}.barcodeToPos.h5
in1=/path/to/data/{lane}_read_1.fq.gz
in2=/path/to/data/{lane}_read_2.fq.gz
encodeRule=ACTG
out={lane}
barcodeReadsCount=/path/to/ouptut/00.mapping/{lane}.barcodeReadsCount.txt
action=4
platform=T10
barcodeStart=0
barcodeLen=25
umiStart=25
umiLen=10
umiRead=1
mismatch=1
```



Run mapping pipeline

Note! If a fatal error has occurred as shown below, you could delete --readNameSeparator \" \" from the command:

```
EXITING: FATAL INPUT ERROR: empty value for parameter "readNameSeparator" in input "Command-Line"

SOLUTION: use non-empty value for this parameter
```

2.1.3 Outputs

The output files of mapping are organized as below:

```
$ ll -Rth /path/to/output/00.mapping
-rw-rw-r-- 1 ubuntu ubuntu 11M Apr 13 20:57 lane.Aligned.sortedByCoord.out.bam.bai
-rw-rw-r-- 1 ubuntu ubuntu 8.9K Apr 13 20:40 lane.Log.out
-rw-rw-r-- 1 ubuntu ubuntu 2.0K Apr 13 20:40 lane.Log.final.out
-rw-rw-r-- 1 ubuntu ubuntu 17K Apr 13 20:40 lane.Log.progress.out
-rw-rw-r-- 1 ubuntu ubuntu 62G Apr 13 20:40 lane.Aligned.sortedByCoord.out.bam
-rw-rw-r-- 1 ubuntu ubuntu 1.1K Apr 13 20:06 lane_barcodeMap.stat
-rw-rw-r-- 1 ubuntu ubuntu 11M Apr 13 20:06 lane.SJ.out.tab
-rw-rw-r-- 1 ubuntu ubuntu 997M Apr 13 20:06 lane.barcodeReadsCount.txt
-rw-rw-r-- 1 ubuntu ubuntu 528 Apr 13 17:46 lane.bcPara
```

If one sample has multiple FASTQ files, you need to run **mapping** for each FASTQ pair. The output files are organized as below (Next page showing example of 2 pairs of FASTQ):

```
•••
$ tree /path/to/multi_lane_output/00.mapping
/path/to/multi_lane_output/00.mapping
└─ 00.mapping

    lane1.Aligned.sortedByCoord.out.bam

     lane1.Aligned.sortedByCoord.out.bam.bai
      lane1_barcodeMap.stat

    lane1.barcodeReadsCount.txt

      - lane1.bcPara
      lane1.Log.final.out
      lane1.Log.out
      lane1.Log.progress.out
      lane1.SJ.out.tab
      lane2.Aligned.sortedByCoord.out.bam

    lane2.Aligned.sortedByCoord.out.bam.bai

      lane2_barcodeMap.stat
```

```
lane2.barcodeReadsCount.txt
lane2.bcPara
lane2.Log.final.out
lane2.Log.out
lane2.Log.progress.out
lane2.SJ.out.tab
```

2.2 merge (optional)

SAW merge pipeline is used to combine the results of mapping. Run merge requires the following files:

- mapping output mapped CID with reads count files (.txt)
- Expected running time for ~1G reads 2 lanes: ~5 min, Memory: ~5G

2.2.1 Arguments and Options

Table 2-3 merge Arguments and Options

Parameter	Function
-i	(Required) Mapped CID list files with reads counts for each CID.
out	(Required) Mapped CID list file which merges all input files.
action	(Required; defaults to 1) Action number. Valid options: 1=map CID to slide, 2=merge CID list, 3=mask format change, or 4=mask merge.

2.2.2 Usage Example

2.2.3 Ouputs

The output file of merge has been organized as below:

```
$ ll -Rth /path/to/multi_lane_output/01.merge
-rw-rw-r-- 1 ubuntu ubuntu 997M Apr 13 20:53 SN.barcodeReadsCount.txt
```



^{*} Please be noted that we use the backward slash "\" to indicate the end of a line in a command that spans multiple lines.

2.3 count

SAW **count** pipeline annotates **Uniquely Mapped Reads** filtered from **mapping** output based on the reference genome annotation records. Through quantification of annotated reads, **count** generates spatial gene expression data after de-duplication reads according to CID, gene ID and MID information.

Run count requires the following files:

- mapping output BAM file (.bam)
- Reference genome annotation GFF/GTF^{9,10} file (.gtf / .gff)
- Expected running time for ~1G reads: 1.5h, Memory: ~50G

2.3.1 Arguments and Options

Table 2-4 count Arguments and Options

Parameter	Function
-i	(Required) mapping output BAM file. Separate multiple files by comma.
-o	(Required) Set the count output BAM file name.
-a	(Required) Gene annotation file.
-s	(Required) Set the count output statistical summary report file name.
-e	(Required) Set the count output gene expression file name.
umi_len	(Required; defaults to 10) MID length.
sn	(Required) STOmics Chip serial number (SN).
-c	(Optional) CPU core number to use.
save_lq	(Optional) Save low quality reads if set.
save_dup	(Optional) Save duplicate reads if set.
umi_on	(Optional) Correct MID if set.
sat_file	(Optional) Set the saturation sampling file name which is prepared for sequencing saturation (requiresumi_on).
-m	(Optional; defaults to detected) Set available memory (GB).

2.3.2 Usage Example

```
$ mkdir -p /path/to/output/02.count
 $ geneExp=/path/to/output/02.count/{SN}.raw.gef
 $ saturationSamplingFile=/path/to/output/02.count/raw_barcode_gene_exp.txt
 $ singularity exec SAW_v4.1.0.sif count \
         -i /path/to/output/00.mapping/{lane}.Aligned.sortedByCoord.out.bam \
         -o /path/to/output/02.count/{SN}.Aligned.sortedByCoord.out.merge.q10.dedup.
*target.bam \
         -a /path/to/genomeDir/genes.gtf \
         -s /path/to/output/02.count/{SN}.Aligned.sortedByCoord.out.merge.q10.dedup.
^{*}target.bam.summary.stat \
         -e ${geneExp} \
         --umi_len 10 \
         --sat_file ${saturationSamplingFile} \
         --sn {SN} \
         --umi_on \
         --save_lq \
         --save_dup \
         -c 8 \
         -m 128
```

^{*} Please be noted that we use the backward slash "\" to indicate the end of a line in a command that spans multiple lines.



For more than one pair of FASTQ files (Here showing an example of 2 pairs of FASTQ),

