DMAP2: User Guide

DMAP2 builds on the original DMAP distribution (Stockwell, *et al.*, 2014) by providing scripts to simplify the selection of processing options and running the programs for the various steps.

Overview:

After installation of the various prerequisite software packages and genomic resources, DMAP has a series of steps:

- a. Genome preparation to generate files in required form: steps that usually only need doing once for a series of samples. bismark mapping needs an indexing step for the genome, whereas bsmapz only requires that the genome is in a single file. Information files are created for steps (d) and (e)
- b. Adaptor trimming of bisulphite sequence reads for each sample
- c. Mapping reads for each sample to the genome with bismark or bsmapz producing .bam or .sam files
- d. Comparing mapped data with the DMAP program diffmeth
- e. Relating genomic fragments or regions of interest to genomic features with the DMAP program identgeneloc

DMAP2 groups these steps into a number of scripts each implementing one or more operations. If prior work has already performed some steps (genome preparation, mapping) then those can be omitted and appropriate information put in the basic parameter file. While the mapping scripts are based on using the bismark or bsmapz (formerly bsmap) bisulphite aligners, other aligners can be used in which case alignment files from them can be processed from the diffmeth point at step (d).

Script details:

(i) Information common to all scripts is specified in a file like dmap_basic_params.sh which defines the following variables:

bismark_build_options - options for building bismark genomic index bismark_run_options - options for bismark at runtime bsmapz exe - where bsmapz executable is, blank if already on your path

- bsmapz_cores number of processor cores for bsmapz run, determined by your CPU, the more available, the faster the run
- bsmapz_mismatch_val if <1 is the proportion of allowable mismatches in a read, if >=1 the actual number of allowable mismatches: setting this too high will reduce the unique mapping and probably increase processing time

- annotation file location location of the annotation file(s)
- annotation_files list or array of the files: SeqMonk, EMBL or Genbank expect 1 file for each chromosome, GTF & GFF3 use a single file

We have found SeqMonk annotation useful when the locations of TSS (Transcription Start Sites) and CpG Islands are needed, since these are contained in the SeqMonk files. It is also possible to include extra features into SeqMonk files if required. See **Prerequisites** section below for descriptions of how to obtain SeqMonk files or Gencode GTF annotation files.

(ii) dmap_index_build.sh - performs the genome preparation (step a) and further sets up details for the annotation step (e). The script takes information from dmap_basic_params.sh For bismark mapping, by default, the indices will be built in the same location as the genome fasta files, but this requires you to have write access to that location. If not then you can specify some other writeable directory in dmap bismark index location in dmap basic params.sh.

Since all the mapping for an experiment would usually be done against the same genome, the index building step should only be needed once.

The annotation preparation depends on the annotation format and produces an information file relating chromosome IDs to files for EMBL, Genbank and SeqMonk formats. This is not needed for GTF/GFF3 formats.

Once the file dmap_basic_params.sh has been set up for your environment the script should be run with a command like:

./dmap index build.sh dmap basic params.sh

noting that this, for bismark mapping, will take some time for the genome indexing process. It may be worth using Linux/Unix tools like nohup to keep such jobs running in the background, especially if you need to disconnect from the machine running this operation. Mapping with bsmapz does not require an indexing operation, but the dmap_index_build step is still required to ensure that genome files are present both as a single file (for bsmapz) and multiple files (for diffmeth) forms as below.

The DMAP program diffmeth needs one genome fasta file per chromosome. In the event that multiple chromosomes or contigs are present in one file, the script split_fasta.awk in Appendix VI will separate them into individual files, each named by the leading part of the

genome file name plus the chromosome fasta ID. The script is generated and run automatically if required.

In order to prevent the bismark indexing and mapping steps from finding multiple files for each chromosome, when a single file is required, it will be written to a directory (created if needed) genome_single. Or if the genome source is a single file it will be split, for diffmeth use, into a directory multi genome.

