

The SaSSY Manual

(Version 0.1.2)

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The SaSSY Manual

(Version 0.1.2)

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2 Installing SaSSY

At the time of writing, SaSSY has an “academic” build environment. That means, it works, but it could be improved a whole lot more. It should just build on any Linux / Unix environment but there may be some bugs here and there. The end goal is to have a `./configure → make → make install` setup. Anyone who wants to help me streamline the process and include libtool, autoconf, etc. is more than welcome to help, and any help will be GREATLY appreciated.

SaSSY has only been tested with different versions of the gcc compiler. Most work well, however, I recommend version 4.5.4 as this seems to have given the best results. See notes below for any compiler specific information.

2.1 Dependencies

There are a few dependencies which will need to be installed before SaSSY can be compiled and run. The versions given here are the versions I have tested but your mileage may vary.

| Dependency | Version | Download |
|-------------------|---------|---|
| String-CamelCase | v-0.02 | http://search.cpan.org/~hio/String-CamelCase-0.02/ |
| pcre | v-8.11 | http://www.pcre.org/ |
| ncurses | v-5.7 | http://ftp.gnu.org/pub/gnu/ncurses/ |
| Google sparsehash | v-1.10 | http://code.google.com/p/google-sparsehash/ |
| readline | v-6.1 | http://tiswww.case.edu/php/chet/readline/rltop.html |

Table 1. SaSSY dependencies, versions and URLs

These need to be installed and tested. If you are installing SaSSY in your home directory then you may still be ok, it depends on what's installed on your system. It's best to just try and install SaSSY and deal with any errors as they occur. Otherwise, see section 2.4 below.

2.2 Get the source

Please visit sassy.mikeimelfort.com

2.3 System Install

Change into the src directory and type:

```
$ make
```

And go get a cup of coffee. This will produce an executable; either Sassy64 or Sassy32, depending on your architecture. Make symlinks from this folder to your bin directory. Also make sure you symlink the 'formatdefs' folder into the same directory as the executable.

2.4 Local Install

This section assumes that you will be installing SaSSY in your home directory which I'll call '\$HOME'. You have extracted the code into a working directory called \$SASSY. The first thing to do is simply try to build it using the 'make' command. However, this may not work because you may not have all the dependencies installed. Check the error messages which are printed when you try to make, to see which dependencies are missing. I'll assume that you have gcc or a similar compiler installed. Do some or all of the following, and then try to make again.

2.4.1 Install dependencies

First install any missing dependencies. I'll assume that you are going to install dependencies in '\$HOME/local/'. If you are going to install them elsewhere, you'll need to modify these code snippets.

NOTE: Do not just type '\$HOME'. On my system my home directory is located at '/home/uqmimelf' so I would write '\$HOME/local/' as '/home/uqmimelf/local/'.

For all the dependencies EXCEPT String-CamelCase and readline use the following commands:

```
$ ./configure --prefix ~/local
$ make
$ make install
```

When you install readline you MUST link it against ncurses. So change the configure line to:

```
$ ./configure --prefix $HOME/local/ --with-curses
```

When you install String-CamelCase you need to use the 'perl' way of doing things:

```
$ perl Makefile.PL prefix $HOME/local
$ make && make install
```

2.4.2 Modify Makefile.inc

Now you will need to modify a file called Makefile.inc. It is located in: `$SASSY_DIR/src/.makefile_pieces`. You need to modify the `BASE_CCFLAGS` and `BASE_LIBS` lines located at the top of this file to make SaSSY use the local versions of the dependencies. Originally, these lines should look like:

```
BASE_CCFLAGS := -O3 -Wall -Werror -Wno-unused-function
BASE_LIBS = -lreadline -lncurses -lpcrc
```

If you installed any of the dependencies locally, you will need to change the `BASE_CCFLAGS` to this:

```
BASE_CCFLAGS := -O3 -Wall -Werror -Wno-unused-function -I $HOME/local/include
```

If you installed readline as one of the local dependencies, you will need to change the `BASE_CCFLAGS` to this:

```
BASE_CCFLAGS := -O3 -Wall -Werror -Wno-unused-function -static -I \
$HOME/local/include
```

If you installed *any* of the dependencies locally, you will need to change the `BASE_LIBS` line to something like this:

```
BASE_LIBS = -Bstatic $HOME/local/lib/libreadline.a -L$HOME/local/lib -lncurses \
-lpcrc
```

2.4.3 Set the PERL5LIB environment variable

If you needed to install String-CamelCase in your home directory you will need to tell the system where to find it. Type the following on the command line before you try to compile SaSSY. Your setup may be slightly different, but this should give you the main idea...

```
$ export PERL5LIB=$HOME/local/lib/perl5/site_perl/5.10.0/
```

3 Running SaSSY

SaSSY typically runs in two stages. First, there is a “load and save” stage where the raw data is loaded, naïve contigs are made and then all the info is saved into binary files. The second is a “re-load and extend” stage where the information is loaded from the binary files and then the naïve (or other) contigs are extended. The following describes a typical work flow.

