

# Analysing Hi-C matrices

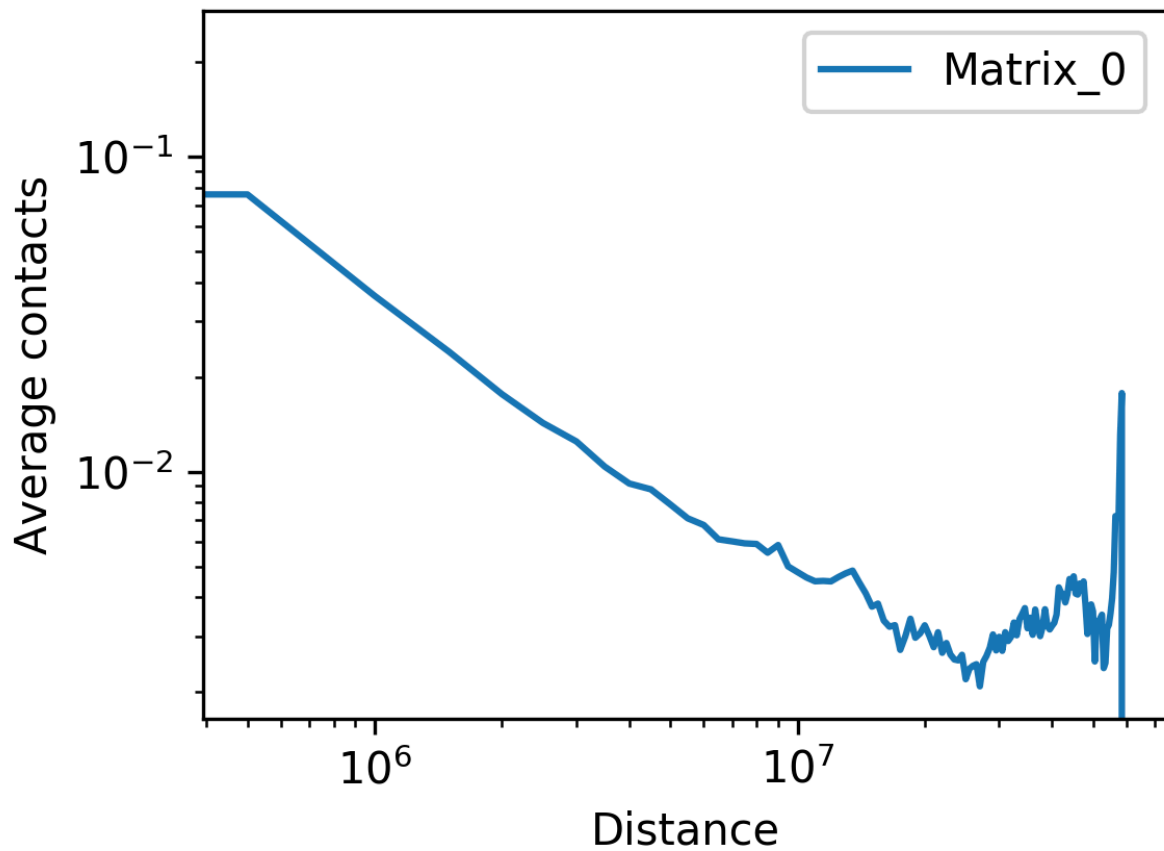
This tutorial assumes that you have gone through the `hic_matrix_generation` tutorial or followed the Getting started instructions from the FAN-C documentation.

## Practical 1: Distance decay of expected contact probability or $P(s)$ curve

The contact intensity in a Hi-C matrix gets progressively weaker the further apart two loci are. The expected values follow a distinctive profile with distance for Hi-C matrices, which can be approximated by a power law and forms an almost straight line in a log-log plot.