The genome can be gzip compressed (.gz extension), but will be decompressed during dmap_index_build.sh operation in order to provide appropriate input to bsmapz and diffmeth. This will also generate genonme_single and multi_genome directories as required.

(iii) map_bisulphite_reads.sh performs the steps of adaptor trimming (b) and bismark or bsmapz mapping (c). Parameters specific to each sample are in a file like sample_params.sh (example in Appendix II) where dmap_run_type is set to RRBS or WGBS, needed here since RRBS library preparation adds a CG dinucleotide to the end of each MspI fragment and these require removal during adaptor trimming. In order to avoid issues with overrepresentation of C's central to short reads with paired end mapping, we routinely used read 1 only. See Appendix I for other possible options.

The input fastq files can be gzip compressed (.gz extension).

map_bisulphite_reads.sh creates and uses an adaptor file contam.fa for the adaptor trimming step and runs cleanadaptors to produce a trimmed file <samplename>_at.fastq where <samplename> is taken from the fastq input file name.

For example: the original untrimmed read file mysample1_R1.fastq would be adaptor trimmed to mysample1_R1_at.fastq then mapped with bismark to produce output files like mysample1_R1_at_bismark_bt2.bam and mysample1_R1_at_bismark_bt2_SE_report.txt. bsmapz mapping would generate mysample1_R1 at.fastq bsmapz.bam.

Running the mapping for each sample would be done with a command like:

./map_bisulphite_reads.sh dmap_basic_params.sh sample_params.sh

again noting that some time will be needed for each sample. The use of nohup or some other utility to run these jobs in the background is useful.

Memory requirements for the mapping depends on the genome size and for *Homo sapiens* is typically around 10Gb for bismark and 8Gb for bsmapz.

(iv) **dmap_run_diffmeth.sh** takes mapped data from bismark or bsmapz and performs the required differential methylation analysis (step d) with parameters from a file which specifies the type of analysis, the input files and the diffmeth output file.

diffmeth performs several different types of analysis:

pairwise Fisher's Exact statistic comparison of two samples

chisquare Chi Square statistic for differentially methylated regions over 2 or more

samples

Anova Analysis of variance (F statistic) for two or more sets of samples

CpGlist List methylation for each CpG for one or more samples
binlist List + and - methylation for each region (bin) for 1 sample
binlist pc Append a % methylation column to binlist lines

An example diffmeth parameter file is in Appendix III and, in this case, is for a 3 group Anova run. The important variables are:

dmap_run_type - defines RRBS or WGBS and should match the setting for the
 map bisulphite reads.sh step

diffmeth_run_type - defines the analysis type (Anova, chisquare, pairwise, CpGlist, binlist or binlist pc) noting that these are case sensitive

file_list1, file_list2, ... each containing a list (array) of bam or sam files for each group.

diffmeth out name sets the name of the output file.

diffmeth bases its operation on regions for WGBS or fragments(usually MspI) for RRBS and the term bin applies to those genomic divisions.

file list requirements for different analyses:

binlist, binlist_pc a single file in file_list1
CpGlist one or more files in file_list1

pairwise two files in file_list1

chisquare two or more files in file list1

Anova two or more sets of files, at least 2 in each in

file list1, file list2, ...

Anova runs also need anova_group_number and anova_detail to be defined as well as additional lists of file names in file_list2, etc.: one such list for each group. If the variable anova_fold_difference is set to "yes" a column of fold methylation differences will be added to the output.

Further control of diffmeth parameters is *via* the variables:

```
diffmeth_fragment_min - minimum length of RRBS fragments diffmeth fragment max - maximum length of RRBS fragments
```

wgbs window - window length for WGBS runs

min_cpg_hits - minimum number of hits for a CpG to be included in analysis

min_valid_cpgs - number of CpGs meeting min_cpg_hits for a fragment/region to be included

min_sample_count - minimum number of samples fulfilling min_cpg_hits and min valid cpgs for a fragment/region to be included

map_init_CpG_to_prev - for 3' matching reads, the initial C is counted in the adjacent RRBS fragment

pr threshold - reject fragments/regions with Pr above this threshold

For SAM mapping files it may be necessary to define max_sam_readlength if the reads exceed 150bp. This is not required for BAM files.