3.1 Create “FileSet” tokens

If you're impatient:

```
$ SassyXX newFileSet
```

...and follow the prompts...

If that approach fails:

SaSSY loads raw data from a variety of different short read formats. Some of these formats store the read data in one text file, others use two. To avoid any confusion, I'll call each set of files a *FileSet*. Typically, each FileSet will contain paired short reads from a single library, with a given insert size and relative orientation. However, sometimes, for example with Illumina mate-paired data, there appears to be two distinct libraries; one with the intended insert size and relative orientation, and another *shadow library* with an insert size of about 200-400bp and a relative orientation in the opposite orientation. There are probably other good reasons for having multiple libraries in one FileSet but the shadow library is the most common one I've seen. SaSSY has been designed to handle FileSets which contain multiple libraries, you just need to tell it that they're there. This is done using a *FileSet token*.

Basically, a FileSet token wraps the raw data files and stores information about any libraries the FileSet contains. To make a FileSet:

```
$ Sassy64 newFileSet
```

```
***Sassy64*** Version: 0.1.0.0
```

```
Copyright (c) 2008, 2009, 2010 Michael Imelfort and Dominic Eales.
```

```
This is free software; see the source for copying conditions. There is NO  
warranty; not even for MERCHANTABILITY or FITNESS FOR A PARTICULAR PURPOSE.
```

```
Creating a new file-set...
```

```
NOTE: If you get an error saying the format does not match when you are sure it does  
then chances are you will need to convert the file from windows style new lines to dos
```

```
Please choose the format of the file-set:
```

1. FASTA-dualfiles (illumina style)
2. FASTA-dualfiles (plain vanilla)
3. FASTA-singlefile (TAGdb output)
4. FASTA-singlefile (illumina style shuffled)
5. FASTA-singlefile (plain vanilla shuffled)
6. FASTQ-dualfiles
7. FASTQ-singlefile
8. Illumina SCARF
9. Unknown file format

Choice: █

Now enter a number from 1 – 9, depending on the type of input file you have.

Choice: 2

** All files must be in the same directory.
** All files must have the same number of lines.

Enter file 1 (FASTA_1 file): █

We have chosen “FASTA-dualfiles (plain vanilla)” so the program expects two files to be entered, one for each end of the paired read. Enter the path to the first data file. This must be an exact path. You will be asked to enter the path to the second file after this.

Enter file 1 (Sequence File 1 file): PATH_TO_FILE_1
Enter file 2 (Sequence File 2 file): PATH_TO_FILE_2

Number of bases to trim from end of read: [0 for none] █

If you want to do permanent trimming on this data file then do it here. NOTE: you can trim on a per-run basis on the command line when executing SaSSY so it may be better to enter 0 here. Next you will be asked to enter *estimates* for the insert size of the library. SaSSY will use these estimates to calculate more exact figures during runtime.

NOTE: SaSSY expects the insert size to be the distance BETWEEN the centres of the reads. This is different from all other assemblers and is something I will rectify when I find the time. For now you need to remember that if you have reads which have come from a library with a fragment size of 580bp using 100bp paired reads then the estimated insert size will be 480bp. In velvet you would use 580bp. This could/will be rectified in future versions of SaSSY.

