SLALOM (v2.1.4) User Manual

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1. About SLALOM

SLALOM (Statistical Analysis of Locus Overlap Method) is a standalone software tool for analysis of positional data. Namely, it compares two distinct lists of site intervals (hereafter – sites; in the associated publication referred to as continuous sequence elements - CSEs) in a given set of sequences to evaluate the overlap quality according to rules set by the user. The sites are presented as sequence coordinates, with the start and end positions, while the sequences can be of any nature (e.g., alphabetical or time series). This manual is written as a tutorial, each section describing a specific aspect; for the complete list of available command line options see the last section. Please, read the associated publication [Prytuliak, et al., SLALOM, a flexible method for the identification and statistical analysis of overlapping continuous sequence elements in sequence- and time-series data, Publication pending] for detailed explanations of the method, specific cases, and comparison with the other tools.

2. Download and Installation

SLALOM is open-source software distributed under the GPLv3 license; its latest version can be freely downloaded from the GitHub repository: https://github.com/BCF-calanques/SLALOM. Alternatively, if git is installed, type in the terminal:

```
>git clone -b master --single branch https://github.com/BCF-calanques/SLALOM --depth 1
```

This will generate the directory <code>slalom/</code> with the source code in the subdirectory <code>slalom/src/</code> and all the mentioned test files in the subdirectory <code>slalom/test data/</code>.

SLALOM is a command-line tool written in Python. It is cross-platform and does not require additional installation beyond Python interpreter of version 3.5 (https://www.python.org/downloads/) scientific library and for computing numpy (https://www.numpy.org). To install numpy from the command line using Python package manager, type:

```
>pip install numpy
```

That's all! To use SLALOM, simply run slalom.py. For example:

```
>python SLALOM/src/slalom.py --help
```

3. Input Files

slalom.py expects four input files: two annotations (i.e., lists of sites in given sequences), the sequence length table and the group-mapping table, the last two being optional:

```
>python slalom.py [options] -a1 <first_annotation_file> -a2 <first_annotation_file> [-s
<sequence length table>] [-m <group mapping table>] -o <output file>
```

The main input files are the two site annotations to compare. In the basic case, they contain 3 columns: the sequence identifier (SID) the start and the end positions of the sites. For example:

```
>cat test anno1.tsv
        10
seqA 6
seqA 8
           12
seqA 9
           18
seqB 15
          25
>cat test anno2.tsv
seqA 5
          11
seqA 7
           10
seqA 9
           15
seqB 15
           19
seqB 21
```

By default, the numeration of symbol positions in a sequence starts from 1. Both start and end positions are considered included. That means that the length of a site that starts at 15 and ends at 25 is 11 symbols. The sites can overlap, contain or be duplications of each other.

The annotation files must be provided with the options -a1/--anno1file and -a2/--anno2file.

To complete the input, the user needs to specify the sequence length (say, all the sequences are 50 symbols long) with the option $-1/--seqlen_value$ and specify the output file with the option -o/--outfile:

```
>python slalom.py -a1 test_anno1.tsv -a2 test_anno2.tsv -1 50 -o output.tsv
```

Alternatively, the sequence lengths can be provided in the file using the option -s/--seqlenfile. This is the only option, if the sequences have different lengths. The file should contain two columns – the SIDs and their lengths:

```
>cat test_seqlenfile1.tsv
seqA 50
seqB 30
seqC 45
seqD 60
>python slalom.py -a1 test_anno1.tsv -a2 test_anno2.tsv -s test_seqlenfile1.tsv -o
output.tsv
```

4. Analysis of Grouped Sequences

Sometimes the sequences and sites are grouped into categories and a separate analysis within each category is required. To enable sequence grouping, the user should provide the file with sequence-to-group mapping. The file should contain two columns: SIDs and respective group identifiers (GIDs). The same sequence can belong to multiple groups; a group can contain arbitrary number of sequences:

```
>cat test_groupmapping.tsv
seqA group0
seqB group0
seqB group1
seqC group1
seqD group2
```

For multi-group analysis, the user should also specify the group for each site. An extra column should be added to the annotation files:

```
>cat test anno3.tsv
seqA group0 8
seqA group0 21
                      15
seqA group0
                 21
                        25
group0
seqB group1
seqC grown
                 6
                 8
seqB group1
seqC group1
                        11
                 18
                        24
seqD group2
                        9
                  1
seqD group2 1 5
                        12
>cat test anno4.tsv
seqB group1 8
                        12
seqC group1
                  11
                        16
seqD group2
                        10
```

Note that in this example, the two sites in <code>seqB</code> in <code>test_anno3.tsv</code> are no longer considered overlapping for the analysis, as they belong to different groups.

The group-mapping table must be provided with the option -m/--mapfile:

```
>python slalom.py -a1 test_anno3.tsv -a2 test_anno4.tsv -s test_seqlenfile1.tsv -m test_groupmapping.tsv -o output.tsv
```

An input annotation with the grouping information can be reused as the group-mapping table (duplicating records are no problem):

```
>python slalom.py -a1 test_anno3.tsv -a2 test_anno4.tsv -s test_seqlenfile1.tsv -m test_anno3.tsv -o output.tsv
```

The four input files specified by the -a1, -a2, -s, and -m options provide flexibility in general cases. However, in many practical situations, there is some additional information available allowing for simpler input. For example, when the groups do not overlap, a site can belong to one group only, and therefore it is no longer necessary to specify GIDs in the annotation files. This can be indicated by the option -nog/--non_overlapping_groups. Furthermore, if each sequence forms its own unique group, the user can activate the option -sg/--sequences_as_groups and skip the group-mapping file. These and other options useful in simplified cases are discussed in more detail further down in this manual.

5. The Main Output File

The main output file begins with several commented (i.e., starting with #) lines, containing information about the input command line as well as about the statistics calculated:

```
# This file was generated at 2017-01-01 00:00:01 with SLALOM
# Command line options (unquoted and unescaped): -a1 test anno3.tsv -a2 test anno4.tsv
-s test seqlenfile1.tsv -m test anno3.tsv -o output.tsv
# The following statistics have been calculated:
    Nseq: Number of sequences
    P1: Fraction of symbols present in the first annotation
    P2: Fraction of symbols present in the second annotation
    PP: Fraction of symbols present in both annotations
    AP: Fraction of symbols absent in the first annotation but present in the second
    PA: Fraction of symbols present in the first annotation but absent in the second
    AA: Fraction of symbols absent in both annotations
    ACC: Symbol-wise accuracy
    MCC: Symbol-wise Matthews correlation coefficient
    F1: Symbol-wise F1 score
    SiteN1: Number of sites in the first annotation
    SiteN2: Number of sites in the second annotation
    SiteN1m: Number of matched sites in the first annotation
    SiteN2m: Number of matched sites in the second annotation
    SiteL1: Total length of sites in the first annotation
    SiteL2: Total length of sites in the second annotation
     SiteF1: Site-wise F1 score
     SitePCV: Site-wise positive correlation value
# All the provided dataset-wide averages are macro-averages of the group-wide metrics
```

This block is followed by the table with the first row being the header containing the column names. The commented block is helpful at the first usages of SLALOM. However, for easier further processing of the output, it can be suppressed with the option -c/-clean:

```
>python slalom.py -a1 test_anno3.tsv -a2 test_anno4.tsv -s test_seqlenfile1.tsv -m test_groupmapping.tsv -o output.tsv -c
```

If this option is provided, only the actual table (with the header) row is returned:

```
>cat output.tsv
Group Nseq
               Ρ1
                          SiteN1 SiteN2 SiteN1m SiteN2m SiteL1
                                                                 SiteL2
       2
               0.2125
                          3
                                  0
                                          0
                                                 0
                                                         17
                                                                 0
group0
               0.1467
                         2
                                  2
                                          1
                                                 1
                                                         11
                                                                 11
group1
                                                                         ...
group2 1
               0.2000
                                  1
                                          2
                                                 1
                                                         12
                                                                 6
Average 1.6667 0.1864 ... 2.3333 1.0000 1.0000 0.6667 13.333 5.6667 ...
```

For metrics like total site length, the sum may be more informative than the average. The option -sum/--calculate sums will add the corresponding row to the output:

Note that sequences belonging to multiple groups are counted the corresponding numbers of times.

6. The Operating Modes

SLALOM supports three different approaches of handling overlapping sites in one annotation: the symbol-resolved, gross, and enrichment modes. In addition, the annotations can be treated symmetrically – if they have similar origins and reliability – or the first annotation can be used as benchmark to evaluate the second one: the equal and benchmarking modes.

In the output file shown in the previous section, one can see that the total length of sites in the first annotation (the column <code>sitel1</code>) for <code>group2</code> equals to 12 or the fraction of 20% (the column <code>P1</code>). This means that implicit merging of the two overlapping sites (1-9 and 5-12) happened. Note, however, that the number of sites in the first annotation (the column <code>siteN1</code>) still equals to 2 and the site-based statistics are calculated accordingly. This behavior is termed 'symbol-resolved mode' and is applied by default.