Run the diffmeth script with a command like:

```
./dmap_run_diffmeth.sh dmap_basic_params.sh my_diffmeth_params.sh
```

noting that the execution time can range from 20 minutes to some hours, depending on the amount of data and the computer performance. diffmeth runs will typically require some 5Gb of RAM or more depending on the genome size, the number of samples and the coverage.

(v) **dmap_run_genloc.sh** implements step (e) on the output from **diffmeth** runs or other appropriately formatted input in which **identgeneloc** looks upstream (in the sense of the feature) for the nearest feature of interest to each region/fragment. The minimum input is 3 columns (chromosome ID, start, stop) but more usually the other columns generated by **diffmeth** would be included.

dmap_run_genloc.sh requires a parameter file (example in Appendix IV) to specify the source of annotation data and to control other aspects of the run. Variables are:

- genloc input file name of input file with 3 or more columns
- genloc_column_number number of columns of input file to include, leave as 0 for automatically including all columns
- show_TSS for SeqMonk annotation, show nearest upstream transcription start sites (TSS)
- TSS_limit if non-zero will limit how far TSS will be scanned from regions, zero means no limit
- show CpGi for SeqMonk annotation, show nearest upstream CpG island
- show CpGi internal allow CpG islands to be internal to regions
- show_TSS_CpGi_ranges show ranges for CpG islands and TSS

- genloc_GTF_featuretypes for GTF annotation, specify the desired features as in column 3 of the GTF file, e.g. gene, multiples allowed as a list or array

Run the identgeneloc script with a command like:

./dmap run genloc.sh dmap basic params.sh geneloc params.sh

identgeneloc runs will typically require some 350Mb of memory, depending on the size of the annotation file(s) and would take some 12 minutes or so for input files of 500,000 regions.

Prerequisites:

In order to run DMAP the following prerequisites should be obtained and installed.

(a) Genomes

DMAP2 requires genomes in FASTA format either a single whole genome file or with each chromosome in an individual file: files can be gzip compressed (.gz externsion). Various sources exist, but one of the most convenient is

 $\label{lem:https://www.ensembl.org/info/data/ftp/index.html following the 'DNA (FASTA)' link. Three different versions of each chromosome are offered: .dna.,$

.dna.rm. and .dna.sm. respectively being complete genome, genome with repeat regions masked as 'N' and genome with repeats in lower case. The first (.dna.) is probably the most useful.

(i) As a single file (for Homo sapiens):

Select the file Homo_sapiens.GRCh38.dna.primary_assembly.fasta.gz for download or execute the following at the command line (all complete on one line):

```
curl -0 ftp.ensembl.org/pub/release-
105/fasta/homo_sapiens/dna/Homo_sapiens.GRCh38.dna.primary_assembly.fasta.gz
```

The complete assembly contains all chromosomes, including mitochondrial, as well as a series of scaffolds with IDs of the form KI270728.1. Mapping reads to these may be inconvenient in which case downloading the required multiple files as below may be better. DMAP2 is capable of using compressed genome files so they do not need to be decompressed.

In dmap_basic_params.sh set dmap_genome_file_location to the directory where this file has been downloaded and fill dmap_genome_fasta_files with the complete genome file name.

(ii) As multiple files (for Homo sapiens):

Files can be downloaded by selecting each individually with the browser or as a group from the command line using a command like:

```
curl -0 ftp.ensembl.org/pub/release-
105/fasta/homo_sapiens/dna/Homo_sapiens.GRCh38.dna.chromosome.[1-22].fa.gz
followed by
```

```
curl -0 ftp.ensembl.org/pub/release-
105/fasta/homo_sapiens/dna/Homo_sapiens.GRCh38.dna.chromosome.[X-Y].fa.gz
```

noting that each command is complete on one line.