Number of bases to trim from end of read: [0 for none] 0

Estimated library insert size #1: [0 to stop] 3500
Estimated library insert size #1: [0 to stop]
Orientation code for insert #1
0 <--- >--->
1 <--- <---
2 >--- <---
3 >--- >--->
: 0

```

Estimated library insert size #2: [0 to stop] 250
Estimated library insert size #2: [0 to stop]
Orientation code for insert #2
0 <--- <--->
1 <--- <---
2 <---> <---
3 <---> <--->
: 2
Estimated library insert size #3: [0 to stop] 0

File-set saved to `READS_DIRECTORY+FILENAME.sassy'

```

Enter the insert sizes and orientation types for all the libraries in this FileSet. Enter 0 to stop. You will find a FileSet token file in the directory which contains the reads. It has local references to the data files so it should stay in that folder. If you need to modify any of the values you entered, it is plain text so this should be straight forward. A typical FileSet token looks like this:

```

name:RANDOM_BAC_1_sequence.txt
format:Illumina SCARF
file_names:RANDOM_BAC_1_sequence.txt,RANDOM_BAC_2_sequence.txt
num_reads:7308118
read_length:50
trim_length:0
insert_lib:M:3500 St:0 0:0 Nr:0
insert_lib:M:250 St:0 0:2 Nr:0
--

```

This is the file which is passed to SaSSY with the '-f' flag in section 2.2

3.2 Load raw data and make naïve contigs

```
$ SassyXX -s -d OUTPUTDIR -f FileSet[,MAX_READS] [-f FileSet[,MAX_READS]]
```

Compulsory parameters:

| | |
|---|--|
| <code>--dir -d WORKING_DIR</code> | The directory to save output including where the assembled contigs are saved |
| <code>--save -s</code> | Load data from FileSets, do basic processing. Then save to output directory and exit |
| <code>--FileSet -f FileSet[,MAX_READS]</code> | Add sequences described by 'FileSetfile' 'MAX_READS' is maximum paired reads to load |

Optional parameters:

| | |
|--------------------------------------|---|
| <code>--loglevel -v LEVEL</code> | Set log level. 0 for no logging [default: 0] |
| <code>--readlength READLENGTH</code> | Set the effective readlength to this value (override values in the FileSet tag) |
| <code>--naive_off OFFSET</code> | Maximum offset used to make naïve contigs [default: 4] |

`--ext_off OFFSET`

Maximum offset used during extension (this value
cannot be exceeded during the extension step)
[default: 12]

... And go make a cup of coffee.

NOTE: SaSSY can take any number of FileSet tokens.

NOTE: The readlength used in the assembly will be equal to the largest odd number which is less than or equal to the shortest readlength of any of the input FileSets. For example: If one FileSet has a readlength of 100bp and another has a readlength of 36bp, SaSSY will perform the assembly with a readlength of 35bp. The last base from reads in the later FileSet and the last 65bp from reads in the the former FileSet will be **DISCARDED**. It is recommended that if you have to use data with readlengths that vary this much then you should cut down the longer reads using the script called 'cutDown.pl' located in the scripts directory. Set -readlength to control this variable.

The output of this command is the saved binary files, and a file called 'out.naive_contigs.fa' which contains the naïve contigs. If a log level was specified, there will also be a file called sassy.log. Logging is not required, but if you like getting a log file, a level of 4 produces a manageable amount of information.

Loading part of a FileSet: If you would only like to load the first 1,000,000 reads of some FileSet, then modify the command line like this:

`-f FileSet_1`

to:

`-f FileSet_1,1000000`

Naïve and extension offsets:

`--naive_off OFFSET`

`--ext_off OFFSET`

Set the maximum allowable offsets for naïve contigs and extended contigs. The offset is the length of the non-overlapping parts of two overlapped reads. By default, SaSSY sets naïve offset to 4 and extension offset to 12. Thus, during naïve contig creation, two reads will be joined together if they are offset from each other by no more than 4 bases. During contig growing, this is extended to up to 12 bases. The greater the cut off, the more edges will be made between the reads (and hence more memory will be used).

You may override these parameters with any EVEN value between 2 and

16 inclusive. For example, if you were trying to assemble a data set with ridiculously high coverage, it may help to set these values to 2bp and 6bp respectively, which will reduce the number of edges made and hence the amount of memory used - hopefully without sacrificing the connectivity of the graph. If you had a very low coverage dataset, you may want to set them to 8bp and 12bp to try induce extra edges in the naïve contig graph.