Sometimes, however, there is a need to count the symbols as many times as they occur in a given annotation, i.e., as they are traversed by a site. This behavior is termed 'gross mode' and is invoked by providing the value gross with the option -E/--enrichment count:

```
>python slalom.py -a1 test anno3.tsv -a2 test anno4.tsv -s test seqlenfile1.tsv -m
test groupmapping.tsv -o output.tsv -E gross -c --quiet
>cat output.tsv
Group Nseq ... SiteN1 SiteN2 SiteN1m SiteN2m SiteL1 SiteL2
group0 2
             ... 3
                            0 0 17
                                                 0
             ... 2
group1 2
                      2
                            1
                                          11
                                                 11
                                   1
group2 1
               2
                            2
                     1
                                          17
                                   1
                                                 6
Average 1.6667 ... 2.3333 1.0000 1.0000 0.6667 15.000 5.6667 ...
```

This time, the total length of sites in the first annotation is 17; the fraction of symbols is not calculated in the gross mode, as the metric could exceed 1 now.

In some other situations, the user may want to consider only those positions that are traversed by a certain minimal number of sites – the 'enrichment mode'. This number must be provided with the option –E/--enrichment_count. In the enrichment mode, the calculated metrics have slightly different names. For example, the fraction of symbols *present* in the first annotation (the column P1) becomes the fraction of symbols *enriched* (the column E1). If it is set, for example, to 2, this results in considering only those positions that belong at least to 2 sites. In the provided example, there are no such positions in the first annotation for group0 and group1; for group2, there are only 5 positions (5-9 in seqD) for 60 symbols, which results in the fraction of 0.0833:

In addition to different strategies of resolving the overlaps within one annotation, the annotations can have different relations to each other. One scenario is comparing annotations, which relate to the same type of sites (e.g., genes in both annotations), come from similar origins, and have comparable quality. This scenario is called 'equal mode' and is assumed by default. However, when sites in different annotations represent different types of regions (e.g., genes and promoters) or have significantly different reliability (e.g., experimentally verified and predicted), a different scenario becomes relevant. This scenario is called 'benchmarking mode' and is assumed if the option -b/--benchmarking is provided:

```
>python slalom.py -a1 test_anno3.tsv -a2 test_anno4.tsv -s test_seqlenfile1.tsv -m test_groupmapping.tsv -o output.tsv -b -c
```

The change from the equal to benchmarking mode involves both, naming conventions, as well as enabling calculation of additional performance measures. The names are chosen to highlight the nonsymmetry in the benchmarking mode: the 'first annotation' becomes 'benchmark', the 'second annotation' becomes 'prediction', the metric 'present in both annotations' becomes 'true positive', etc. Nevertheless, changes in names do not change the calculations: the values of the returned metrics are equivalent in both modes. In the equal mode, however, only symmetric performance measures – those unaffected by flipping the annotation files – are calculated. Accuracy (ACC) and F1 score are examples of symmetric measures. In the benchmarking mode, additional, non-symmetric measures are calculated, for example, specificity (SPC) and precision (PPV). Thus, the equal mode is effectively just a subset of the benchmarking mode and brings only small advantages with respect to speed and output file size; however, it was implemented to prevent the user from misinterpreting certain metrics in the case of symmetric annotations.

The operating modes should not be confused with the simplified modes for easier processing of certain file formats (see next two sections) and can be used independently of those.

7. Annotations in GenBank Format

Although SLALOM works primarily with tabular input, it has a special simplified mode to compare genome annotations in GenBank format. This mode works under the assumption that both annotations are of the same sequence. The sequence length is automatically read from the files and must not be provided separately. To start the program in the GenBank mode, the user needs to provide the --genbank option:

```
>python slalom.py --genbank -a1 N_pharaonis_annotation1.gb -a2 N_pharaonis_annotation2.gb -o comparison_stats.tsv -c
```

In the given example, there will be one data row in the table, as only one sequence (although perhaps a very long one) is analyzed:

```
>cat comparison_stats.tsv
Nseq ... SiteN1 SiteN2 SiteN1m SiteN2m SiteL1 SiteL2 SiteF1 SitePCV
```

However, the user can analyze each strand or reading frame separately. For this, the value strand or frame respectively must be provided with the option -d/--detect:

```
>python slalom.py --genbank -a1 N_pharaonis_annotation1.gb -a2 N_pharaonis_annotation2.gb -o comparison_stats.tsv -c -d frame -sum
```

Now, the output file contains eight rows: the header, statistics for each of the six reading frames, and the bottom lines with the averages and sums:

>cat comparison stats.tsv										
	_		SiteN1	SiteN2	CitaN1m	CitaNom	SiteL1	SiteL2	SiteF1	SitePCV
Frame	Nseq									
+1	1		475	458	453	453	419436	410952	0.9711	0.9711
+2	1		473	460	459	459	416979	411405	0.9839	0.9839
+3	1		459	441	439	440	400059	394497	0.9766	0.9767
-1	1		456	438	436	436	384372	382003	0.9754	0.9754
-2	1		382	376	372	372	344138	339013	0.9815	0.9815
-3	1		449	435	430	431	402931	401475	0.9740	0.9740
Average	1.0000		449.00	434.67	431.50	431.83	394652	389891	0.9771	0.9771
Sum	6		2694	2608	2589	2591	2367915	2339345		

From this table, one can see that there are a total of 475 genes in the reading frame +1 of the first annotation but only 458 genes for the second annotation. Moreover, 453 genes in each annotation overlap with a gene from another annotation (at this stage, it is not possible to conclude that all the matches are reciprocal). The total length of the genes is 419,436 bp for the first annotation and 410,952 bp for the second annotation. If for some reason, there are any overlapping genes annotated, they are merged for this residue calculation, as SLALOM is operating in the default symbol-resolved mode. Also because of the merging, the total gene length without the frame separation (2,359,033 bp) is less than the sum of the frame-wise gene lengths for the first annotation (2,367,915 bp).

8. Annotations in BED Format

SLALOM offers a simplified mode to process annotations in BED format. To use it, the user has to provide the option --bed, as well as the two annotation files, the sequence length table, and the output file:

```
>cat test bed1.bed
      0
chr1
               10
       15
                25
chr1
       40
                50
chr1
       12
               27
chr2
>cat test bed2.bed
      Ω
               12
chr1
       25
               42
chr1
>cat test seqlenfile2.tsv
       50
chr1
chr2
        40
>python slalom.py --bed -a1 test bed1.bed -a2 test bed2.bed -s test seqlenfile2.tsv -o
output.tsv -c
```

Note that although SLALOM by default counts residues starting from 1 with both start and end residues being included, it is automatically adjusted to the BED format specifications, which require counting starting from 0 with only the start being included.

The default output table contains only the summary line:

As the BED format does not explicitly specify group identifiers, grouping of sequences is not possible in this simplified mode. However, with the option <code>-sg/--sequences_as_groups</code>, each sequence will be treated as a group on its own and the statistics for each sequence will be printed separately:

```
>python slalom.py --bed -a1 test bed1.bed -a2 test bed2.bed -s test seqlenfile2.tsv -o
output.tsv -c -sg -sum --quiet
>cat output.tsv
                         SiteN1
                                 SiteN2
                                        SiteN1m SiteN2m SiteL1
                                                               SiteL2
                                                                       SiteF1
Seq. Nseq
              Р1
      1
                         3
                                 2
               0.6000
                                        2
                                                2
                                                        30
                                                               29
                                                                       0.8000
chr1
                                                        15
chr2 1
              0.3750
                         1
                                 Ω
                                        Ω
                                                \cap
                                                               \cap
                                                                       nan
                                        1.0000
                         2.0000
                                1.0000
Average 1.0000 0.4875
                                               1.0000 22.500
                                                               14.500
                                                                       0.8000
                                       2
     2
                      ... 4
                                2
                                                2
                                                        45
                                                               29
```

If the BED files contain strand information, the statistics for each strand separately can be calculated:

```
>cat test bed3.bed
chr1
     0
              10
                       featA
                               0
       15
chr1
               25
                       featB
                               0
chr1
       40
               50
                       featC
                               0
     12
chr2
              27
                       featD
                               0
>cat test bed4.bed
      0
               12
                       featE
chr1
       25
               42
                       featF
                               0
chr1
>python slalom.py --bed -a1 test bed3.bed -a2 test bed4.bed -s test seqlenfile2.tsv -o
output.tsv -c -d strand -sum --quiet
>cat output.tsv
       Nseq
               Ρ1
                          SiteN1 SiteN2
                                         SiteN1m SiteN2m SiteL1
                                                                 SiteL2
                                                                         SiteF1
Seq.
               0.4000
chr1+
       1
                          2
                                  1
                                          1
                                                  1
                                                          20
                                                                 12
                                                                         0.6667
                                          0
chr1-
       1
               0.2000
                          1
                                  1
                                                  0
                                                          10
                                                                 17
                                                                         nan
chr2- 1
               0.3750
                          1
                                  0
                                         0
                                                  0
                                                         15
                                                                 0
                                                                         nan
Average 1.0000 0.3250
                          1.3333 0.6667 0.3333
                                                 0.3333 15.000
                                                                 9.6667
                                                                        0.6667
```

Note that in case of strand or frame detection, the option of treating sequences as groups is activated automatically.