```
$ mkdir -p /path/to/multi_lane_output/02.count
 $ geneExp=/path/to/multi_lane_output/02.count/{SN}.raw.gef
 $ saturationSamplingFile=/path/to/multi_lane_output/02.count/raw_barcode_gene_exp.txt
 $ singularity exec SAW_v4.1.0.sif count \
         -i /path/to/multi_lane_output/00.mapping/{lane1}.Aligned.sortedByCoord.out.
{}^{lack}bam,/path/to/multi_lane_output/00.mapping/\{lane2\}.Aligned.sortedByCoord.out.bam \setminus
         -o /path/to/multi_lane_output/02.count/{SN}.Aligned.sortedByCoord.out.merge.
□q10.dedup.target.bam \
         -a /path/to/genomeDir/genes.gtf \
         -s /path/to/multi_lane_output/02.count/{SN}.Aligned.sortedByCoord.out.merge.
^{
m L}q10.dedup.target.bam.summary.stat \
         -e ${geneExp} \
         --umi_len 10 \
         --sat_file ${saturationSamplingFile} \
         --sn {SN} \
         --umi_on \
         --save_lq \
         --save_dup \
         -c 8 \
         -m 128
```

2.3.3 Ouputs

The **count** output files are organized as below:

```
$ ll -Rth /path/to/output/02.count
-rw-rw-r-- 1 ubuntu ubuntu 636M Apr 13 22:11 SN.raw.gef
-rw-rw-r-- 1 ubuntu ubuntu 2.6G Apr 13 22:11 SN_raw_barcode_gene_exp.txt
-rw-rw-r-- 1 ubuntu ubuntu 393 Apr 13 22:11 SN.Aligned.sortedByCoord.out.merge.
q10.dedup.target.bam.summary.stat
-rw-rw-r-- 1 ubuntu ubuntu 46G Apr 13 22:11 SN.Aligned.sortedByCoord.out.merge.
q10.dedup.target.bam
```

2.4 register (optional)

SAW **register** pipeline aligns the microscopic tissue staining image with the plot of the gene expression matrix generate by **count** based on the tracklines on the chip while establishing the mapping relationship between images and spatial expression.

Run register requires the following files:

- count output gene expression matrix file (.raw.gef)
- ImageQC processed microscopic tissue staining image file (.tar.gz)
- ImageQC image information report (.json)
- Expected running time for ~1G reads: ~20 min, Memory: ~20G

^{*} Please be noted that we use the backward slash "\" to indicate the end of a line in a command that spans multiple lines.

2.4.1 Arguments and Options

Table 2-5 register Arguments and Options

Parameter	Function
-i	(Required) ImageQC processed staining image TAR.GZ file.
-c	(Required) ImageQC JSON file.
-v	(Required) count output gene expression matrix GEF file.
-o	(Required) Path to the directory where to store the register outputs.

2.4.2 Usage Example

2.4.3 Outputs

register output files are organized as below:

```
$ ll -th /path/to/output/03.register
drwxrwxr-x 2 ubuntu ubuntu 4.0K Apr 13 22:15 7_result/
drwxrwxr-x 2 ubuntu ubuntu 4.0K Apr 13 22:15 6_analysis/
drwxrwxr-x 2 ubuntu ubuntu 4.0K Apr 13 22:15 3_vision/
drwxrwxr-x 2 ubuntu ubuntu 4.0K Apr 13 22:15 4_register/
drwxrwxr-x 2 ubuntu ubuntu 4.0K Apr 13 22:13 5_mask/
drwxrwxr-x 2 ubuntu ubuntu 4.0K Apr 13 22:12 2_stitch/
drwxrwxrwx 3 ubuntu ubuntu 4.0K Apr 13 22:11 1_origin/
$ ll -Rth /path/to/output/03.register/4_register
-rw-rw-r-- 1 ubuntu ubuntu 668M Apr 13 22:15 fov_stitched_regist.tif
-rw-rw-r-- 2 ubuntu ubuntu 2.1M Apr 13 22:14 transform_thumb.png
-rw-rw-r-- 2 ubuntu ubuntu 39 Apr 13 22:14 attrs.json
-rw-rw-r-- 1 ubuntu ubuntu 458M Apr 13 22:13 fov_stitched_transformed.tif
                           50 Apr 13 22:13 im_shape.txt
-rw-rw-r-- 1 ubuntu ubuntu
$ ll -Rth /path/to/output/03.register/5_mask
-rw-rw-r-- 1 ubuntu ubuntu 458M Apr 13 22:13 fov_stitched_transformed_tissue_cut.tif
$ ll -Rth /path/to/output/03.register/7_result
-rw-rw-r-- 1 ubuntu ubuntu 321M Apr 13 22:15 merge_SN.tif
-rw-rw-r-- 1 ubuntu ubuntu 176M Apr 13 22:15 SN.rpi
-rw-rw-r-- 1 ubuntu ubuntu 4.9K Apr 13 22:15 SN_20211125.json
-rw-rw-r-- 1 ubuntu ubuntu 38 Apr 13 22:15 SN_tissue_bbox.csv
-rw-rw-r-- 1 ubuntu ubuntu 668M Apr 13 22:15 SN_tissue_cut.tif
-rw-rw-r-- 1 ubuntu ubuntu 668M Apr 13 22:15 SN_regist.tif
```

2.5 tissueCut

SAW tissueCut pipeline could delineate and extract the tissue coverage area based on the aligned image generated from register or from the plot of gene expression matrix (if microscopic tissue staining images are not available). tissueCut outputs expression data in GEF format. Users may generate registered image in TIFF or JPG format from image pyramid RPI file using python package Stereopy ¹¹.

If the output of tissueCut doesn't match the morphology of the tissue, user could use Stereopy to do lasso selection interactively to extract the expression matrix of tissue-covered region. Please check the tutorial, Stereopy->Examples->Interactive.

Run tissueCut requires the following files:

- Mapped CID with reads count file (.txt)
- count output gene expression matrix file (.raw.gef)
- Directory stores aligned microscopic staining image (optional)
- Expected running time for ~1G reads: ~1h, Memory: ~35G

2.5.1 Arguments and Options

Table 2-6 tissueCut Arguments and Options

Parameter	Function
dnbfile	(Required) Mapped CID list file with reads counts for each CID. Use the merged mapped CID with reads count file if multiple pairs of FASTQ files are involved in the analysis.
-i	(Required) count output gene expression matrix GEF file.
-o	(Required) Path to the directory where to store the tissueCut outputs.
-s	(Optional) Path to the directory where the register outputs are stored. Only valid when register has performed.
-t	(Required) Run for tissue segmentation. Valid options: tissue.
snId	(Required) STOmics Chip serial number (SN).
platform	(Required, default to T1) Sequencing platform.

2.5.2 Usage Example

Run tissueCut if register aligned microscopic staining image is provided:

Run tissueCut if image is not available:

2.5.3 Outputs

tissueCut output files:

Image is provided:

```
$ ll -th /path/to/output/04.tissuecut
-rw-rw-r-- 1 ubuntu ubuntu 1.3K Apr 13 22:39 tissuecut.stat
drwxrwxr-x 2 ubuntu ubuntu 4.0K Apr 13 22:39 tissue_fig/
-rw-rw-r-- 1 ubuntu ubuntu 526M Apr 13 22:19 SN.tissue.gef
drwxrwxr-x 2 ubuntu ubuntu 4.0K Apr 13 22:17 dnb_merge/
-rw-rw-r-- 2 ubuntu ubuntu 5.6G Apr 13 22:17 SN.gef
$ ll -Rth /path/to/output/04.tissuecut/tissue_fig
-rw-rw-r-- 1 ubuntu ubuntu 136K Apr 13 22:39 violin_200x200_MID_gene.png
-rw-rw-r-- 1 ubuntu ubuntu 27K Apr 13 22:39 scatter_200x200_MID_gene_counts.png
-rw-rw-r-- 1 ubuntu ubuntu 185K Apr 13 22:36 violin_150x150_MID_gene.png
-rw-rw-r-- 1 ubuntu ubuntu 32K Apr 13 22:36 scatter_150x150_MID_gene_counts.png
-rw-rw-r-- 1 ubuntu ubuntu 295K Apr 13 22:32 violin_100x100_MID_gene.png
-rw-rw-r-- 1 ubuntu ubuntu 29K Apr 13 22:32 scatter_100x100_MID_gene_counts.png
-rw-rw-r-- 1 ubuntu ubuntu 514K Apr 13 22:28 violin_50x50_MID_gene.png
-rw-rw-r-- 1 ubuntu ubuntu 29K Apr 13 22:28 scatter_50x50_MID_gene_counts.png
-rw-rw-r-- 2 ubuntu ubuntu 176M Apr 13 22:17 SN.ssDNA.rpi
```

Image is not provided:

```
$ tree /path/to/output/04.tissuecut
/path/to/output/04.tissuecut
— dnb_merge
   ├─ bin200.png
  - SN.gef
 SN.tissue.gef
 tissue_fig
    — scatter_100x100_MID_gene_counts.png
      scatter_100x100_MID_gene_counts.png
     scatter_150x150_MID_gene_counts.png
     scatter_200x200_MID_gene_counts.png
     scatter_50x50_MID_gene_counts.png
     — violin_100x100_MID_gene.png
     — violin_150x150_MID_gene.png
      violin_200x200_MID_gene.png
     violin_50x50_MID_gene.png
   tissuecut.stat
```

2.6 spatialCluster

SAW **spatialCluster** pipeline perform clustering analysis for spots at bin200 (bin size of 200) using Leiden algorithm.

Run **spatialCluster** requires the following files:

- tissueCut output GEF file for the tissue-covered region (.tissue.gef)
- Expected running time for ~1G reads: ~5 min, Memory: ~5G



2.6.1 Arguments and Options

Table 2-7 spatialCluster Arguments and Options

Parameter	Function
-i	(Required) tissuCut output GEF file for the tissue coverage area.
-o	(Required) Output path for the clustering result in H5AD format.
-s	(Required) Bin size, recommend to use 200.

2.6.2 Usage Example

```
$ mkdir -p /path/to/output/05.spatialcluster
$ singularity exec SAW_v4.1.0.sif spatialCluster \
    -i /path/to/output/04.tissuecut/{SN}.tissue.gef \
    -o /path/to/output/05.spatialcluster/{SN}.spatial.cluster.h5ad \
    -s 200
```

2.6.3 Outputs

spatialCluster output files are:

```
$ ll -Rht /path/to/output/05.spatialcluster
-rw-rw-r-- 2 ubuntu ubuntu 231M Apr 13 22:40 SN.spatial.cluster.h5ad
```

2.7 Saturation

SAW **saturation** pipeline is performed to compute the sequencing **saturation** for the tissue coverage area.

Run **saturation** requires the following files:

- count output saturation sampling file (.txt)
- tissueCut output GEF file for the tissue coverage area (.tissue.gef)
- mapping output statistical report of CID mapping (.stat)
- count output statistical report of annotation (.stat)
- Expected running time for ~1G reads: ~15 min, Memory: ~30G

2.7.1 Arguments and Options

Table 2-8 saturation Arguments and Options

Parameter	Function
-i	(Required) count output saturation sampling file.
tissue	(Required) tissueCut output GEF file for the tissue coverage area.
-о	(Required) Path to the directory where to store the saturation outputs.
bcstat	(Required) mapping output statistical report of CID mapping. Separate multiple files by comma.
summary	(Required) count output statistical report of annotation.



2.7.2 Usage Example

```
$ mkdir -p /path/to/output/06.saturation
$ singularity exec SAW_v4.1.0.sif saturation \
    -i /path/to/output/02.count/raw_barcode_gene_exp.txt \
    --tissue /path/to/output/04.tissuecut/{SN}.tissue.gef \
    -o /path/to/output/06.saturation \
    --bcstat /path/to/output/00.mapping/{lane}_barcodeMap.stat \
    --summary /path/to/output/02.count/{SN}.Aligned.sortedByCoord.out.merge.q10.

* dedup.target.bam.summary.stat
```

For more than one pair of FASTQ files (Here showing an example of 2 pairs of FASTQ),

2.7.3 Outputs

saturation output files are:

```
$$ ll -Rht /path/to/output/06.saturation
-rw-rw-r-- 1 ubuntu ubuntu 77K Apr 13 22:54 plot_200x200_saturation.png
-rw-rw-r-- 1 ubuntu ubuntu 36K Apr 13 22:54 plot_1x1_saturation.png
-rw-rw-r-- 1 ubuntu ubuntu 780 Apr 13 22:53 sequence_saturation.txt
```

2.8 report

SAW **report** pipeline is performed to integrate analysis report from each step and generate the report in JSON format as well as a web report in HTML format. HTML analytical report integrate genes' spatial expression distribution, key statistical metrices, sequencing saturation plots, and clustering analysis result plots.

Run **report** requires the following files:

- mapping output statistical reports of CID mapping (.stat) and STAR alignment (.out)
- count output statistical report of annotation (.stat)
- **tissueCut** output GEF file (.tissue.gef), statistical report of tissue-covered region (.stat), plots (.png) and image pyramid RPI file (.rpi)
- spatialCluster output clustering file (.h5ad)
- saturation output bin200 sequence saturation plot (.png)
- Expected running time for ~1G reads: ~1 min, Memory: 1G



^{*} Please be noted that these two lines belong to the same line of command.

^{*} Please be noted that we use the backward slash "\" to indicate the end of a line in a command that spans multiple lines.

2.8.1 Arguments and Options

Table 2-9 report Arguments and Options

Parameter	Function
-т	(Required) Statistical report of CID mapping. Separate multiple files by comma.
-a	(Required) Statistical report of STAR alignment. Separate multiple files by comma.
-g	(Required) Statistical report of annotation.
-1	(Required) Statistical report of tissue-covered region.
-n	(Required) tissueCutoutputGEF file.
-b	(Required) tissueCut output bin 200 scatter plot.
-v	(Required) tissueCut output bin 200 violin plots.
-i	(Optional) The image pyramid RPI file.
-d	(Required) spatialCluster output H5AD file.
-t	(Required) saturation output bin 200 sequence saturation plot.
-r standard_version	(Required) Set to specifying report version.
-s	(Required) The STOmics Chip serial number.
pipelineVersion	(Required) Set to specifying analysis pipeline version.
-0	(Required) The directory to store outputs.

2.8.2 Usage Example

Run report if register aligned microscopic staining image is provided:

```
$ mkdir -p /path/to/output/07.report
 $ singularity exec SAW_v4.1.0.sif report \
     -m /path/to/output/00.mapping/{lane}_barcodeMap.stat \
     -a /path/to/output/00.mapping/{lane}.Log.final.out \
     -g /path/to/output/02.count/{SN}.Aligned.sortedByCoord.out.merge.q10.dedup.
target.bam.summary.stat \
     -l /path/to/output/04.tissuecut/tissuecut.stat \
     -n /path/to/output/04.tissuecut/{SN}.gef \
     -d /path/to/output/05.spatialcluster/{SN}.spatial.cluster.h5ad \
     -t /path/to/output/06.saturation/plot_200x200_saturation.png \
     -b /path/to/output/04.tissuecut/tissue_fig/scatter_200x200_MID_gene_counts.png \
     -v /path/to/output/04.tissuecut/tissue_fig/violin_200x200_MID_gene.png \
     -r standard_version \
     -i /path/to/output/04.tissuecut/tissue_fig/{SN}.ssDNA.rpi \
     -s {SN} \
     --pipelineVersion SAW_v4.1.0 \
     -o /path/to/output/07.report
```



^{*} Please be noted that we use the backward slash "\" to indicate the end of a line in a command that spans multiple lines.

For more than one pair of FASTQ files (Here showing an example of 2 pairs of FASTQ),

```
$ mkdir -p /path/to/multi_lane_output/07.report
 $ singularity exec SAW_v4.1.0.sif report \
     -m /path/to/multi_lane_output/00.mapping/{lane1}_barcodeMap.stat,/path/to/multi_
tane_output/00.mapping/{lane2}_barcodeMap.stat \
     -a /path/to/multi_lane_output/00.mapping/{lane1}.Log.final.out,/path/to/multi_
lane_output/00.mapping/{lane2}.Log.final.out \
     -g /path/to/multi_lane_output/02.count/{SN}.Aligned.sortedByCoord.out.merge.q10.
* dedup.target.bam.summary.stat \
     -l /path/to/multi_lane_output/04.tissuecut/tissuecut.stat \
     -n /path/to/multi_lane_output/04.tissuecut/{SN}.gef \
     -d /path/to/multi_lane_output/05.spatialcluster/{SN}.spatial.cluster.h5ad \
     -t /path/to/multi_lane_output/06.saturation/plot_200x200_saturation.png \
     -b /path/to/multi_lane_output/04.tissuecut/tissue_fig/scatter_200x200_MID_gene_
*counts.png \
     -v /path/to/multi_lane_output/04.tissuecut/tissue_fig/violin_200x200_MID_gene.png
     -r standard_version \
     -i /path/to/multi_lane_output/04.tissuecut/tissue_fig/{SN}.ssDNA.rpi \
     -s {SN} \
     --pipelineVersion SAW_v4.1.0 \
     -o /path/to/multi_lane_output/07.report
```

Run **report** if **register** aligned microscopic staining image is not provided (Here showing an example of just one pair of FASTQ, similar to multiple pairs),

```
$ mkdir -p /path/to/output/07.report
 $ singularity exec SAW_v4.1.0.sif report \
     -m /path/to/output/00.mapping/{lane}_barcodeMap.stat \
     -a /path/to/output/00.mapping/{lane}.Log.final.out \
    -g /path/to/output/02.count/{SN}.Aligned.sortedByCoord.out.merge.q10.dedup.
*target.bam.summary.stat \
     -l /path/to/output/04.tissuecut/tissuecut.stat \
     -n /path/to/output/04.tissuecut/{SN}.gef \
     -d /path/to/output/05.spatialcluster/{SN}.spatial.cluster.h5ad \
     -t /path/to/output/06.saturation/plot_200x200_saturation.png \
     -b /path/to/output/04.tissuecut/tissue_fig/scatter_200x200_MID_gene_counts.png \
     -v /path/to/output/04.tissuecut/tissue_fig/violin_200x200_MID_gene.png \
    -r standard_version \
     -s {SN} \
     --pipelineVersion SAW_v4.1.0 \
     -o /path/to/output/07.report
```

2.8.3 Outputs

report output files are organized as below:

```
$ ll -Rth /path/to/output/07.report
-rw-rw-r-- 2 ubuntu ubuntu 1.2M Apr 13 22:54 SN.report.html
-rw-rw-r-- 2 ubuntu ubuntu 5.8K Apr 13 22:54 new_final_result.json
```

^{*} Please be noted that we use the backward slash "\" to indicate the end of a line in a command that spans multiple lines.

^{*} Please be noted that we use the backward slash "\" to indicate the end of a line in a command that spans multiple lines.

2.9 Other Applications

2.9.1 CIDcount

CIDcount is a small program for computing the number of CIDs in the STOmics Chip Mask file and roughly estimating how much memory will be needed to do **mapping**.