3.3 *Extend naïve contigs / other sequences*

```
$ SassyXX -l -d OUTPUTDIR
```

Compulsory parameters:

| | |
|-----------------------------------|--|
| <code>--dir -d WORKING_DIR</code> | The directory where saved data is stored and where the assembled contigs are saved |
| <code>--load -l</code> | Load the data from saved state |

Optional parameters:

| | |
|---|---|
| <code>--loglevel -v LEVEL</code> | Set log level. 0 for no logging [default: 0] |
| <code>--no_strict</code> | Relax the growing algorithm, longer contigs but causes more over assemblies / chimeras [default: STRICT = true] |
| <code>--no_olap</code> | Skip overlaperation [default: OLAP = true] |
| <code>--upper_coverage_percent PERCENT</code> | Set the maximum coverage allowable for a naïve Contig to be used in stretching [default: true (use all contigs)] |
| <code>--ext_off OFFSET</code> | Maximum offset used during extension [default: 12] |
| <code>--olap_min_size MIN_SIZE</code> | The minimum number of bases two Contexts must overlap [default: highest upper cut off] |
| <code>--olap_max_wastage MAX_WASTAGE</code> | The maximum number of bases which can be wasted during an overlaperation event [default: 2 * read length] |
| <code>--scaff_link_cut_off NUM_LINKS</code> | Minimum number of links needed to scaffold 0 to skip scaffolding [default: 5] |
| <code>--scaff_insert_cut_off CUT_OFF</code> | For scaffolding, use only reads with insert greater than CUT_OFF [default: 0] |
| <code>--ignore_trap TRAP_INSERT</code> | When using a small supplementary insert size to clean the data, ignore this in extensions. 0 to use all InsertLibs. [default: Ignore = 0] |
| <code>--graph</code> | Output the scaffold graph to stdout in format [default: no graph] |
| <code>--csv</code> | Output the information used in scaffolding to stdout in CSV format [default: no csv] |

This should / will run significantly faster than the load & save run. This stage MUST be able to find a multiple fasta file, called out.naive_contigs.fa, in the working directory. It will output three files; out.advanced_contigs.fa, out.final_contigs.fa and out.scaffolds.fa. The first is the extended contigs before

final overlapping, the second contains the same contigs after overlapping and the final one contains the overlapped contigs after scaffolding. If a log level was specified, there will also be a file called sassy.log

Using “alien” seeds: This stage MUST be able to find a multiple fasta file, called out.naive_contigs.fa in the working directory. BUT this doesn't mean that this file has to contain contigs from SaSSY. You *could* use any fasta file here...

Extension options: These options are used during the extension phase of the algorithm.

`--no_strict`

By default, the extension phase runs in a “strict” mode. This means that during the growing phase it will try to extend the input contigs as best it can but it will not try to “push” things too far, as this can create chimeric contigs. If you have a low coverage data set then it may help to turn this feature off. To do this use the '--no_strict' flag

NOTE: This will nearly always let you produce longer contigs, you just need to spend more time after assembly to make sure that they are correct.

`--upper_coverage_percent PERCENT`

SaSSY uses the naïve contigs generated in the first stage to “stretch” extended contigs. For example, when an extension fails it may be that SaSSY has extended the seed contig and is part way through another long naïve contig. Because SaSSY “trusts” naïve contigs, especially longer ones, it will stretch the extension to the end of that contig and then try to continue extending. This can cause problems if SaSSY tries to stretch across a highly repetitive region. Setting this parameter will force SaSSY to use contigs for stretching only if they are within PERCENT percent of the mean coverage of the ten longest naïve contigs. This option will not work if you are using alien seeds (see above).

`--ignore_trap -I TRAP_INSERT`

Illumina paired-end data sets typically have the insert size distribution shown in Figure 1 below.

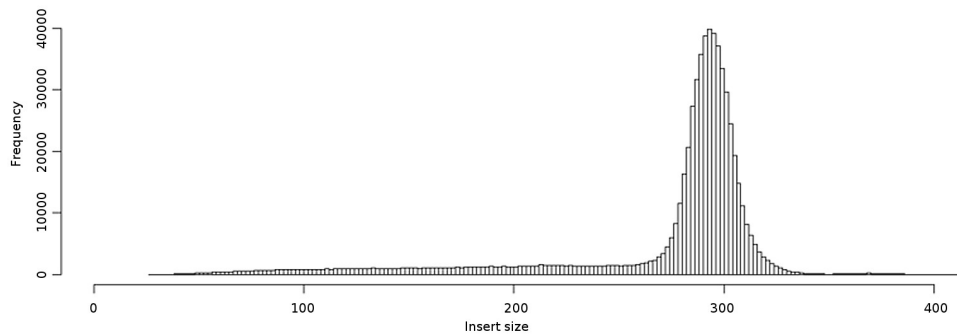


Figure 1. Typical distribution of inserts for Illumina Paired-end data.