9. Averaging Approaches

In the previous two sections, one could have noticed that some metrics – for example, the site-wise F1 score – change their value merely on turning on and off a sequence grouping option. This is not a bug in SLALOM, but a consequence of differences between micro and macro averaging. With micro averaging, all the symbol and site counts are summed up before the performance metrics formulas are applied. With macro averaging, the formulas applied first and the results are simple-averaged to produce the displayed metrics. Averaging is applied twice – to the sequence data at the group level as well as to the group data at the dataset level – and is regulated through the option –a/--averaging. Providing value sequence will trigger macro averaging on both levels, while providing dataset will trigger micro averaging; the default value group sets micro averaging at the group-level and macro averaging at the dataset-level.

Thus, unless dataset-wise averaging is applied, the average value for a column is the simple average of the corresponding group-wide values (nans are ignored):

```
>python slalom.py --bed -a1 test bed3.bed -a2 test bed4.bed -s test seqlenfile2.tsv -o
output.tsv -c -d strand -sum --benchmarking --quiet
>cat output.tsv
Seq. Nseq
                SiteNB SiteNP SiteNPm SiteLB SiteLP SiteTPR SitePPV ...
                       1
                                            20
      1
                2
                              1 1
                                                   12
                                                          0.5000 1.0000
chr1+
               1
chr1- 1
                       1
                              0
                                     0
                                            10
                                                   17
                                                          0.0000 0.0000
             ... 1
                                                          0.0000 nan
                       0
                              0
                                     0
                                            15
                                                   0
chr2- 1
Average 1.0000 ... 1.3333 0.6667 0.3333 0.3333 15.000 9.6667 0.1667 0.5000
                                            45
```

In the benchmarking mode, the site-wise TPR (true positive rate; a.k.a. recall or sensitivity) is the ratio of the number of matched benchmark sites (SiteNBm) to the total number of benchmark sites (SiteNB); the site-wise PPV (positive predictive value; a.k.a. precision) is the ratio of the number of matched predicted sites (SiteNPm) to the total number of predicted sites (SiteNP). These numbers are calculated for each group and then averaged.

Under dataset-wise averaging, however, the respective calculations are performed on the summed-up numbers of sites:

```
>python slalom.py --bed -a1 test bed3.bed -a2 test bed4.bed -s test seqlenfile2.tsv -o
output.tsv -c -d strand -sum -b -a dataset --quiet
>cat output.tsv
Seq. Nseq ...
                SiteNB SiteNP SiteNPm SiteLB SiteLP SiteTPR SitePPV ...
chr1+
                     1
                             1
                                 1
                                           20
                                                  12
                                                        0.5000 1.0000
                                    0
                                                  17
chr1-
                              0
                                           10
                                                         0.0000 0.0000
chr2- 1
                1
                      0
                              0
                                    0
                                           15
                                                  0
                                                         0.0000 nan
Average 1.0000
             ... 1.3333 0.6667 0.3333 0.3333 15.000 9.6667 0.2500 0.5000
                       2
                              1
                                     1
                                           45
```

Only one site out of four was recalled for the whole dataset, and therefore the site-wise TPR is now 0.25. The site-wise PPV remains at 0.5, although it would also change for a different dataset.

Another aspect of averaging is sequence length adjustment. Depending on the underlying question, a site of length 5 in a sequence of length 100 may be weighted differently than another site of length 5 in a sequence of length 25. As a result, it may happen that, if only one of such two sites in a benchmark annotation was recalled/predicted correctly in another annotation, the user would like to see different values depending on which site exactly is predicted correctly. Under the default settings, the performance metrics are not adjusted for the sequence length, and therefore the results are equivalent for both situations:

```
>cat test seqlenfile3.tsv
seqA
       100
     25
seqB
>cat test anno benchmark.tsv
     21 25
seqA
seqB 6
                10
>cat test anno prediction1.tsv

\begin{array}{cccc}
    & - & - & 25 \\
    & 16 & 20 \\
\end{array}

seqA
seqB
>cat test anno prediction2.tsv
     41 45
seqA
       6
                 10
seqB
>python slalom.py -a1 test anno benchmark.tsv -a2 test anno prediction1.tsv -s
test seqlenfile3.tsv -o output.tsv -c -b --quiet
```

```
>cat output.tsv
                  FP FN
Nseq ... TP
                                    TPR
                                           PPV
                                                   SPC
                                                              SiteTPR SitePPV ...
                                 ...
         0.0400 0.0400 0.0400 ...
                                   0.5000 0.5000 0.9565
>python slalom.py -a1 test anno benchmark.tsv -a2 test anno prediction2.tsv -s
test seqlenfile3.tsv -o output.tsv -c -b --quiet
>cat output.tsv
                                ... TPR
       ... TP
                                                           ... SiteTPR SitePPV ...
Nseq
                  FΡ
                                            PPV
                                                    SPC
                         FN
       ... 0.0400 0.0400 0.0400 ... 0.5000 0.5000 0.9565 ... 0.5000 0.5000 ...
```

The output changes accordingly, if the option <code>-A/--adjust_for_seqlen</code> is provided. It acts like a variation of sequence-wise (macro) averaging at the group level. While by default the sequence lengths and TP (true positive), FP (false positive), etc. counts are summed-up for the whole group and then used to calculate the respective fractions (columns <code>TP</code>, <code>FP</code>, etc.), the fractions are calculated for each sequence and then simple-averaged when adjusting for sequence length:

```
>python slalom.py -a1 test anno benchmark.tsv -a2 test anno prediction1.tsv -s
test seqlenfile3.tsv -o output.tsv -c -b -A --quiet
>cat output.tsv
                 FP
                                 ... TPR
                                           PPV
                                                     SPC
                                                           ... SiteTPR SitePPV ...
       ... TP
                         FN
2
       \dots 0.0250 0.1000 0.1000 \dots 0.2000 0.2000 0.8857 \dots 0.5000 0.5000 \dots
>python slalom.py -a1 test anno benchmark.tsv -a2 test anno prediction2.tsv -s
test seqlenfile3.tsv -o output.tsv -c -b -A --quiet
>cat output.tsv
                                 ... TPR
       ... TP
                  FP
                          FN
                                           PPV
                                                     SPC
                                                            ... SiteTPR SitePPV ...
Nseq
     ... 0.1000 0.0250 0.0250 ... 0.8000 0.8000 0.9714 ... 0.5000 0.5000 ...
```

Note that this adjustment may have a quite significant influence on performance metrics, if the sequence lengths are very diverse. However, it affects only the symbol-wise measures and not the site-wise ones. Note also that the adjustment for sequence length differs from sequence-wise averaging:

```
>python slalom.py -a1 test_anno_benchmark.tsv -a2 test_anno_prediction1.tsv -s test_seqlenfile3.tsv -o output.tsv -c -b -a sequence --quiet 
>cat output.tsv 
Nseq ... TP FP FN ... TPR PPV SPC ... SiteTPR SitePPV ... 
2 ... 0.0400 0.0400 0.0400 ... 0.5000 0.5000 0.9737 ... 0.5000 0.5000 ...
```

10. Resolving Overlaps within the Annotations

In some annotation, the sites overlap within one annotation, for example:

```
>cat test anno1.tsv
seqA 6
          1.0
seqA 8
           12
seqA 9
           18
seqB 15
           25
>cat test anno2.tsv
          11
     5
seqA
     7
           10
seqA
           15
seqA
     9
     15
           19
seqB
seqB 21
           25
```

By default, all the sites are treated separately, even if they are overlapping or duplicating each other. Therefore, for example, in the first/benchmark annotation of the sequence seqA, if counted gross, there are 3 sites with the total length 20 symbols:

```
>python slalom.py -a1 test anno1.tsv -a2 test anno2.tsv -s test seglenfile1.tsv -o
output.tsv -sum -b -E gross -sg --quiet
>cat output.tsv
     SiteNB: Number of sites in the benchmark annotation
     SiteNP: Number of sites in the prediction annotation
#
     SiteNBm: Number of matched sites in the benchmark annotation
#
     SiteNPm: Number of matched sites in the prediction annotation
     SiteLB: Total length of sites in the benchmark annotation
     SiteLP: Total length of sites in the prediction annotation
                              SiteNB SiteNP SiteNBm SiteNPm SiteLB SiteLP
Seq.
                   TPR
        Nseq
                   0.8500
                                       3
                                                       3
                                                               20
                              3
                                               3
                                                                       18
                                                                                1.0000
seqA
        1
                                                                                1.0000
                                       2
                                                       2
        1
                   0.9091
                              1
                                               1
                                                               11
                                                                        10
seqB
                ...
                           ...
                   . _ _ _ _ _ _
                           ...
                ...
Average 1.0000
                                                                       14.000
                   0.8795
                              2.0000
                                       2.5000 2.0000 2.5000 15.500
                                                                                1.0000
                           ...
                ...
        2
                              4
                                               4
                                                                        28
                                                               31
```

If this is not desired behavior, the overlaps can be resolved. Resolving is performed for each annotation individually and is regulated through the options <code>-alr/--annolfile_resolve</code> and <code>-a2r/--anno2file_resolve</code> respectively. There are three alternatives to the default behavior.