- (1) To calculate the expected values of any FAN-C compatible matrix, you can use the `fanc expected` command on `juicer`, `cooler` or `fan-c` matrices:

```
cd ~/examples/
fanc expected -p \
  architecture/expected/fanc_example_500kb_expected.png \
  -c chr19 \
  output/hic/binning/fanc_example_500kb.hic \
  architecture/expected/fanc_example_500kb_expected.txt
```



The raw expected values are stored in `fanc_example_500kb_expected.txt`

```
less architecture/expected/fanc_example_500kb_expected.txt
```

```
distance    Matrix_0
0           0.24442297400748084
500000      0.07759323503191953
1000000     0.03699383283713825
1500000     0.02452933204893787
2000000     0.017725227895561607
2500000     0.014272302693312262
3000000     0.011708011997703627
3500000     0.010125456912234796
...
```

It is advisable to run the `fanc expected` command before the steps below so that the calculated values are stored in the `.hic` object for future reference. Expected contact frequencies away from the diagonal will be contrasted with the observed values at each position to estimate enrichments over background. Use `--recalculate` to force a re-calculation of expected values, for whatever reason. Use `-N` to plot the unnormalised expected values.

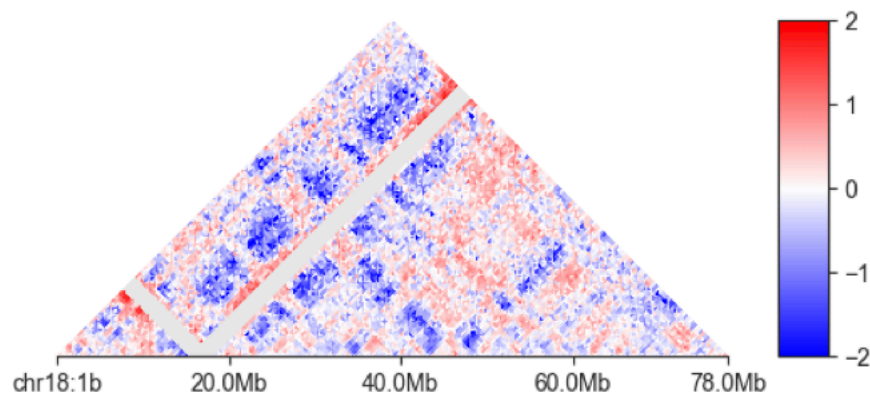
(2) We can calculate the expected value of multiple matrices at the same time in this way:

```
fanc expected \
-l "HindIII 5M" "HindIII 100k" \
  "MboI 1M" "MboI 100k" "MboI 50k" -c chr19 \
```

```
-p architecture/expected/expected_multi.png \
architecture/other-hic/lowc_hindiii_5M_1mb.hic \
architecture/other-hic/lowc_hindiii_100k_1mb.hic \
architecture/other-hic/lowc_mboi_1M_1mb.hic \
architecture/other-hic/lowc_mboi_100k_1mb.hic \
architecture/other-hic/lowc_mboi_50k_1mb.hic \
architecture/expected/expected_multi.txt
```

- (3) Having pre-calculated expected values, we can easily plot observed/expected matrices that highlight Hi-C interaction patterns more easily. Here, we are showing a log2-transformed O/E matrix:

```
fancplot \
-o architecture/expected/fanc_example_500kb_chr18_oe.png \
chr18:1-78mb -p triangular \
-e output/hic/bin/binned/fanc_example_500kb.hic \
-vmin -2 -vmax 2
```



- (4) Which layers of genome folding do you see in this picture?

## Practical 2: AB compartment analysis

Regions in a Hi-C matrix can generally be assigned to either the ‘active’ or the ‘inactive’ compartment, also called ‘A’ and ‘B’ compartments, respectively. The traditional way of calling compartments aims to enhance the plaid or chess-like pattern in the Hi-C matrix to identify regions that interact in similar ways to each other.

### Correlation matrix

This is first achieved by calculating an (auto) correlation matrix in which each entry  $i,j$  corresponds to the Pearson correlation between row  $i$  and column  $j$  of the Hi-C matrix.