Typically, there will be a lot of rubbish on the left hand side of the mean. Because SaSSY relies on insert size stability as opposed to coverage stability this can cause problems. To reduce the effect that this has on the assembly you can set a “trap” insert when defining the FileSet. For example, in the image above you would tell SaSSY that this data file has two insert sizes; one with a mean of approximately 300bp and one with a mean of approximately 50bp. The result is that many of the rubbish reads get separated from the “real” reads into their own little space, and the standard deviation about the upper mean gets a lot tighter. You can tell SaSSY to ignore these reads during the extension process by setting the ignore flag. For the diagram above, it may improve the assembly to ignore all reads with an insert size less than say 220bp.

NOTE: This is a global setting. Thus, for the example just given, if you had a “real” library with an insert size of less than 220bp, most of its reads would be ignored.

```
--ext_off OFFSET
```

You can override the value set for `ext_off` during naïve contig creation, however, it must be less than or equal to the value used for `ext_off` during the first stage.

Overlapation options: These options concern the overlapation stage which follows extension.

```
--no_olap -o
```

Overlapation can sometimes be a little naïve. If your Contigs are looking very chimeric it may be useful to turn this feature off. During the scaffolding stage, the advanced contigs will be used instead of the final ones.

```
--olap_min_size MIN_SIZE
--olap_max_wastage MAX_WASTAGE
```

Overlapation occurs when two contigs to have a **perfect overlap** (after end trimming) at least as long as the `olap_min_size` variable. By default this value is based on the largest insert size. If you are lucky enough to have a

data set with a number of different insert sizes, say 200bp, 2000bp and 10000bp you may want to set this parameter to something slightly larger than 2000bp so you can make use of these reads during overlaperation. Otherwise SaSSY will require two extended contigs to share a perfect overlap of about 10000bp for overlaperation to occur. Use `olap_max_wastage` to override the maximum number of bases which can be trimmed from the ends of the contigs to make a perfect overlap.

Scaffolding options: These options concern the scaffolding stage which is the final stage of the algorithm.

`--scaff -S NUM_LINKS`

Minimum number of links needed to make scaffold. By default this is set to 5. During scaffolding there needs to be a minimum of 5 paired links between two contigs for there to be any chance of a scaffold occurring. For lower or higher coverage datasets you may wish to lower or raise this amount, or let NUM_LINKS be 0 to skip scaffolding altogether.

`--scaff_insert_cut_off CUT_OFF`

Perhaps you don't want to reads from a 200bp insert size library to make scaffolds. You can control the minimum insert size used for scaffolding here.

Output options: Generating more output...

`--graph -g`

Output the scaffolds as a GraphViz formatted graph to STDOUT. Can be useful for seeing how the final contigs were formed into scaffolds. Using this flag will output two graphs. One containing all the possible ways SaSSY could have built the scaffolds and one containing the only the configuration which SaSSY used to create the final 'out.scaffolds' file.

`--csv`

Output all the links which could be used to make scaffolds in csv format to STDOUT. **This may generate a serious amount of output.** This will give every link between every contig which SaSSY identified during the scaffolding stage. It may be useful to examine this output if you feel that SaSSY made some bad scaffolding choices.

The fields of the csv file are given below:

| Column | Description |
|------------|---|
| Base | The first contig used in the link. |
| Linker | The second contig used in this link. |
| IsSameDirn | Is the Linker facing in the same direction as the Base? |
| IsBefore | Does the Linker lie before or after the Base? |

| | |
|------------|--|
| FileId | Identifies which file this linking pair is from. |
| BasePos | The position of the read in the Base contig. |
| BaseDist | The Distance from the read to the end which closest to the Linker. |
| LinkerPos | The position of the read in the Base contig. |
| LinkerDist | The Distance from the read to the end which closest to the Base. |
| PairKey | A sorting key which is made from the IDs of the two contigs. |
| TypeKey | A sorting key which is made from the PairKey and the relative directionality of the Linker with respect to the base. |

Table 2. Description of the scaffold csv columns.

NOTES:

1. The ID of the Base is always numerically lower than the ID of the Linker.
2. All directions are calculated relative to the directions that the contigs have been written in the file 'out.final_contigs.fa' (or out.advan_contigs.fa if -no_olap is set).
3. All distances are calculated relative to the centre of the reads. Thus, the first read in a contig lies at position 0 and the last read lies at position L - RL where L is the length of the contig and RL is the read length.

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