First, all the overlapping sites can be merged prior to further calculations. For this, the value merge should be passed. The merging has the same affect on symbol-wise statistics as counting symbol-resolved. However, unlike the symbol-resolved mode, the explicit merging affects also the site-wise statistics (for sake of calculating symbol fractions, the displayed total site length is also affected by the operating mode; however, there is no impact on site-wise statistics, like TPR or PPV). Consecutively, there is only one site 6-18 of length 13 upon merging in the first annotation of seqA:

```
>python slalom.py -a1 test anno1.tsv -a2 test anno2.tsv -s test seqlenfile1.tsv -o
output.tsv -sum -c -b -sg --quiet
>cat output.tsv
                       ... SiteNB SiteNP SiteNBm SiteNPm SiteLB SiteLP
Seq. Nseq
                                                                     SiteTPR ...
                TPR
                0.7692
                          3
                                 3
                                         3 3 13
                                                              11
                                                                     1.0000
      1
seqA
                        ...
     1
                0.9091
                          1
                                 2
                                         1
                                                2
                                                       11
                                                              10
                                                                     1.0000
seqB
Average 1.0000
             ... 0.8392
                          2.0000
                                 2.5000 2.0000 2.5000 12.000 10.500
                                                                     1.0000
                        ... 4
                                 5
                                               5
                                        4
     2
                                                       24
>python slalom.py -a1 test anno1.tsv -a2 test anno2.tsv -s test seqlenfile1.tsv -o
output.tsv -sum -c -b -sg -a1r merge -a2r merge --quiet
>cat output.tsv
                          SiteNB SiteNP SiteNBm SiteNPm SiteLB SiteLP SiteTPR ...
    Nseq
                 TPR
              ...
                       ...
      1
                          1
                                 1
                                         1 1 13
                                                              11
                                                                     1.0000
seqA
                0.7692
                        ...
                0.9091
                                  2
                                                2
                          1
                                         1
                                                       11
                                                              10
                                                                     1.0000
seqB
     1
                                                                     1.0000
Average 1.0000
                 0.8392
                          1.0000
                                 1.5000 1.0000 1.5000 12.000
                                                              10.500
                          2
                                  3
                                         2
                                                3
                                                       24
```

Second, from each overlapping group, only the last site can be retained, while the others are discarded. For this, the value last should be passed. In this case, there is only the site 9-18 of length 10 is left:

```
>python slalom.py -a1 test anno1.tsv -a2 test anno2.tsv -s test seqlenfile1.tsv -o
output.tsv -sum -c -b -sg -a1r last -a2r last --quiet
>cat output.tsv
        Nseq
                   TPR
                               SiteNB
                                      SiteNP
                                               SiteNBm SiteNPm SiteLB
                                                                        SiteLP
                                                                                SiteTPR ...
Seq.
                           ...
                   0.7000
                              1
                                       1
                                               1
                                                       1
                                                                10
                                                                        7
                                                                                1.0000
seaA
                           ...
seqB
      1
                   0.9091
                               1
                                       2
                                               1
                                                       2
                                                                11
                                                                        10
                                                                                1.0000
                ...
                           ...
Average 1.0000
                   0.8045
                              1.0000 1.5000 1.0000 1.5000 10.500
                                                                        8.5000
                                                                                1.0000
                           ...
                               2
                                       3
                                                        3
                                                                21
                                                                        17
```

Finally, only the first sites of each overlapping group can be retained, by providing the value first. This time, only the site 6-10 of length 5 is retained:

```
>python slalom.py -a1 test anno1.tsv -a2 test anno2.tsv -s test seqlenfile1.tsv -o
output.tsv -sum -c -b -sg -a1r first -a2r first --quiet
>cat output.tsv
                TPR
Seq. Nseq
                          SiteNB SiteNP SiteNPm SiteLB SiteLP SiteTPR ...
                       ... 1 1
     1
             ... 1.0000
                                        1
                                               1
                                                      5
                                                             7
                                                                    1.0000
seqA
seqB 1
                                        1
             ... 0.9091 ... 1
                                 2
                                               2
                                                      11
                                                             10
                                                                    1.0000
             ... 0.9545 ... 1.0000 1.5000 1.0000 1.5000 8.0000
Average 1.0000
                                                             8.5000 1.0000
                                                                           ...
     2
                       ... 2
                                               3
                                                      16
                                                             17
```

Note that there are no overlapping sites in the sequence seqB. Consecutively, its metrics are affected neither by overlap resolving nor by mode switching. Note also that after any of three kinds of resolving there are no overlapping sites left. Therefore, both the symbol-resolved and gross modes will produce equivalent results upon resolving.

11. Overlap Criteria between the Annotations

Overlap criteria are of crucial importance for calculating site-wise metrics. By default, any overlap – even a single symbol – is sufficient to count a pair of sites from different annotations as matched. This is reason, why the site-wise TPR always remained 1.0 in the previous section: regardless of which sites were retained, there was always an overlap. However, often a more sophisticated approach is needed.

Consider the following two annotations:

The site 2-6 from the first/benchmark annotation corresponds to the site 3-7 from the second/prediction annotation fairly good: the overlap is 80% (but which still may be not good enough depending on the context). The site 20-22 is completely covered; however, the partner 16-30 is much longer: 3 symbols vs. 15. Such situations are typical for weak predictors, which are trying to capture the benchmark sites just by chance. Such a match may be considered good in some context but not in the other. The site 37-47 is covered on 10 symbols out of 11; however, the coverage is in two patches of length 5 each. Again, it is not possible to say, if this site is recalled without knowing the context and the specific question asked about the data.

SLALOM offers considerable flexibility while considering site matching criteria by allowing to set several types of constraints on overlap of putatively matched sites.

The easiest constraint is the minimal number of symbols in the overlap. The desired value should be passed with the option <code>-Os/--overlap_symbols</code>. In this example, at least 4 symbols are required:

```
>python slalom.py -a1 test anno5.tsv -a2 test anno6.tsv -1 50 --single sequence -o
output.tsv -b -Os 4 --quiet
>cat output.tsv
     SiteNB: Number of sites in the benchmark annotation
#
    SiteNP: Number of sites in the prediction annotation
    SiteNBm: Number of matched sites in the benchmark annotation
#
    SiteNPm: Number of matched sites in the prediction annotation
#
                          ... SiteNB SiteNP SiteNBm SiteNPm ... SiteTPR SitePPV ...
                  PPV
         TPR
Nseq
       ... 0.8947 0.5484 ... 3
                                     4
                                            2
                                                3 ... 0.6667 0.7500 ...
```

For the benchmark, the sites 2-6 and 37-47 are matched but not the site 20-22: although it is completely covered, it is only 3 symbols long and therefore can never match with the constraint of at least 4 symbols in overlap. This leads to the site-wise recall of 2/3. For the prediction, all the sites except for 16-30 got their overlap of at least 4 symbols. This leads to the site-wise precision of 3/4.

Another type of constraint is the minimal part of a site to overlap. The desired value should be passed with the option <code>-op/--overlap_part</code> as a fraction (not percentage). In this example, at least 80% overlap is required:

This time, the prediction is reported site-wise perfect. This is because by default SLALOM calculates overlapping parts from the shortest of the two sites. Thus, the sites 20-22 and 16-30 score 100% overlap, although it would be significantly less of the length of the latter. This is the most permissive approach of applying the overlapping constraints.

To change the way, how the constraints are applied, the user should provide the option -Oa/--overlap apply with the corresponding value. There are three alternative values to shortest.

The most rigorous approach is applying to the longer site of the two, by providing the value longest:

This time, only the sites 2-6 and 3-7 could match. The site 20-22 was not considered, because the overlap part was calculated by the longer partner 16-30: it is now only 3/15, or 20%.

When applying to either the shorter or the longer site, matches are always symmetric. However, it is not the case, when applying to the site of the annotation currently considered, by providing the value current:

Now, two of the sites from the benchmark annotation got their match: 20-22 was considered but 37-47 still not. This is because the latter may be covered to 80% but not with a single site. For each of the sites 37-41 and 43-48 the overlap is only 5 symbols, or 45.45%. For the prediction, however, the situation was the opposite: no match registered in the pair 20-22 and 16-30 but instead both 37-41 and 43-48 passed the 80% threshold. Note that although the recall and precision have the same value as in the example with the requirement of 4 symbols, the contributing sites are now different.

Finally, if patched coverage – the coverage of one site by many – is also good, the value patched should be provided:

This time, the site 37-47 was also matched by satisfying the 80% threshold with the total coverage of 10 symbols from two sites contributing 5 symbols each.

In addition to setting constraints on overlap length, one may also consider the order of sites in a putative pair. Sometimes, additional information about the relation between sites in the annotations is available, which makes only those site pairs meaningful, in which the predicted site appears earlier or later than the benchmark one. To impose such constraints, the user can use the option <code>-on/--overlap_nature</code> with the value either <code>leading</code> or <code>lagging</code>. The option is compatible only with the benchmarking mode; a leading prediction can start no later than the benchmark, and vice versa for the lagging one:

If the prediction is leading, the sites 2-6 and 3-7 can no longer match, even if they overlap sufficiently, because the latter begins later than the former. Similarly, if the prediction is declared lagging, the sites 20-22 and 16-30, as well as 37-47 and 43-48 cannot match, because the predicted site begins earlier.