```
$ singularity exec SAW_v4.1.0.sif CIDcount \
    -i /path/to/data/{SN}.barcodeToPos.h5 \ ## STOmics Chip Mask file path
    -s {speciesName} \ ## a string of species name
    -g {genomeSize} ## Genome file size in GB, can be acquired by "ls -l --block-size=GB ${Genome file of the species after STAR indexing}"
```

The output of **CIDcount**is shown as below,

```
$ singularity exec SAW_v4.1.0.sif CIDcount -i SN.barcodeToPos.h5 -s mouse -g 3 645784920 ## CID count 62 ## estimated memory for mapping
```

If users wich to run **CIDcount**, they may add "bcNum" to the required {lane}.bcPara file for mapping as shown below,

```
$ vim /path/to/output/00.mapping/{lane}.bcPara
in=/path/to/data/{SN}.barcodeToPos.h5
in1=/path/to/data/{lane}_read_1.fq.gz
in2=/path/to/data/{lane}_read_2.fq.gz
encodeRule=ACTG
out={lane}
barcodeReadsCount=/path/to/ouptut/00.mapping/{lane}.barcodeReadsCount.txt
action=4
platform=T10
barcodeStart=0
barcodeLen=25
umiStart=25
umiLen=10
umiRead=1
mismatch=1
bcNum=645784920 ## first line from output of CIDcount
```

2.9.2 gefTools

gefTools¹² is an application for manipulating GEF file. SAW contains this tool to convert GEF format gene expression matrix to plain table or complete a GEF. Users may also manipulate the GEF files using its python encapsulated package **gefpy**¹³.

Function 1: GEF to plain table GEM format

```
$ singularity exec SAW_v4.1.0.sif gefTools view \ ## convert GEF that only contains
bin1 geneExp
    -i /path/to/output/02.count/{SN}.raw.gef \
    -o {SN}.raw.gem
$ singularity exec SAW_v4.1.0.sif gefTools view \ ## convert a whole GEF
    -i /path/to/output/04.tissuecut/{SN}.gef \
    -o {SN}.gem
$ singularity exec SAW_v4.1.0.sif gefTools view \ ## convert tissue GEF that only
contains bin1 geneExp
    -i /path/to/output/04.tissuecut/{SN}.tissue.gef \
    -o {SN}.tissue.gem
```



Function 2: completion of a GEF

Function 3: converting GEM to GEF

```
$ singularity exec SAW_v4.1.0.sif gefTools bgef \ ## convert GEM to GEF in specific
bin size. Separate multiple bin sizes with comma
    -i {SN}.gem \
    -o {SN}.gef \
    -b 1,20,50
```

Example of GEF to GEM conversion using gefpy, users may specify the bin size.

```
$ python
>>> from gefpy.bgef_reader_cy import BgefR
>>> bgef=BgefR(filepath=' /path/to/output/04.tissue/{SN}.tissue.gef' ,bin_size=200,n_
thread=4)
>>> bgef.to_gem( '{SN}.tissue.bin200.gem' )
```

CHAPTER 3 TEST DATA DEMONSTRATION

Users may refer to this section as a format for testing SAW process. This chapter includes the statistical results and examples of critical files for each key step.

SN: SS200000135TL_D1

..." in the demo stands for some lines of log information that can be omitted.

3.1 mapping

3.1.1 Statistical Report for CID Mapping and Filtering

```
$ cat /path/to/output/00.mapping/E100026571_L01_trim_read_barcodeMap.stat
getBarcodePositionMap_uniqBarcodeTypes: 645784920
total_reads: 1002214171
fixed_sequence_contianing_reads:
                                                                 0.00%
pass_filter_reads: 1002214171
mapped_reads: 845170516 84.33%
reads_with_adapter: 8137401 0.81%
reads_with_dnb: 45284608 4.52%
barcode_exactlyOverlap_reads: 698287595
                                                                69.67%
barcode_misOverlap_reads:
barcode_withN_reads: 0
                                          146882921
                                                                14.66%
                                           0.00%
Q10_bases_in_barcode: 99.54%
Q20_bases_in_barcode: 97.49%
Q30_bases_in_barcode: 91.74%
Q30_bases_in_barcode: 91.74%
Q10_bases_in_umi: 99.26%
Q20_bases_in_umi: 96.32%
Q30_bases_in_umi: 89.45%
Q10_bases_in_seq: 99.47%
Q20_bases_in_seq: 97.12%
Q30_bases_in_seq: 91.08%
umi_filter_reads: 8451821 0.84%
umi_with_N_reads: 13355 0.00%
umi_with_polyA_reads: 13044 0.00%
umi_with_low_quality_base_reads:
                                                      8425422 0.84%
mapped_dnbs: 78023582
```

3.1.2 Statistical Report for Reference Genome Alignment

```
$ cat /path/to/output/00.mapping/E100026571_L01_trim_read.Log.final.out
                          Number of input reads |
                                                         783296686
                      Average input read length |
                                                         100
                                    UNIQUE READS:
                   Uniquely mapped reads number |
                                                         624859097
                        Uniquely mapped reads %
                                                         79.77%
                          Average mapped length |
                                                         98.02
                       Number of splices: Total |
                                                         68091842
            Number of splices: Annotated (sjdb) |
                                                         65321302
                       Number of splices: GT/AG |
                                                         66183758
                       Number of splices: GC/AG |
                                                         462810
                       Number of splices: AT/AC |
                                                         43237
               Number of splices: Non-canonical |
                                                         1402037
                      Mismatch rate per base, % |
                                                         0.58%
                         Deletion rate per base |
                                                         0.07%
                        Deletion average length |
                                                         3.87
                        Insertion rate per base |
                                                         0.04%
                       Insertion average length |
                                                         1.26
                             MULTI-MAPPING READS:
        Number of reads mapped to multiple loci
                                                         81769070
             % of reads mapped to multiple loci
                                                         10.44%
        Number of reads mapped to too many loci
                                                         2071180
             % of reads mapped to too many loci |
                                                         0.26%
                                  UNMAPPED READS:
 Number of reads unmapped: too many mismatches |
                                                         0
      % of reads unmapped: too many mismatches
                                                         0.00%
            Number of reads unmapped: too short
                                                         73977028
                 % of reads unmapped: too short |
                                                         9.44%
                Number of reads unmapped: other
                                                         620311
                     % of reads unmapped: other
                                                         0.08%
                                  CHIMERIC READS:
                       Number of chimeric reads |
                            % of chimeric reads |
                                                         0.00%
```

3.1.3 Example of BAM mapping