In dmap_basic_params.sh set dmap_genome_file_location to the directory where these files have been downloaded and fill dmap_genome_fasta_files with the chromosome file names.

(iii) Decompressing downloaded files:

Not actually necessary since DMAP2 will generate these as required during the dmap_index_build.sh phase, however it may be helpful to check the file contents which can be done with commands like:

gzip -dc Homo_sapiens.GRCh38.dna.chromosome.1.fa.gz | more
or

gunzip Homo_sapiens.GRCh38.dna.chromosome.1.fa.gz

which will replace the compressed file with the uncompressed version. Compressed fasta files are about 1/3 the size of the uncompressed versions.

(iv) Other genomes:

Various releases are available at ftp.ensembl.org/pub/ noting that the fasta links will return files in the required formats for DMAP2 mapping.

- **(b) Annotation** files in one of the following formats:
- (i) SeqMonk obtained through the SeqMonk mapped sequence data analyser, available from

https://www.bioinformatics.babraham.ac.uk/projects/seqmonk/

by starting the application and choosing File > New Project then 'Import Genome From Server' and selecting then downloading the genome of choice. This should create a directory seqmonk_genomes, probably in your top level directory, containing the organism and genome version which contain a series of files 1.dat, 2.dat,... - these contain the annotations for each chromosome. Set the location of these files in dmap_basic_params.sh as annotation_file_location and fill annotation_files with the list of chromosome files. feature_annotation_type is set to SeqMonk.

SeqMonk annotation has proven particularly useful when transcription start sites and CpG islands are needed and allows the choice of specific biotypes for the features of interest. Further, it is possible to append other feature information (e.g. enhancers) to each chromosome: Appendix V contains examples of added features.

(ii) GTF - obtained from https://www.gencodegenes.org/ for mouse or human, by choosing the organism and genome version, then downloading the comprehensive gene annotation file as GTF which will download a file with a name like: gencode.v39.annotation.gtf.gz which can be moved to an appropriate location. The dmap index build.sh step will decompress the file, or you can do this with:

gunzip gencode.v39.annotation.gtf.gz

producing gencode.v39.annotation.gtf. Set the location in dmap_basic_params.sh as annotation_file_location and annotation files to the filename. feature annotation type is set to GTF.

- (iii) GFF3 obtained as for GTF, with similar basic parameter settings.
- (iv) EMBL obtained from https://www.ensembl.org/info/data/ftp/index.html as 'Annotated sequence (EMBL)' or consider using utilities like ftp or curl to transfer the files at the command line, with a command (for Homo sapiens) like:

```
curl -0 ftp.ensembl.org/pub/release-
105/embl/homo_sapiens/Homo_sapiens.GRCh38.105.chromosome.[1-22].dat.gz
```

(all complete on 1 line). X and Y chromosomes will need [1-22] replaced by [X-Y].

The dmap_index_build.sh step will decompress the files or you can decompress them with a command like:

```
gunzip Homo sapiens.GRCh38.105.chromosome.*.dat.gz
```

Set the location of these files in dmap_basic_params.sh as annotation_file_location and fill annotation_files with the list of chromosome files. feature annotation type is set to EMBL.

Note that EMBL (and Genbank) feature tables obtained from ENSEMBL only return ENSEMBL gene IDs with DMAP.

(v) Genbank - available from NCBI, though hard to find there and comes with awkward chromosome IDs. Better obtained similarly to EMBL from Ensembl as 'Annotated sequence (GenBank)' or at the command line with something like:

```
curl -0 ftp.ensembl.org/pub/release-
105/genbank/homo_sapiens/Homo_sapiens.GRCh38.105.chromosome.[1-22].dat.gz
```

as above. feature_annotation_type is set to Genbank.

For SeqMonk, EMBL and Genbank annotation formats the list of file names and chromosome IDs are written to the file set in dmap_annot_info_file, which defaults to dmap_annot_info.txt.