- (1) The `fanc compartments` command can produce a correlation matrix (AB compartment object) from a FAN-C matrix file. Example:

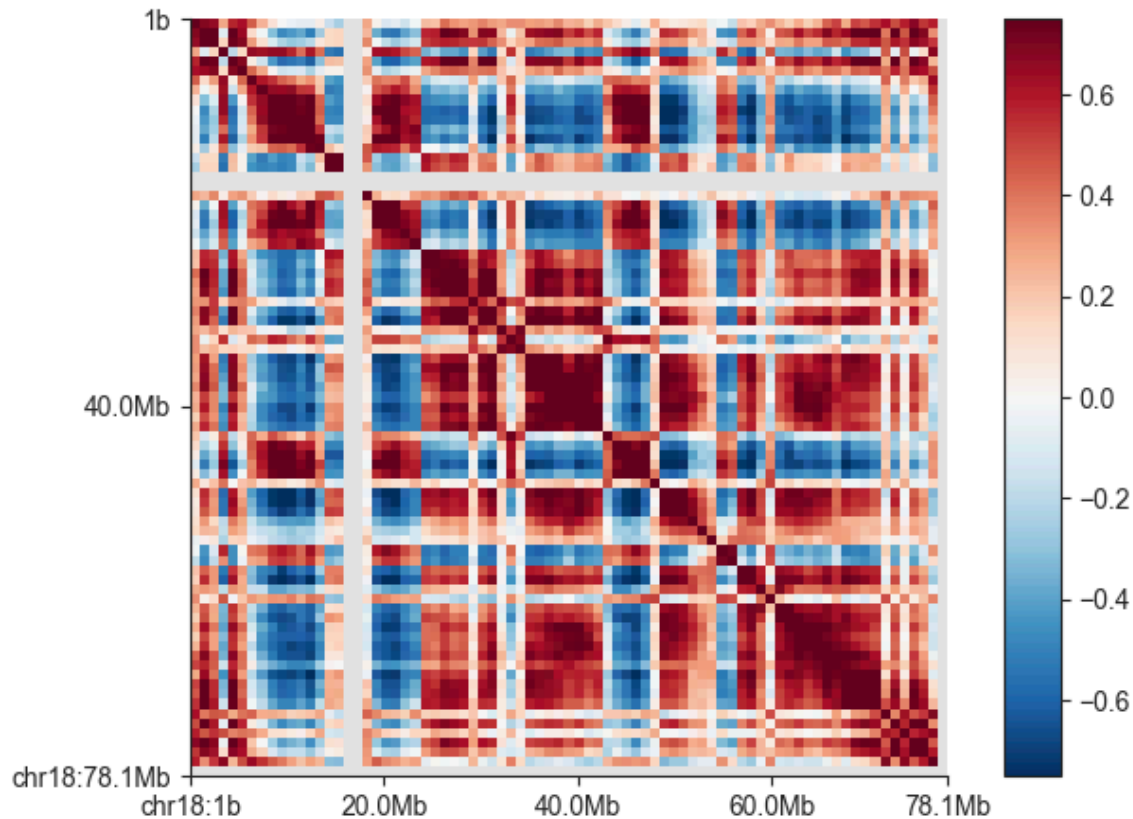
```
fanc compartments \
output/hic/bin/binned/fanc_example_1mb.hic \
```

```
architecture/compartments/fanc_example_1mb.ab
```

By default, compartment matrices are calculated on a per-chromosome basis, since each chromosome might be normalised differently and is hard to estimate expected values inter-chromosomally. To force the AB calculation on the whole genome, use the `-w` option.