The three constraints can be applied simultaneously; the AND logic will be used.

Note that the three discussed types of constraints do affect only the site-wise metrics and do not influence residue-wise ones.

12. Detailed, Site-wise Statistics as Output

In previous sections of this manual, it was demonstrated that the main output file contains statistics at the group or whole dataset levels. However, in some situations, information on each annotated site is desirable. This may be useful for two purposes. First, the user may wand to understand, how performance metrics on the group and dataset levels arise or why the performance is worse/better

than expected. Second, some sort of mapping between sites in different annotations may be required. Mapping is discussed in more detail in the next section.

SLALOM can provide site-wise statistics in two different output files.

The first file is called 'detailed output'. It is a plain text file and represents a log of SLALOM operations, as it works through the input annotations. The filename should be provided with the option -od/--outfile detailed. Using an example from the previous section:

```
>python slalom.py -a1 test_anno5.tsv -a2 test_anno6.tsv -l 50 --single_sequence -o
output.tsv -od output detailed.txt -b -Op 0.8 -Oa current -c --quiet
>cat output_detailed.txt
Information on the unnamed sequence (length 50 symbols):
    17 symbols are present in both annotations (true positives)
    2 symbols are present exclusively in the benchmark (false negatives)
    14 symbols are present exclusively in the prediction (false positives)
    17 symbols are absent in both annotations (true negatives)
    Site 2-6 of the benchmark: overlaps with site 3-7 of the prediction by 4 symbols
(80% and 80% of the site lengths respectively)
    Site 20-22 of the benchmark: overlaps with site 16-30 of the prediction by 3
symbols (100% and 20% of the site lengths respectively)
    Site 37-47 of the benchmark: no sufficient overlap found
    Site 3-7 of the prediction: overlaps with site 2-6 of the benchmark by 4 symbols
(80% and 80% of the site lengths respectively)
    Site 16-30 of the prediction: no sufficient overlap found
    Site 37-41 of the prediction: overlaps with site 37-47 of the benchmark by 5
symbols (100% and 45% of the site lengths respectively)
    Site 43-48 of the prediction: overlaps with site 37-47 of the benchmark by 5
symbols (83% and 45% of the site lengths respectively)
    There are 3 sites in the benchmark with total length 19 unique symbols
    2 benchmark sites are matched in the benchmark
    1 benchmark site has no match in the benchmark
    There are 4 sites in the prediction with total length 31 unique symbols
    3 prediction sites are matched in the benchmark
    1 prediction site has no match in the benchmark
```

The second file contains essentially the same information, but in a tabular form (the TSV format). It lists all the sites, one site per row. The column names are the following: sequence (SID; empty, if single sequence without an SID is processed), Annotation (b for the benchmark, p for the prediction; in the equal mode: 1 and 2), Site begin, Site end, Overlapped symbols, Overlapped perc., Partner overlapped perc., Partner begin, Partner end. They comprise the header row. The filename should be provided with the option -os/--outfile_sites:

```
>python slalom.py -a1 test anno5.tsv -a2 test anno6.tsv -1 50 --single sequence -o
output.tsv -os output sitewise.tsv -b -Op 0.8 -Oa current -c --quiet
>cat output sitewise.tsv
                                4
                                        80
                                                80
       b
                        6
       b
               20
                        22
                                3
                                        100
                                                20
                                                       16
                                                                30
       b
                37
                        47
                               0
                                       0
                                                0
                                               80
                                                        2
                3
                        7
                               4
                                       80
                                                                6
       р
                        30
               16
                               0
                                       0
                                               0
       р
                                                       37
               37
                        41
                                5
                                       100
                                               45
                                                               47
       р
                                                       37
               43
                       48
                                5
                                               45
                                       83
                                                                47
       р
```

13. Mapping of Site Names/Identifiers between the Annotations

Sometimes, the input annotations contain the same sites (for example, genes in a certain organism), which, however, are derived from different sources with each source providing its own version of the site names (a.k.a. identifiers or accessions). Matching the names in such a situation is known as mapping. As the sites do not necessarily match perfectly between the annotations, the mapping is often ambiguous. Nevertheless, a reasonably good mapping can be produced, if the data quality is good enough and overlapping criteria are set wisely.

Consider the following annotations (assuming both sequences have the length 150):

```
>cat test anno7.tsv
         40
seqA 21
                geneA
seqA 61
           85
                 geneB
seqA 101
           130
                geneC
seqB 16
           30
                geneD
>cat test_anno8.tsv
         40
seqA 21
                 gene1
           88
                 gene2
seqA 61
          150
seqA 121
                gene3
```

In them, geneA and gene1 match perfectly, geneB and gene2 match reasonably, geneC and gene3 overlap only modestly, while geneD cannot be matched in the second annotation at all. To read in also the site names and consequently show them in the site-wise output, the user should provide the option $-n/-site_names$. The overlap quality criterion can be, for example, at least 90% of the current gene:

```
>python slalom.py -a1 test anno7.tsv -a2 test anno8.tsv -1 150 -o output.tsv -os
output mapping.tsv -n -Op 0.9 -Oa current -c --quiet
>cat output mapping.tsv
seqA
        1
                 21
                         40
                                  geneA
                                           20
                                                   100
                                                            100
                                                                             40
                                                                                      gene1
seqA
        1
                 61
                         85
                                  geneB
                                           25
                                                   100
                                                            89
                                                                    61
                                                                             88
                                                                                      gene2
        1
                 101
                         130
                                  geneC
                                           0
                                                   0
                                                            0
seqA
        2
                 21
                          40
                                  gene1
                                           20
                                                   100
                                                            100
                                                                    21
                                                                             40
                                                                                      geneA
seqA
        2
                 61
                          88
                                  gene2
                                          0
                                                   0
                                                            0
seqA
        2
                 121
                         150
                                           0
                                                   0
                                                            0
                                  gene3
seqA
                          30
                 16
                                  geneD
seqB
```

In comparison to the case, when no site names are provided, the table has two additional columns: column 5 (Site name) and column 11 (Partner name). By default, the mapping lists all the sites from the input annotations. For sites without a match, the partner name is empty. Nevertheless, the user can limit the output by setting the option <code>-osd/--outfile_sites_diff</code> to either <code>matched</code> or <code>discrepant</code> (the latter will list all but perfect matches, regardless of the overlapping criteria set):

```
>python slalom.py -a1 test anno7.tsv -a2 test anno8.tsv -1 150 -o output.tsv -os
output_mapping.tsv -osd matched -n -Op 0.9 -Oa current -c --quiet
>cat output mapping.tsv
        1
                 21
                         40
                                  geneA
                                          20
                                                   100
                                                           100
                                                                    21
                                                                            40
seqA
                                                                                     gene1
        1
                 61
                         85
                                          25
                                                   100
                                                           89
                                                                    61
                                                                            88
seqA
                                  geneB
                                                                                     gene2
        2
                21
                         40
                                          20
                                                  100
                                                           100
seqA
                                  gene1
                                                                    21
                                                                            40
                                                                                     geneA
>python slalom.py -a1 test_anno7.tsv -a2 test_anno8.tsv -l 150 -o output.tsv -os
output mapping.tsv -osd unmatched -n -Op 0.9 -Oa current -c --quiet
>cat output mapping.tsv
        1
                 101
                         130
                                  geneC
                                          0
                                                   0
                                                           0
seaA
                                  gene2
                                                   0
seqA
        2
                 61
                         88
                                          0
                                                           0
        2
                                                   0
                                                           0
seqA
                 121
                         150
                                  gene3
                                          0
seqB
        1
                 16
                         30
                                  geneD
                                                   0
                                                           0
```

```
>python slalom.py -a1 test anno7.tsv -a2 test anno8.tsv -1 150 -o output.tsv -os
output mapping.tsv -osd discrepant -n -Op 0.9 -Oa current -c --quiet
>cat output mapping.tsv
        1
                 61
                         85
                                  geneB
                                          25
                                                   100
                                                            89
                                                                    61
                                                                             88
                                                                                     gene2
seqA
        1
                 101
                         130
                                          0
                                                   0
                                                            0
                                  geneC
seqA
        2
                 61
                         88
                                          0
                                                   0
                                                            0
seqA
                                  gene2
                                  gene3
        2
                 121
                         150
                                          0
                                                   0
                                                            0
seqA
                                                   0
                                                            0
                 16
                         30
                                          0
seqB
                                  geneD
```

Discrepant and unmatched sites are important, if one wants to concentrate on differences: for example, when comparing two versions of the same genome annotation.

Note that in the simplified modes (GenBank or BED) the decision on reading the site name is done automatically on the basis of the structure of the input files.