Note that Genbank (and EMBL) feature tables obtained from ENSEMBL only return ENSEMBL gene IDs with DMAP.

(c) Bismark: we have generally found bismark to be a good performer for vertebrate genomes, available from https://github.com/FelixKrueger/Bismark or from https://www.bioinformatics.babraham.ac.uk/projects/bismark/. Installation is as per instructions, which amount to putting the perl executables into an appropriate directory: either system ones (/usr/local/bin - system privilege required) or your own top level bin.

The variable bismark_run_options in dmap_basic_params.sh can be set to reflect your needs: we have found that '-N 1' restricts the permitted seed mismatches to 1 (defaults to 2) works well for our data and gives significantly faster mapping. The option '-- bowtie2' is not now necessary for recent bismark versions, but remains for completeness.

- (d) bsmapz: replaces the original bsmap and is available as source code from https://github.com/zyndagj/BSMAPz. bsmapz runs much faster than earlier versions of bsmap and works well with a wide range of genomes. Note that the methods of resolving multiply-mapped reads differ between bsmapz and bismark so they do not produce identical results it is your choice which you prefer. Building bsmapz requires C and C++ compilers and the install process is described in the distribution.
- (d) bowtie2: recent versions of bismark require the aligner bowtie2. If not already installed in your systems it is available from

https://github.com/BenLangmead/bowtie2 or

https://sourceforge.net/projects/bowtie-bio/files/bowtie2/, the latter providing precompiled binaries which can be unzipped and installed in an appropriate directory.

(e) samtools: required by Bismark in order to generate bam rather than sam files. (bam files are about 1/3 the size of the corresponding sam files.) If not already installed, it is available from https://sourceforge.net/projects/samtools/ as source code which will need unpacking and compiling using instructions like:

```
bzip2 -dc samtools-1.14.tar.bz2 | tar -xvf -
cd samtools-1.14
./configure
make
make install
```

The latter instruction may require system privilege (e.g. sudo). Alternatively you can copy the executable to a location on your PATH with something like:

```
cp samtools ~/bin/
```

(f) DMAP: available as source code from

https://github.com/peterstockwell/DMAP either by downloading as a zip archive from 'Code->Download ZIP' and using unzip or from the command line with:

```
git clone https://github.com/peterstockwell/DMAP
```

producing either DMAP-master or DMAP which contains instructions for building in README.md.

The source code should compile with any reasonable C compiler, but note that the library zlib is needed for full functionality with .bam files. The main executables needed are diffmeth and identgeneloc which should be copied to an appropriate directory on your PATH or set path_to_dmap in dmap_basic_params.sh to the actual location.

```
(g) DMAP2: available from:
```

https://github.com/peterstockwell/DMAP2

with a web browser or from the command line with:

```
git clone https://github.com/peterstockwell/DMAP2
```

which will produce a directory DMAP2 master or DMAP2 containing the following:

README . md - distribution notes

DMAP2_UserGuide.pdf - this documentation. Further documentation is present in the script files as commments

Scripts - directory containing the main DMAP2 scripts:

dmap_run_genloc.sh - to relate sequence regions or fragments to genomic features dmap basic params.sh - to provide basic project details to all the above scripts

Example_params - directory containing example parameter scripts for the various steps:

sample_params.sh - typical sample parameters for the mapping step for

map bisulphite reads.sh

dmap_anova2_params.sh - parameters for diffmeth with 2 group Anova statistic dmap_anova3_params.sh - for 3 group Anova statistic dmap_chisq_params.sh - for differential methylation analysis by Chi Square statistic

dmap_prwise_params.sh - for Fisher's Exact pairwise analysis dmap_cpglist_params.sh - list CpG counts and positions for a single sample for each fragment or region

dmap_binlist_params.sh - list methylation counts for a single sample for each fragment or bin

Extra_scripts - directory containing some other useful scripts:

split fasta.awk - to split multi entry fasta files into individual files

You may find it helpful to copy the scripts into the working directory for your project, especially dmap_basic_params. sh and the example parameter scripts which will need editing to suit your data. The example run commands shown above assume that these files are all in your project working directory.