```
•••
$ samtools view /path/to/output/00.mapping/E100026571_L01_trim_read.Aligned.
sortedByCoord.out.bam | head -3
E100026571L1C009R00301275185
                                    3000095 255
                                                26M121066N74M
                        16
                              1
            GTATATTAATTTGCACTGACTGTCATAACAAAATAC
                               G+:GFFGGFGFFGFGFGFFFFFCFGFCFGGGF-
GGFGFFFFGGFGFFGFFFGFGFFFFFFFFFFFFFFFGGFFGFFGFF
                                                  NH:i:1 HI:i:1
AS:i:88 nM:i:0 Cx:i:4826
                        Cy:i:11598
                                   UR:Z:6FA29
E100026571L1C008R02501359039
                        256
                                    3000101 3
                                                11S33M329303N56M
                              1
                  0
            0
7G=5F2A9F2,3,F8EDFGGD@8GFGGFFAFFG@E?EFFCGF8GGFGFEA7G@FGECFFGGDF;FA<G
HI:i:2 AS:i:66 nM:i:9 Cx:i:13900
                              Cy:i:20099
                                          UR:Z:BA677
E100026571L1C007R00303973559 256
                             1
                                    3000644 3
                                                100M
      GCCTCATTGTGCCCCATATGTTTGCCTATGTTGTGGACTTATTTTCATTAAAACTTTTAAAACATCTTTA-
ATTTTTTTTTTTTTTCATCATTGACCAAGCT
                           -FCA9D?GFFD<-D<CGFEGD-DG*FGFDFBE;E(9BGGE38FFF-
G9GG;0?GGFGB?E@G:GGG3GF79F0GGDG?G<D>F;EG,G?<<CD4>G=>B+C NH:i:2 HI:i:2 AS:i:94
nM:i:2 Cx:i:8839
              Cy:i:7539
                            UR:Z:120CF
```



3.2 merge

3.2.1 Example of Mapped CID List with Reads Count File

```
$ head /path/to/output/01.merge/SS200000135TL_D1.barcodeReadsCount.txt
12286
        19289
2055
        21686
                3
        14086
10005
                2
5040
        12492
                1
        10095
15271
                 6
6032
        10419
7498
        14163
                1
15553
        7772
3206
        13520
                1
13437
        11123
```

3.3 count

3.3.1 Statistical Report for MID Filtering and Gene Annotation

```
$ cat /path/to/output/02.count/SS200000135TL_D1.Aligned.sortedByCoord.out.merge.q10.dedup.
target.bam.summary.stat
## FILTER & DEDUPLICATION METRICS
TOTAL_READS
             PASS_FILTER
                             ANNOTATED_READS UNIQUE_READS
                                                             FAIL_FILTER_RATE
                                                                                    FAIL_
               DUPLICATION_RATE
ANNOTATE_RATE
706628167
              624859097
                              506676745
                                             104709429
                                                             11.57 18.91
                                                                           79.33
## ANNOTATION METRICS
TOTAL_READS
             MAP
                      EXONIC INTRONIC
                                             INTERGENIC
                                                             TRANSCRIPTOME
                                                                            ANTISENSE
624859097
              624859097
                              456866088
                                             49810657
                                                             118182352
                                                                            506676745
109864594
100.0 100.0
               73.1
                      8.0
                              18.9
                                      81.1
                                              17.6
```

3.3.2 Example of Annotated BAM

```
$ samtools view /path/to/output/02.count/SS200000135TL_D1.Aligned.sortedByCoord.out.merge.q10.
dedup.target.bam | grep GE:Z | head -3
E100026571L1C002R00703943265
                                                                                 1040
                                                                                                     1
                                                                                                                          3082766 255
                                                                                                                                                                  11M132671N89M
                    \tt CTGCTGCAGCTTTTTTTTTTTGAGATTTATTTTTATGCTATGTGTATGGGTATTTTGCCTGCATATATGTCTATGCAC-INCOME. THE transfer of the
                                                                      FFFFFECGFDCFGDGDFEE@EEGIBFGGCGFFGACGFCGFFDGDGFFFFFFEGCDFCGFF-
CATGTGTGCAGTGCTTGAG
                                                                                                                NH:i:1 HI:i:1
GG@FFF=EFFDGGGGGFDGFFFGGGFFGGGDFG
                                                                                                                                                             AS:i:88 nM:i:0 Cx:i:7767
                                                                                 XF:i:0 GE:Z:Xkr4
                                                                                                                                              GS:Z:- UB:Z:79E49
                                        UR:Z:7AE49
E100026571L1C006R01702839878
                                                                                 16
                                                                                                     1
                                                                                                                         3108680 255
                                                                                                                                                                  11M187430N89M
                    GTCCTTTTTTTTTTTTTCAAGTTATTCCCTCCATTTCCTCTGGTTCTGTGTTTATGACCAATTGCAAATTG-
TAGAAAATTTGCATGTGTACTGAAC
                                                                                 5;E4FGCGGGGEGFDFFGGG>GFGFDGBFDFDF:EGEFFGFGGGGFFGGGGFGGF-
GFCFGFFGGGGGFGGGGGGGGGGGGGGGFGBGFGGEFFD
                                                                                                                NH:i:1 HI:i:1
                                                                                                                                                                    AS:i:88 nM:i:0 Cx:i:7159
Cy:i:12715
                                        UR:Z:88DB5
                                                                                 XF:i:1 GE:Z:Xkr4
                                                                                                                                              GS:Z:-
E100026571L1C002R04102972006
                                                                                 16
                                                                                                     1
                                                                                                                          3110829 255
                                                                                                                                                                  18M211964N82M
                    ATTGTTTTTTTTTTTTTTTCATGGGAGCTAAAAAATGTTTAATTGTTTGAATAAGAAAAATGTTTCTGATCAAATGTCTCATA-
                                                       CAGCTCATATAAAAT
FGGGGGGGGGGGGGGGGG
                                                                                   NH:i:1 HI:i:1
                                                                                                                                AS:i:88 nM:i:0 Cx:i:11283
UR:Z:887BF
                                 XF:i:1 GE:Z:Xkr4
                                                                                                     GS:Z:-
```

3.3.3 Example of count Gene Expression Matrix