SLALOM can also perform mapping of elements that do not overlap at all. This becomes useful, for example, if one wants to associate genes with other genomic elements, like, e.g., promoters. If these are known to precede the associated genes locating to a window of certain length before a gene start position, the start positions can be imaginary 'shifted' in order to register an 'overlap'. The desirable amount of symbols should be provided with the respective option <code>-albs/--annolfile_begin_shift</code> or <code>-albs/--annolfile_begin_shift</code>. As the shift in the direction of the sequence start is needed, a negative value should be provided:

```
>cat test anno9.tsv
seqA 16
         19
                  elemA
seqA 53
            57
                 elemB
seqA 95
            98
                 elemC
seqB 7
          12
                 elemD
seqB 61
          64
                 elemE
>python slalom.py -a1 test anno7.tsv -a2 test anno9.tsv -l 150 -o output.tsv -os
output mapping.tsv -n -albs -10 -c --quiet
>cat output mapping.tsv
        1
                11
                         40
                                         4
                                                 13
                                                          100
                                                                  16
                                                                          19
                                 geneA
                                                                                  elemA
seqA
       1
                51
                        85
                                 geneB
                                         5
                                                 14
                                                         100
                                                                  53
                                                                          57
                                                                                  elemB
seqA
       1
                91
                        130
                                 geneC
                                         4
                                                 10
                                                         100
                                                                  95
                                                                          98
                                                                                  elemC
seqA
        2
                16
                        19
                                 elemA
                                         4
                                                 100
                                                         13
                                                                  11
                                                                          40
seqA
                                                                                  geneA
        2
                53
                        57
                                         5
                                                 100
                                                         14
                                                                  51
                                                                          85
                                 elemB
seqA
                                                                                  geneB
        2
                95
                         98
                                 elemC
                                                         10
                                                                  91
seqA
                                         4
                                                 100
                                                                          130
                                                                                  geneC
        1
                         30
                                                 24
                                                         100
                                                                  7
seqB
                6
                                         6
                                                                          12
                                 geneD
                                                                                  elemD
        2
                7
                        12
                                                 100
seqB
                                         6
                                                          2.4
                                                                  6
                                                                          30
                                 elemD
                                                                                  geneD
        2
                61
seqB
                        64
                                 elemE
                                                 0
                                                          0
```

The same effect can be also reached by extending the elements in the second annotation through the similar option -a2es/--anno2file_end_shift.

In this case, the outputted overlapping percentages do not make much sense any longer, as the 'gene length' was artificially changed. However, the shift options can be also used to correct the input: for example, if symbol counting starts from 0 instead from 1, the value 1 should be passed to adjusting the sites to the SLALOM conventions. Note that 1 is automatically added to the read start positions in the BED simplified mode; independently of the provided shift values.

14. Circular Sequences as Input

In some areas of science, like, e.g., bacterial genomics, circular sequences are very common.

A usual approach to annotate such a sequence is to introduce an imaginary cut at some position, thus converting it to a linear sequence. Working with a circular sequence is therefore the same, as long as the cut does not go through an annotated site. If that happens, however, the cut-through site must be still represented as a single entity. To achieve this, one normally marks the end position with a symbol numbers that are greater than the sequence length. Alternatively, one can a use non-positive start symbol number.

Let us consider the following annotations of one site each in a sequence of length 50. The last annotations actually refer to the same site but with different notation:

```
>cat test_anno_circular1.tsv
1    10
>cat test_anno_circular2.tsv
48    59
>cat test_anno_circular3.tsv
-2    9
```

By default, SLALOM considers input sequences linear and raises an error, when a symbol number out of scope of the sequence is detected:

```
>python slalom.py -a1 test_anno_circular1.tsv -a2 test_anno_circular2.tsv -l 50 -- single_sequence -o output.tsv -b -c --quiet
Error: Error while parsing the line 1 of the file "test_anno_circular2.tsv". Site end position cannot exceed the sequence length
```

To treat the input as circular, the user should provide the value circular with the option -e/-- end overflow policy:

```
>python slalom.py -a1 test anno circular1.tsv -a2 test anno circular2.tsv -1 50 --
single sequence -o output.tsv -od output detailed.txt -e circular -b -c --quiet
>cat output.tsv
                 PPV
Nseq ... TPR
                       ... SiteNB SiteNP SiteNBm SiteNPm SiteLB SiteLP ...
       ... 0.9000 0.7500 ... 1 1 1 1 1
>cat output detailed.txt
   Site 1-10 of the benchmark: overlaps with site 48-59 of the prediction by 9 symbols
(90% and 75% of the site lengths respectively)
   Site 48-59 of the prediction: overlaps with site 1-10 of the benchmark by 9 symbols
(75% and 90% of the site lengths respectively)
>python slalom.py -a1 test_anno circular1.tsv -a2 test anno circular3.tsv -1 50 --
single_sequence -o output.tsv -e circular -b -c --quiet
>cat output.tsv
                PPV ... SiteNB SiteNP SiteNBm SiteNPm SiteLB SiteLP ...
Nseq ... TPR
      ... 0.9000 0.7500 ... 1 1 1 1 1 10 12 ...
```

The overlap of the two sites and performance metrics are calculated as usual, as if there were no cut through the second one.

15. Custom Tabular Files as Input

In the earlier sections of this manual, it was shown that SLALOM can process input files in some predefined formats without further specifications, as well as tabular files, which satisfy certain conditions. These conditions are the following:

- The column delimiter is tab
- The columns are following the order SID GID start end name (non-specified columns omitted) for the annotation files, SID length for the sequence length table, and SID GID for the group mapping file
- There are no further columns before or between the specified ones (additional columns to the right will be ignored)
- There are no header rows
- Quotes are not part of the identifiers and names (although tabs may be if quoted)

Therefore, the user always has an option to convert the input files in one of the supported formats to process them with SLALOM in an easy way. However, this is not always convenient, as there are plenty of tabular data files – available for download or outputted by software tools – that do not strictly conform to requirements of the supported format. Furthermore, the user may encounter situations, when the annotations come from different sources and therefore have different formats. To also offer flexibility for these cases, SLALOM has extended options allowing it to read virtually any kind of tabular data files.

The format options are combinations of prefixes and suffixes. The prefixes correspond to the file name options: -a1 for the first/benchmark annotation, etc. The suffixes are given in the following table:

Short version	Long version	Description
d	delim	Specifies the column delimiter (the following delimiters are
		supported: tab, space, comma, dot, semicolon, colon, slash).
		Consecutive delimiters mark empty values, unless empty string is
		passed as delimiter. In that case, space is used but with leading and
		trailing spaces removed and consecutive spaces collapsed
С	colnumbers	Specifies the column numbers in the following order: start - end -
		SID - GID - name for the annotation files; SID - length (SID - start -
		end for time series) for the sequence length table; SID - GID for the
		group mapping
h	headers	Specifies the number of header rows at the beginning of the file
d	quotes	Triggers the literal treatment of quotes (quoted delimiters are
		treated as delimiters; single – but not double – quotes can be part of
		identifiers and names)

A prefix is combined with a suffix to form one of the 16 possible options. For the long options, the extra underscore is added as separator. Therefore, to specify, for example, that the second annotation file has three header rows that need to be skipped, the user should provide the value 3 with the option $-a2h/--anno2file_headers$ (e.g., -a2h 3). The column numbers are integers, so that the leftmost column has the number 1. They must be provided comma-separated without additional spaces. For example, if the sequence lengths are read from a file that contains the identifiers in the fifth column and the length values in the third column, the user should provide the value 5,3 with the option $-sc/-seglenfile_colnumbers$ (e.g., -sc 5,3).

This approach allows that same files and/or same columns may be used several times if the input format requires so. In the following example, the sequence lengths and the group mapping are read from the annotation files:

```
>cat test format1.txt
Number; ID; Quality; Score; Start; End
1; ABC; Good; 0.76; 17; 55
2; ABC; Good; 1.02; 58; 70
3; DEF; Good; 1.21; 12; 28
4; XYZ; Bad; 0.18; 101; 108
>cat test_format2.csv
\# This file was generated by ...
Origin, Sequence, Length, Peak Value, Average value, Start, End
Natural, ABC, 100, 8.4, 2.0, 48, 62
Artif., DEF, 120, 9.1, 1.9, 26, 40
Artif., DEF, 120, 6.2, 0.9, 69, 85
Artif., XYZ, 150, 11.5, 3.1, 108, 122
>python slalom.py -a1 test format1.txt -a1h 1 -a1c 5,6,2 -a1d ";" -a2 test format2.csv
-a2h 2 -a2c 6,7,2 -a2d "," -s test format2.csv -sh 2 -sc 2,3 -sd "," -m
test format1.txt -mh 1 -mc 2,3 -md ";" -o output.tsv -nOg -c --quiet
>cat output.tsv
Group Nseq P1
                        P2
                                   SiteN1 SiteN2 SiteN1m SiteN2m SiteL1
                                                                            SiteL2
      2
                0.3136 0.2136
                                   3 3
                                                    3 2 69
                                                                             47
Good
                0.0533 0.1000
                                            1
                                                    1
                                                            1
                                                                    8
                                                                             15
       1
                                   1
Average 1.5000 0.1835 0.1568 ... 2.0000 2.0000 2.0000 1.5000 38.500
                                                                            31.000 ...
```

For the first calculation, the column <code>Quality</code> of the file <code>test_format1.txt</code> was used for grouping the sequences. Therefore, the file was provided twice: as the first annotation (<code>-a1 test_format1.txt</code>) and as the group mapping (<code>-m test_format1.txt</code>); the file <code>test_format2.csv</code> was also provided twice: as the second annotation (<code>-a2 test_format2.csv</code>) and as the sequence length table (<code>-s test_format2.csv</code>). The column <code>Quality</code> in the former file has number 3, while the SiDs are in the column <code>ID</code> with the number 2, which results in <code>-mc 2, 3</code>. Providing the option <code>-nog/-non_overlapping_groups</code> allows for omitting the GIDs in the annotation files (in fact, it would be not possible to specify these GIDs in the second annotation file, as the quality information is missing there). Therefore, only three columns are read from each of the annotations: start position, end position, and SID. In the file <code>test_format1.txt</code>, the start positions are in column number 5, the end positions are in column number 6, and the SIDs are in column number 2, which results in <code>-alc 5, 6, 2</code>. Following the same logic, the option for the second annotation is <code>-a2c 6, 7, 2</code>. The sequence length values are provided in the same file in column number 3, which results in <code>-sc 2, 3</code>.