References:

Stockwell, P.A., Chatterjee, A., Rodger, E.J. and Morison, I.M. "DMAP: Differential Methylation Analysis Package for RRBS and WGBS data" *Bioinformatics* (2014) DOI: 10.1093/bioinformatics/btu126.

Krueger, F., and Andrews, S.R., Bismark: a flexible aligner and methylation caller for Bisulfite-Seq applications, *Bioinformatics*,

(2011), https://doi.org/10.1093/bioinformatics/btr167

Yuanxin Xi and Wei Li, "BSMAP: whole genome bisulfite sequence MAPping program" (2009) BMC Bioinformatics 2009, 10:232.

Appendix I: Options for mapping paired end reads.

The authors of bismark (Felix Krueger and Simon Andrews, 2011) advise that the use of paired end reads with RRBS fragments will result in over-representation of the overlapping portion of short fragments, given that the 40-220 bp size selection will generate many reads shorter than read lengths presently available. A possible solution is to use tools like FLASH (http://www.cbcb.umd.edu/software/flash) to generate longer reads where paired end reads clearly overlap. This process generates 3 fastq output files: longer overlapped reads, R1 non-overlapped and R2 non-overlapped. Since bismark is only configured to accept either single ended files (or a list of them) or paired end files for R1 & R2 (or a list for each), a strategy would be for two bismark runs, one single-ended for joined reads, the other paired-ended for unjoined reads, then use samtools merge to combine the results before continuing with DMAP analysis. Our experience with this was that it did not improve the mapping efficiency sufficiently to justify the extra work, hence our decision to work with single-ended data.

Appendix II: An example sample parameter file for mapping:

```
# sample params.sh: parameters to descibe a single bisulphite
# sequence sample, including the raw fastq file(s) and any
# trimming that might be justified by the read qualities.
# identify if this was RRBS or WGBS: this determines a number
# of downstream operations, including adaptor trimming and
# differential methylation calculations. Values are RRBS or WGBS.
dmap run type="RRBS";
# dmap run type="WGBS";
# name of adaptor-trimmed output dir:
adtrimmed out dir="./";
# sample file name(s): we usually only map read 1, so
# read 2 name would omitted.
dmap sample_files=("CRC-P-10_1.fastq.gz" "CRC-P-10_2.fastq.gz");
# hard trim length, leave 0 for no trimming.
dmap sample trim length=0;
# minimum read length for retention: usually set this to 20
dmap min read length=20;
# mapping output directory
bismark out dir="bismark out/";
```