```
$ h5dump -n /path/to/output/02.count/SS200000135TL_D1.raw.gef
HDF5 "/path/to/output/02.count/SS200000135TL_D1.raw.gef" {
FILE_CONTENTS
 group
 group
               /geneExp
 group
               /geneExp/bin1
               /geneExp/bin1/expression
 dataset
               /geneExp/bin1/gene
 dataset
 }
$ h5dump -d /geneExp/bin1/expression /path/to/output/02.count/SS200000135TL_D1.raw.gef | head -15
HDF5 "/path/to/output/02.count/SS200000135TL_D1.raw.gef" {
DATASET "/geneExp/bin1/expression" {
   DATATYPE H5T_COMPOUND {
       H5T_STD_U32LE "x";
H5T_STD_U32LE "y";
       H5T_STD_U8LE "count";
   DATASPACE SIMPLE { ( 73956787 ) / ( 73956787 ) }
   DATA
    (0): {
           4888
           10392,
    },
(1): {
$ h5dump -d /geneExp/bin1/gene /path/to/output/02.count/SS200000135TL_D1.raw.gef |
head -20
HDF5 "/path/to/output/02.count/SS200000135TL_D1.raw.gef" {
DATASET "/geneExp/bin1/gene" {
   DATATYPE H5T_COMPOUND {
       H5T_STRING {
STRSIZE 32;
STRPAD H5T_STR_NULLTERM;
           CSET H5T_CSET_ASCII;
           CTYPE H5T_C_S1;
       } "gene";
H5T_STD_U32LE "offset";
H5T_STD_U32LE "count";
   DATASPACE SIMPLE { ( 24606 ) / ( 24606 ) }
   DATA {
    (0):
           "Gm1992",
           0,
           133
    },
(1): {
```

3.3.4 Example of count Sampling File

```
$ head -8 /path/to/output/02.count/raw_barcode_gene_exp.txt
10392 4888 10551 665954 4
7096 8901 10551 881671 1
7096 8901 10551 357383 20
18783 7397 10551 355789 1
13032 9155 10551 297666 1
13032 9155 10551 298690 1
11778 10617 10551 686313 4
11152 6947 10551 322978 1
```

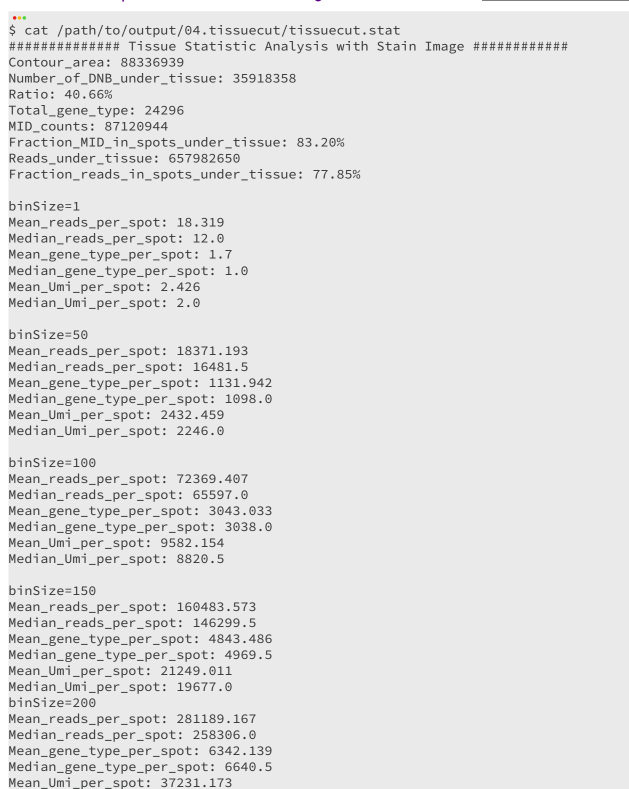
3.4 register

3.4.1 Registered image

File /path/to/output/03.register/ 7_result/SS200000135TL_D1_regist.tif

3.5 tissueCut

3.5.1 Statistical Report for Tissue Covered Region





Median_Umi_per_spot: 34612.5

3.5.2 Example of Gene Expression Matrix for Tissue Covered Region

```
•••
$ h5dump -n /path/to/output/04.tissuecut/SS200000135TL_D1.tissue.gef
HDF5 "/path/to/output/04.tissuecut/SS200000135TL_D1.tissue.gef" {
FILE_CONTENTS {
group
group
           /geneExp
           /geneExp/bin1
group
           /geneExp/bin1/expression
dataset
           /geneExp/bin1/gene
dataset
}
$ h5dump -d /geneExp/bin1/expression /path/to/output/04.tissuecut/SS200000135TL_D1.
tissue.gef | head -15
HDF5 "/path/to/output/04.tissuecut/SS200000135TL_D1.tissue.gef" {
DATASET "/geneExp/bin1/expression" {
  DATATYPE H5T_COMPOUND {
     H5T_STD_U32LE "x";
     H5T_STD_U32LE "y";
     H5T_STD_U8LE "count";
  DATASPACE SIMPLE { ( 60922074 ) / ( 60922074 ) }
  DATA {
   (0): {
        5426,
        2662,
        4
     },
   (1): \{
$ h5dump -d /geneExp/bin1/gene /path/to/output/04.tissuecut/SS200000135TL_D1.tissue.
gef | head -20
HDF5 "/path/to/output/04.tissuecut/SS200000135TL_D1.tissue.gef" {
DATASET "/geneExp/bin1/gene" {
  DATATYPE H5T_COMPOUND {
     H5T_STRING {
        STRSIZE 32;
        STRPAD H5T_STR_NULLPAD;
        CSET H5T_CSET_ASCII;
        CTYPE H5T_C_S1;
     } "gene";
     H5T_STD_U32LE "offset";
     H5T_STD_U32LE "count";
  }
  DATASPACE SIMPLE { ( 24268 ) / ( 24268 ) }
  DATA {
   (0): \{
        "0610005C13R
0,
        23
     },
   (1): {
```

3.5.3 Example of Gene Expression Matrix for Maximum Area Enclosing Rectangle