The next two calculations are conducted in a similar way:

```
>python slalom.py -a1 test format1.txt -a1h 1 -a1c 5,6,2 -a1d ";" -a2 test format2.csv
-a2h 2 -a2c 6,7,2 -a2d "," -s test format2.csv -sh 2 -sc 2,3 -sd "," -m
test format2.csv -mh 2 -mc 2,1 -md "," -o output.tsv -nOg -c --quiet
>cat output.tsv
                                  SiteN1 SiteN2 SiteN1m SiteN2m SiteL1
Group Nseq P1
                       P2
                                                                         SiteL2
                               ...
              0.5200 0.1500
                                                                 52
                                                  2
Natural 1
                                  2
                                          1
                                                     1
                                                                         15
Artif. 2
               0.0926 0.1741
                                          3
                                                  2
                                  2
                                                          2
                                                                  25
                                                                         47
Average 1.5000 0.3063 0.1620
                                 2.0000 2.0000 2.0000 1.5000 38.500 31.000
>python slalom.py -a1 test_format1.txt -a1h 1 -a1c 5,6,2 -a1d ";" -a2 test_format2.csv
-a2h 2 -a2c 6,7,2 -a2d "," -s test format2.csv -sh 2 -sc 2,3 -sd "," -o output.tsv --
sequences as groups -c --quiet
>cat output.tsv
                                  SiteN1 SiteN2 SiteN1m SiteN2m SiteL1
     Nseq
               Р1
                       Ρ2
                                                                         SiteL2
Seq.
                               ...
               0.5200 0.1500
                                                  2
                                                                 52
ABC
       1
                                  2
                                          1
                                                       1
                                                                         15
                                          2
                                                                 17
DEF
               0.1417
                       0.2667
                                  1
                                                  1
                                                          1
                                                                         32
                               ...
                                                                                 ...
               0.0533
                       0.1000
                                  1
                                          1
                                                  1
                                                          1
                                                                  8
                                                                         15
                       0.1722 ...
Average 1.0000 0.2383
                                 1.3333 1.3333 1.3333 1.0000 25.667 20.667 ...
```

16. Time Series as Input

SLALOM is a generic statistical tool. Although it was originally developed to work with SCEs in protein and DNA sequences, its abstract design allows applying it to virtually any kind of sequences, in biology and beyond.

Working with time series follows the same logic as analyzing the ranges in alphabetical sequences. Indeed, time intervals can be viewed as SCEs in the sequences of time units. For processing time series, SLALOM is started with providing the option <code>-t/--time_unit</code>. The four following time units with the corresponding values are supported: second (<code>sec</code>), minute (<code>min</code>), hour (<code>hour</code>), and day (<code>day</code>). Now, the start and end positions in the annotation files are replaced with time stamps. For the sequence length table, the length values are replaced with pairs of sequence start and end time stamps (so that the sequence length table now contains three columns instead of two). The following two date formats are supported: <code>dd.mm.yyyy</code> and <code>mm/dd/yyyy</code>. The optional time is expected after the space in the format <code>hh:MM</code> or <code>hh:MM:SS</code>; <code>AM/PM</code> classification is currently not supported. If the time is not specified, <code>00:00:00:00</code> is assumed.

After the internal conversion of time intervals to symbols, the workflow and the output are the same as for a general SLALOM case. Time sequences as well as individual intervals can also overlap and the usual rules for overlap resolving will be applied:

```
>cat test time lines.tsv
PlaceA 02/01/2015 22:45
                              02/02/2015 09:15
PlaceB 02/02/2015 02:10
                             02/03/2015 00:45
PlaceC 02/03/2015 17:05
                             02/03/2015 22:10
>cat test anno events1.tsv
PlaceA 02/01/2015 23:05
PlaceA 02/02/2015 03:10
PlaceB 02/02/2015 03:15
                             02/01/2015 23:30
                             02/02/2015 04:00
                             02/02/2015 04:05
>cat test anno events2.tsv
PlaceA 02/01/2015 23:10
                             02/01/2015 23:40
PlaceB 02/02/2015 03:15
                             02/02/2015 04:10
PlaceB 02/02/2015 03:45
                             02/02/2015 05:00
PlaceC 02/03/2015 18:35
                             02/03/2015 19:10
>python slalom.py -t min -s test time lines.tsv -a1 test anno events1.tsv -a2
test anno events2.tsv -o output.tsv --sequences as groups -sum -c --quiet
>cat output.tsv
                             ... SiteN1 SiteN2 SiteN1m SiteN2m SiteL1 SiteL2
                 P2
Seq. Nseq P1
PlaceA 1
                                        1
                                                                     30
              0.1190 0.0476
                                2
                                               1 1 75
                                                       2
             0.0369 0.0775 ... 1
                                        2
                                                              50
                                                                     105
PlaceB 1
                                               1
PlaceC 1 0.0000 0.1148 ... 0
                                                      0
                                               0
                                        1
                                                              0
                                                                      35
Average 1.0000 0.0520 0.0800
                               1.0000 1.3333 0.6667 1.0000 41.667
                                                                      56.667
                            ... 3
                                            2
                                                   3
                                                              125
     3
                                    4
                                                                      170
```

Sequence grouping and all the previously discussed associated option work also for time series.

17. Summary of Available Command Line Options for SLALOM

Туре	Default	Description	
Main input/output files			
-s,seqlenfile			
string	(empty)	Sequence length table file. Maps SIDs to corresponding sequence lengths (or, in case of time series, start and finish points). Must be in	

		tabular format. The file shall not be provided, if the length of all sequences is specified elsewhere
-m,mapfile		
string	(empty)	Mapping file. Maps GIDs and SIDs. Must be in tabular format
-a1,anno1f	ile	
string	(required)	First/benchmark annotation file. Lists all sites from the first annotation, one site per line. Must be in tabular format, unless genbank is specified
-a2,anno2f	ile	
string	(required)	Second (prediction) annotation file. See the option -a1 for details
-o,outfile		
string	(required)	Output TSV file with calculated performance measures, for each group and dataset-wide
		Simplified modes
genbank		5p
bool	false	Read the annotations in GenBank format. Sequence length will be read automatically
bed		
bool	false	Read the annotations in BED format
		Operation mode setup
-b,benchma	rking	
bool	false	Activate the benchmarking mode, i.e., treat the first annotation as benchmark and the second as prediction (default: treat the annotations symmetrically)
-E,enrichm	ent_count	
integer or string	0	Switch between symbol-resolved, gross, and enrichment modes. Must be a non-negative integer or gross. If 0, activates the symbol- resolved mode. If >=1, activates the enrichment mode with minimal number of occurrences of a symbol position to count it as enriched. If gross, activates the gross mode
		Core algorithm controls
-Os,overla	p_symbols	
integer	1	Minimal number of symbols required for a site from the annotation currently considered to be overlapping with a site/sites from the other annotation to be counted as a match. Must be a positive integer
-Op,overla	p_part	.,
float	0.0	Minimal fraction of the length of a site from the annotation currently considered required to overlap with site/sites from the other annotation to be counted as a match. Must be in range [0,1]
-Oa,overla	p_apply	
enum	shortest	The principle to apply the minimal number of symbols and the minimal part: - shortest: to the shortest of two sites (one site from the first annotation and the other one from the second) - longest: to the longest of two sites - current: to the site from the annotation currently considered; sites from another annotation are considered one at a time