Appendix III: Example parameter file for diffmeth runs

In this case an Anova run with three groups.

```
# diffmeth_run_params.sh: parameters to define differential
# methylation run type and mapped files for the DMAP diffmeth
# program.
# identify if this was RRBS or WGBS: this determines a number
# of downstream operations, including adaptor trimming and
# differential methylation calculations. Values are "RRBS" or "WGBS"
# "RRBS" compares methylation on fragments (usually MspI) in size range
# "WGBS" compares methylation on defined windows or regions
# this is previously specified for the adaptor trimming and mapping step
# and obviously the values here should be consistent with those.
dmap_run_type="RRBS";
# dmap run type="WGBS";
# type of diffmeth run: values are one of "pairwise" "chisquare" "Anova"
"CpGlist" "binlist" or "binlist_pc"
# where:
# "pairwise" runs Fisher's Exact statistic on two samples
# "chisquare" runs a Chi Square test on multiple samples to identify
fragments/regions with greatest variation
            run Analysis of Variance on two sets of multiple samples,
producing F statistic
# "CpGlist"
              lists methylation counts for each CpG
# "binlist"
            lists methylation information for each fragment/region
# "binlist pc" same as binlist but appends % methylation to each line
diffmeth_run_type="Anova";
case $diffmeth run type in
pairwise)
# "pairwise" requires two mapping output files
file_list1=("./rundata/bismark_out/CRC-P-10_1_at_bismark_bt2_pe.bam"
"./rundata/bismark_out/NC-P-20_1_at_bismark_bt2.bam");
chisquare)
# "chisquare" requires a list of at least 2 mapping output files
file list1=(
 './rundata/comb D1.bam"
 "./rundata/comb_D2.bam"
 "./rundata/comb_D3.bam"
);
;;
Anova)
# Anova requires a group number to define how many groups are involved
# and a list of at least 2 mapping output files for each group
# a further parameter defines the amount of information returned, being
# one of: "anova_simple" "anova_medium" or "anova_full"
# where
# "anova simple" returns F statistic and its probability
# "anova medium" adds sample counts and greater methylated group
# "anova full"
                 further adds methylation figures for each CpG in
fragment/region
anova_group_number=3;
```

```
file list1=(
 "./rundata/comb_D1.bam"
 "./rundata/comb_D2.bam"
 "./rundata/comb_D3.bam"
"./rundata/comb D4.bam"
file list2=(
 "./rundata/comb_D5.bam"
 "./rundata/comb_D6.bam"
"./rundata/comb_D7.bam"
);
file list3=(
 "./rundata/comb D8.bam"
 "./rundata/comb_D9.bam"
 "./rundata/comb_D10.bam"
anova_detail="anova_full";
CpGlist | binlist | binlist_pc)
\# "CpGlist" requires 1 or more mapping files
# "binlist" or "binlist_pc" require a single mapping file
file_list1=("./rundata/comb_D1.bam")
;;
esac
# diffmeth requires a fragment size specification, even for WGBS (ignored
then)
diffmeth_fragment_min=40;
diffmeth_fragment_max=220;
# "WGBS" requires a window (tile) length
wgbs_window=1000;
# output file name for diffmeth
diffmeth out name="diffmeth output prwise NC CP not.txt";
# threshold criteria for including fragment/regions
# No. of hits per CpG for fragment/region to be valid
# default is 1
#min cpg hits=1;
# No. of CpG of fragment/region that must meet min_cpg_hits criterion
# 0 indicates no check for this
min valid cpgs=0;
# Minimum No. samples reaching criteria for a fragment/region to be
included
# defaults to 2
min sample count=6;
```

```
# For RRBS 3' mapping reads, count initial CpG to previous fragment.
map init CpG to prev="yes";
# Probability threshold: omit fragments/regions with Pr greater.
# default (1.0) implies no filtering
#pr_threshold=1.0;
# Max read length for .SAM files (not needed for .BAM, though)
# defaults to 150bp, for 0
#max_sam_read_length=150;
Appendix IV: Example parameter file for dmap run genloc.sh
# genloc run params.sh: parameters to define gene location
# operation with identgeneloc. Some relevant parameters
# are already set with the basic parameters, this file
# relates to the specific settings required for a project.
# file of chromosome & coordinate data (min 3 columns) as:
# chrid start stop
genloc input file="diffmeth output binlist nz.txt";
# name of output file: identgeneloc writes to stdout, we need
# to direct that somewhere, else will still got to stdout
genloc output file="genloc binlist nz chrchk gc.txt";
# the number of columns can be identified automatically
# putting a non-zero value here will override the automatically
# chosen value
genloc_column_number=0;
# Control whether regions internal to genes are allowed
# and whether they are to be related to exon/intron
# boundaries.
# values for the parameter internal_to_genes are:
# "" = disallow
# "internal" = allow
# "exon intron" = relate to exon/intron boundaries
internal to genes="exon intron";
# SeqMonk annotations define TSS and CpG islands
# ignored for other annotation formats
show TSS="yes";
# limit for TSS ranges, zero=nolimit
TSS limit=0;
```

```
show_CpGi="yes";
# CpG islands can be shown within regions: "yes" permits this
show CpGi internal="yes";
show TSS CpGi ranges="yes";
# For SegMonk annotations, can restrict to particular biotypes;
# valid values are: 3prime overlapping ncrna IG C gene IG C pseudogene
IG D gene IG J gene IG J pseudogene
# IG V gene IG V pseudogene Mt rRNA Mt tRNA Mt tRNA pseudogene TEC
TR C gene TR D gene TR J gene TR J pseudogene
# TR V gene TR V pseudogene ambiguous orf antisense lincRNA miRNA
miRNA pseudogene misc RNA misc RNA pseudogene
# ncrna host non coding non stop decay nonsense mediated decay
polymorphic pseudogene processed transcript
# protein coding pseudogene rRNA rRNA pseudogene retained intron
scRNA pseudogene sense intronic sense overlapping
# snRNA snRNA pseudogene snoRNA snoRNA pseudogene tRNA pseudogene
transcribed unprocessed pseudogene
# protein coding is probably one of the most useful.
genloc segmonk biotypes=(
"protein coding"
# GTF & GFF3 annotations allow selection of attributes and feature
# Attributes default to "transcript name, transcript id, gene_name" if
# not specified. Actual attributes can vary widely for GTF format
genloc_GTF_attributes=(
"transcript_name"
"transcript_id"
"gene_name"
)
# feature types default to "exon, CDS, start_codon, stop_codon, transcript"
# not specified. Valid values for GTF are are any of:
CDS, start codon, stop codon, exon, intron CNS, intron, 5UTR, 3UTR, CNS, inter, t
ranscript, gene
# GFF3 values are far more diverse
genloc_GTF_featuretypes=(
"gene"
)
```