(2) We can easily plot the correlation matrix using the `fancplot` command:

```
fancplot \  
-o architecture/compartments/fanc_example_1mb.ab.png chr18 \  
-p square architecture/compartments/fanc_example_1mb.ab \  
-vmin -0.75 -vmax 0.75 -c RdBu_r
```



## AB Eigenvector

To translate this enhanced chess-like pattern into a 1-dimension bed file that captures the binary nature of the correlation we typically use a PCA decomposition and keep the first eigenvector to calculate compartments. The eigenvector of the correlation matrix is used to derive compartment type and strength for each matrix bin. Generally, regions with positive values are assigned the 'A', regions with negative values the 'B' compartment.

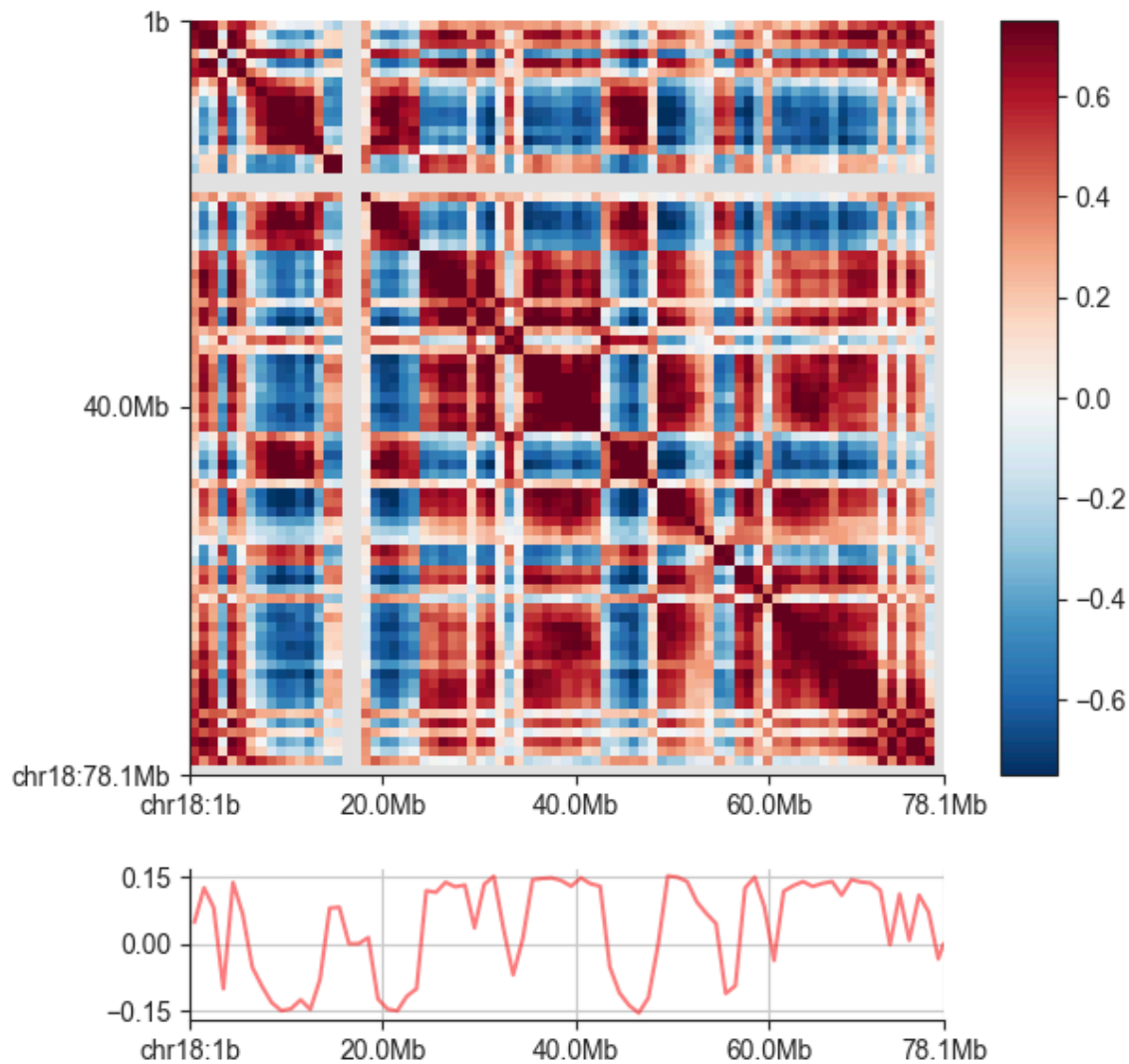
(3) To calculate the eigenvector in addition to the correlation matrix, simply add the `-v <file_name>` option to the previous command:

```
fanc compartments \  
-v architecture/compartments/fanc_example_1mb.ev.txt \  
output/hic/binned/fanc_example_1mb.hic \  
architecture/compartments/fanc_example_1mb.ab
```