	T	- patched: to the sites from the annotation currently
		considered; allowing multiple sites from the other annotation
		to contribute simultaneously
-On,overla	p nature	,
enum	neutral	The required overlap nature of the predicted sites:
Cituin		- neutral: count site overlaps regardless of the order of their start positions
		- leading: only count a predicted site as match, if its start
		position is earlier or the same as that of the respective
		benchmark site
		 lagging: only count predicted site as match, if the begin position is later or the same as that of the respective benchmark site
		The default value can be changed only in the benchmarking mode
		and only for non-circular sequences
-a,averagi	ng	
enum	group	Order of averaging when calculating performance measures: - sequence: calculate the measures for each sequence individually, then simple-average group-wide, then simple-
		average dataset-wide (macro-macro averaging)
		- group: sum the counts group-wide, then calculate the
		measures, then simple-average dataset-wide (micro-macro averaging)
		- dataset: sum the counts dataset-wide and calculate the
		measures (micro-micro averaging); for results on individual
		groups, apply macro averaging
-A,adjust_	for_seqlen	
bool	false	Adjust the counts for each sequence for its length, then average the adjusted values (default: sum the counts). Affects only symbol-wise
		measures. Not compatible with sequence-wide averaging.
		Input file format
-sd,seqlen	T	y
char	tab	Column delimiter in the sequence length table. Allowed delimiters:
		space, tab, comma, dot, colon, semicolon, slash. If an empty string is
		passed, space will be used, multiple spaces will be collapsed, and
md	0 401	leading, as well as trailing spaces will be removed removed
-md,mapfil	_	
-a1d,anno1	_	
-a2d,anno2		
(see the option -s		
-sh,seqlenfil	T	Manches Charles and Assessment Charles and As
integer	0	Number of header rows to skip in the sequence length table. Must be
1 6'3	1 1	a non-negative integer
-mh,mapfil	_	
-a1h,anno1	_	
-a2h,anno2		rs
(see the option -s	_	
-sc,seqlen		,
list of integers	(adjusted)	Comma-delimited list of column numbers (left-most column has

		number 1) in the sequence length table with SIDs and sequence length in that order. In case of time series, start and finish points
		must be provided instead of the length. The default value is adjusted according to the expected columns
-mc,mapfil	e colnumbe	rs
list of integers	1,2	Comma-delimited list of column numbers (left-most column has number 1) in the group-mapping file with SIDs and GIDs in that order
-a1c,anno1	file_colnu	mbers
list of integers	(adjusted)	Comma-delimited list of column numbers (left-most column has number 1) in the first annotation file with site start positions, site end positions, SIDs, GIDs, and site names in that order (with those not provided skipped). The default value is adjusted according to the expected columns
-a2c,anno2		mbers
(see the option -a		
-sq,seqlen		
bool	false	Treat single quotes (apostrophes) in the sequence length table file literally. By default, delimiters inside quotes (both single and double) are treated as belonging to fields and quotes themselves are ignored. Double quotes can never be part of read input; their presence in the file triggers an error, if this option is true
-mq,mapfil	e quotes	7 1
-a1q,anno1	_	.s
-a2q,anno2	_	
(see the option -s		
(CCC SEC SPICE	-19	Alternative input options
-1,seglen v	alue	•
integer	0	Length of all sequences. Must be a non-negative integer. If 0, this option is ignored. If positive, the sequence length table shall not be provided; the length of every sequence is set to the specified value; all SIDs encountered in the annotations and the group-mapping file will be used to build the sequence length table. With time series options -ts and -tf shall be used instead
-ss,single_	sequence	
bool	false	Process single sequence. SIDs shall not be provided if this option is activated
-ts,timeser	ies_start	-
string	(empty)	Start of all time series. If empty, this option is ignored. Must be used together with the option -tf. For details, see the option -1
-tf,timeser	ies_finish	1
string	(empty)	Finish of all time series. For details, see the option -ts
-alas,annol	false	Consider all the sites from the first annotation as belonging to all the sequences. SIDs shall not be provided if this option is activated. This option cannot be selected for both annotations simultaneously
-a2as,anno2 (see the option -a		sequences
-a1ag,anno2		roins
aray,aiii102	.TTTE	lr orbs

bool	false	Consider all the sites from the first annotation as belonging to sequences with provided SIDs in all the groups. If this option is activated, GIDs shall not be provided, although the group mapping must be provided. This option can be selected for both annotations, but it is not compatible with considering all the sites as belonging to all the sequences
-a2ag,anno2 (see the option -a		groups
-sg,sequenc	es as grou	ıps
bool	false	Generate respective 1-sequence groups for all the SIDs. If this option is activated, neither the group-mapping file nor GIDs shall be provided
-nOg,non_ov	erlapping	groups
bool	false	Define the groups provided in the group-mapping file as non- overlapping. If this option is activated, GIDs in the annotation files shall not be provided
	1	Input controls
-n,site nam	ies	*
bool	false	Also read in site names from the annotation files. This option does not affect calculations, but the names will be shown in the detailed and site-wise output files. The names can be duplicated, but cannot be empty or contain double quotes
-t,time_uni	t	
enum	none	Time unit if the sequences are time series (none otherwise). Supported values: sec, min, hour, day
-a1r,anno1f enum	all	The rule to resolve site overlaps within the first annotation: - all: leave all sites untouched - first: retain only the first site from an overlapping group - last: retain only the last site from an overlapping group - merge: merge an overlapping group to single site If first or last is selected, only one site will be retained from the whole group, even if the first and last sites of the group do not overlap. For circular sequences, only all and merge are supported
-a2r,anno2f	ile resolv	
(see the option -a		.~
-a1bs,anno1		n shift
integer	0	Constant shift in symbols of site start positions in the first annotation; this number will be added to all the start positions. If strand detection is activated, this option affects actual end positions in the reverse strand. If frame detection is activated, the frames are not affected by this value
-a2bs,anno2	file begir	shift
(see the option -a		
-ales,annol	-	hift
integer	0	Constant shift in symbols of site end positions in the first annotation. For details, see the option -albs
-a2es,anno2	file_end_s	hift
(see the option -a	1es)	

,	63 3 4	
-e,end_over	, 	
enum	forbid	The rule for overflowing site end positions, which is required if a site start is non-positive or a site end exceeds the sequence length: - forbid: terminate program with the error message - trim: trim the site to fit the sequence; ignore, if both, begin and end exceed the sequence length - ignore: ignore the site completely - circular: make overflowing sites reappear at the other end of the sequence; the difference between the end and the begin must still be non-negative and less than the sequence length For time series, circular is not supported
-z,zero_for		
bool	false	Treat nan values as zeros to calculate averages (default: exclude from averaging)
-min,min_gr	coup_size	
integer	1	Minimal number of sequences of a group. Smaller groups will be ignored. Must be a positive integer
-max,max_gr	oup_size	·
integer	0	Maximal number of sequences of a group. Larger groups will be ignored. Must be a non-negative integer. If 0, the size is unlimited
-d,detect	+	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,
enum	none	If strand, detect DNA strand based on the information in the file. If frame, in addition detect reading frames on the basis of the remainder of division by 3 of the gene start position in the corresponding strand. The frames are not adjusted by user-specified shifts (see the option <code>-albs</code> for details). The default value can be changed only in the simplified <i>GenBank</i> or <i>BED</i> modes
		Additional output files
-od,outfile	detailed	
string		Output file with details at the sequence and group levels
-os,outfile		
string	(empty)	Output file with site-wise statistics including match information
-ou,outfile	_union	
string	(empty)	Output TSV file with the union of the two input annotations
-oi,outfile	_intersect	ion
string	(empty)	Output TSV file with the intersection of the two input annotations
-oc1,outfil	T	y
string	(empty)	Output TSV file with the complement of the first/benchmark annotation
-oc2,outfil	e_compleme	ent_2
string	(empty)	Output TSV file with the complement of the second/prediction annotation
-ore1,outfi	le_rel_enr	,
string	(empty)	Output TSV file with the sites of relative enrichment in the first/benchmark annotation
-ore2,outfi	le_rel_enr	ichment_2
string	(empty)	Output TSV file with the sites of relative enrichment in the second/prediction annotation

Output options					
-osd,outfile sites diff					
enum	all	Limit the site-wise statistics to:			
		- all: do not limit			
		- matched: show only sites that do have a match in the other annotation under the provided criteria			
		- unmatched: show only sites that do not have a match in the other annotation under the provided criteria			
		- discrepant: show only sites that do not have a perfect			
		reciprocal match in the other annotation. Not applicable, if			
		overlap criteria are applied to patched			
		This option applies only in a non-enrichment mode			
-c,clean	T	,,			
bool	false	Produce cleaned output TSV without the commented lines. This file will contain a single header line with column names and will be			
		ready for import into various data analysis software packages			
-sort,sort_	output				
bool	false	Sort the main output file by the group names (default: sort by the order of appearance in the input). Applicable only, if groups are defined			
-sum,calcul	ate sums				
bool	false	Calculate sums of integer counts in addition to averages. Adds additional row in the main output table. Applicable only, if groups are defined			
		Other options			
-preparse,p	reparse_ma	apfile			
bool	false	Pre-parse the group-mapping file before parsing the sequence length file. This option improves performance, if the number of SIDs in the sequence length file is much larger than in the mapping file			
-wwarning	-w,warning level				
integer	11	Warning level:			
integer		- 0: switch off warnings			
		- 1: show all warnings			
-q,quiet	1	2. 55			
bool	false	Do not print the progress in the command line			
DUUI		bo not print the progress in the command inte			