Appendix V: Example of additional SeqMonk annotation appended to a chromosome feature file

```
FT
     gene
                     856539..856757
                     /description="enhancer chr1:856539-856757"
FT
                     /name="enh 856539"
FT
                     /score="2"
FT
                     /biotype="enhancer"
FT
FT
     gene
                     858256..858648
FT
                     /description="enhancer chr1:858256-858648"
                     /name="enh 858256"
FT
                     /score="5"
FT
                     /biotype="enhancer"
FT
FT
                     868565..868684
     gene
                     /description="enhancer chr1:868565-868684"
FT
                     /name="enh_868565"
FT
                     /score="2"
FT
FT
                     /biotype="enhancer"
```

Appendix VI: split_fasta.awk script to split multi-sequence fasta files.

```
# split fasta.awk: script to break multi-entry fasta
# files into separate files, each containing 1 entry.
# files named by the fasta IDs, with optional header
# Peter Stockwell: 21-Feb-2022
#
# usage:
# awk -f split_fasta.awk <multi_fasta_file>
# or piped, e.g.
# cat <multi_fasta_file> | awk -f split_fasta.awk
# Can define optional useful parameters in command line:
# namehdr - prefixed to each output file name, can be a directory or
string or both (e.g. namehdr="singles_fasta/")
# extension - use as file extension, default is ".fa"
# infofile - will write table of Chr IDs and full file paths to this
file
# use as: e.g.
# awk -f split_fasta.awk namehdr="genome_singles/" extension=".fasta"
infofile="chr_info.txt" <multi_fasta_file>
# to produce chr info.txt which is suitable for the diffmeth -G option
# If namehdr is a directory, it must already exist.
BEGIN {namehdr="";
extension=".fa";
outfile="";
infofile=""
$0~/>/{if (outfile != "")
  close(outfile);
outfile = sprintf("%s.%s%s",namehdr,substr($1,2),extension);
printf("%s\n",$0) > outfile;
if (infofile != "")
  printf("\"%s\"\t\"%s\"\n",substr($1,2),outfile) > infofile;
}
$0!~/>/{
printf("%s\n",$0) > outfile;
END{
if (outfile != "")
 close(outfile);
if (infofile != "")
  close(infofile);
}
```