```
$ h5dump -n /path/to/output/04.tissuecut/SS200000135TL_D1.gef
HDF5 "/path/to/output/04.tissuecut/SS200000135TL_D1.gef" {
FILE_CONTENTS {
 group
 group
            /geneExp
            /geneExp/bin1
 group
 dataset
            /geneExp/bin1/expression
 dataset
           /geneExp/bin1/gene
 group
           /geneExp/bin10
           /geneExp/bin10/expression
 dataset
           /geneExp/bin10/gene
 dataset
           /geneExp/bin100
 group
 dataset
           /geneExp/bin100/expression
           /geneExp/bin100/gene
 dataset
 group
           /geneExp/bin20
           /geneExp/bin20/expression
 dataset
           /geneExp/bin20/gene
 dataset
 group
           /geneExp/bin200
 dataset
           /geneExp/bin200/expression
           /geneExp/bin200/gene
 dataset
           /geneExp/bin50
 group
 dataset
           /geneExp/bin50/expression
           /geneExp/bin50/gene
 dataset
 group
           /geneExp/bin500
 dataset
           /geneExp/bin500/expression
 dataset
           /geneExp/bin500/gene
 group
           /stat
 dataset
           /stat/gene
           /wholeExp
 group
 dataset
           /wholeExp/bin1
 dataset
           /wholeExp/bin10
 dataset
           /wholeExp/bin100
 dataset /wholeExp/bin20
 dataset
           /wholeExp/bin200
 dataset
           /wholeExp/bin50
 dataset
           /wholeExp/bin500
 }
$ h5dump -d /stat/gene /path/to/output/04.tissuecut/SS200000135TL_D1.gef | head -20
HDF5 "/path/to/output/04.tissuecut/SS200000135TL_D1.gef" {
DATASET "/stat/gene" {
   DATATYPE H5T_COMPOUND {
      H5T_STRING {
         STRSIZE 32;
         STRPAD H5T_STR_NULLTERM;
         CSET H5T_CSET_ASCII;
         CTYPE H5T_C_S1;
      } "gene";
      H5T_STD_U32LE "MIDcount";
      H5T_IEEE_F32LE "E10";
   DATASPACE SIMPLE { ( 24606 ) / ( 24606 ) }
   DATA {
   (0): \{
         "Gm42418",
         5851160,
         60.11
      },
   (1): {
```

3.6 saturation

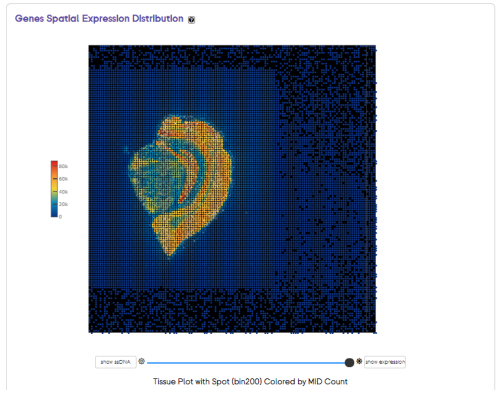
3.6.1 Example of Sequence Saturation File

```
$ head /path/to/output/06.saturation/sequence_saturation.txt
#sample bar_x bar_y1 bar_y2 bar_umi bin_x bin_y1 bin_y2 bin_umi
0.05 21857012 0.2451227 1 16499361 21857012 0.2706192 2851 6616
0.1 43714024 0.3835907 1 26945730 43714024 0.4051051 3822 10805
0.2 87428049 0.5359986 1 40566739 87428049 0.5518293 4724 16267
0.3 131142073 0.6192704 1 49929674 131142073 0.631755 5160 20021
0.4 174856098 0.6722583 1 57307641 174856098 0.6826324 5483 22980
0.5 218570122 0.7092118 1 63557618 218570122 0.7181397 5762 25486
0.6 262284147 0.7365943 1 69087139 262284147 0.7444621 5939 27704
0.7 305998171 0.757775 1 74120410 305998171 0.7648404 6068 29722
0.8 349712196 0.7746872 1 78794642 349712196 0.7811282 6188 31596
```

3.7 report

3.7.1 Example of Statistical Summary Report 3.7.2HTML Report

3.7.2 HTML Report



1.00G Total Reads 37.23K Mean MID per Bin200 **6.34K** Mean Gene per Bin200

Sunburst @	
Total Reads	1.00G
Valid CID Reads	836.72M
Clean Reads	783.30M
Reads Mapped to Genome	708.70M

Sequencing	
Total Reads	1.00G
Rate of Q30 Bases in CID	91.74%
Rate of Q30 Bases in MID	89.45%
Rate of Q30 Bases in Seq	91.08%

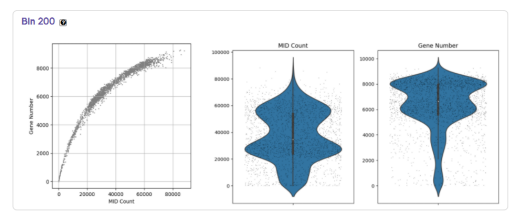


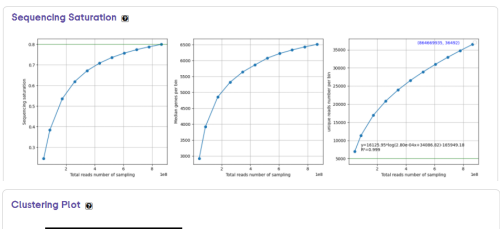
RNA Mapping	
Clean Reads	783.30M
Unique Mapping Reads	624.86M
Multiple Mapping Reads	83.84M
RNA Unmapping Reads	74.60M

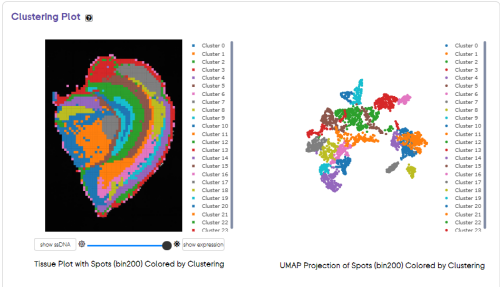
Annotation	
Exonic	456.87M
Intronic	49.81M
Intergenic	118.18M
Transcriptome	506.68M
Antisense	109.86M

TissueCut Total Stat				
Contour Area	88.34M			
Number of DNB Under Tissue	35.92M			
Ratio	40.66%			
Total Gene Type	24.30K			
MID Under Tissue	87.12M			
Fraction MID in Spots Under Tissue	83.20%			
Reads Under Tissue	657.98M			
Fraction Reads in Spots Under Tissue	77.85%			

TissueCut Bin Stat							
Mean Reads	Median Reads	Mean Gene Type	Median Gene Type	Mean MID	Median MID		
18.37K	16.48K	1.13K	1.10K	2.43K	2.25K		
72.37K	65.60K	3.04K	3.04K	9.58K	8.82K		
160.48K	146.30K	4.84K	4.97K	21.25K	19.68K		
281.19K	258.31K	6.34K	6.64K	37.23K	34.61K		
	Mean Reads 18.37K 72.37K	Mean Reads Median Reads 18.37K 16.48K 72.37K 65.60K 160.48K 146.30K	Mean Reads Median Reads Mean Gene Type 18.37K 16.48K 1.13K 72.37K 65.60K 3.04K 160.48K 146.30K 4.84K	Mean Reads Median Reads Mean Gene Type Median Gene Type 18.37K 16.48K 1.13K 1.10K 72.37K 65.60K 3.04K 3.04K 160.48K 146.30K 4.84K 4.97K	Mean Reads Median Reads Mean Gene Type Median Gene Type Mean MID 18.37K 16.48K 1.13K 1.10K 2.43K 72.37K 65.60K 3.04K 3.04K 9.58K 160.48K 146.30K 4.84K 4.97K 21.25K		







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