If `architecture/compartments/fanc_example_1mb.ab` already exists, it will not be recalculated but the matrix is loaded from file. You can use the `-f` option to overwrite the existing file in any case.

(4) We can plot the eigenvector using `fancplot`:

```
fancplot \  
-o architecture/compartments/fanc_example_1mb.ab_and_ev.png \  
chr18 \  
-p square architecture/compartments/fanc_example_1mb.ab \  
-vmin -0.75 -vmax 0.75 -c RdBu_r \  
-p line architecture/compartments/fanc_example_1mb.ev.txt
```



NOTE: `fanc compartments` outputs the first eigenvector by default. In some cases (like organisms with non-acrosomal chromosomes) it might be useful to choose a different eigenvector (sometimes the first eigenvector identifies chromosomal arms rather than compartments). To change the eigenvector use the `-i` option, e.g. `-i 2` to keep the second instead of the first correlation matrix eigenvector.

- (5) The sign of the eigenvector does not necessarily correspond well to the A or B compartment. Often, the eigenvector is “flipped” (inverted signs on its entries). Mathematically, if  $\mathbf{x}$  is an eigenvector, so is  $-\mathbf{x}$ . You can use external information to “orient” the eigenvector, so that it most likely corresponds to the active and inactive compartments.

Specifically, you can supply a FASTA file with the genomic sequence to `fanc compartments` using the `-g <fasta_file>` argument. This is generally a good idea: `fanc compartments` then calculates the average GC content of regions with positive and those with negative eigenvector entries. As GC content has previously been shown to correlate well with compartmentalisation, the eigenvector is oriented in such a way that

negative entries correspond to ‘B’ (low GC content) and positive entries to ‘A’ (high GC content).

```
fanc compartments \
  -g hg19_chr18_19.fa \
  -v architecture/compartments/fanc_example_1mb.ev_gc.txt \
  architecture/compartments/fanc_example_1mb.ab
```

## AB Domains

Consecutive matrix bins with the same eigenvector sign are considered part of a “domain”.

(6) You can use the `-d <domain_file>` option to write the AB domains to a BED file:

```
fanc compartments -g hg19_chr18_19.fa \
  -d architecture/compartments/fanc_example_1mb.domainsgc.bed \
  architecture/compartments/fanc_example_1mb.ab
```

The domains BED file merges all consecutive bins in the same domain, which is why A and B are always alternating. It contains the domain type in the “name” field and the average eigenvector entry values of all bins in the domain in the “score” field:

## AB Enrichment profiles or ‘Saddle plots’

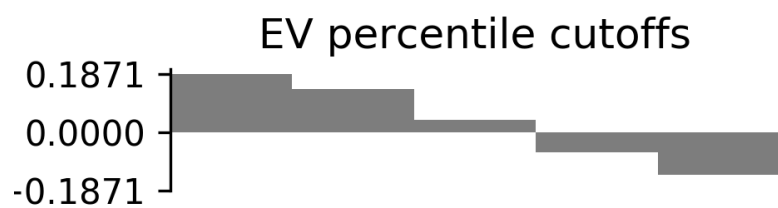
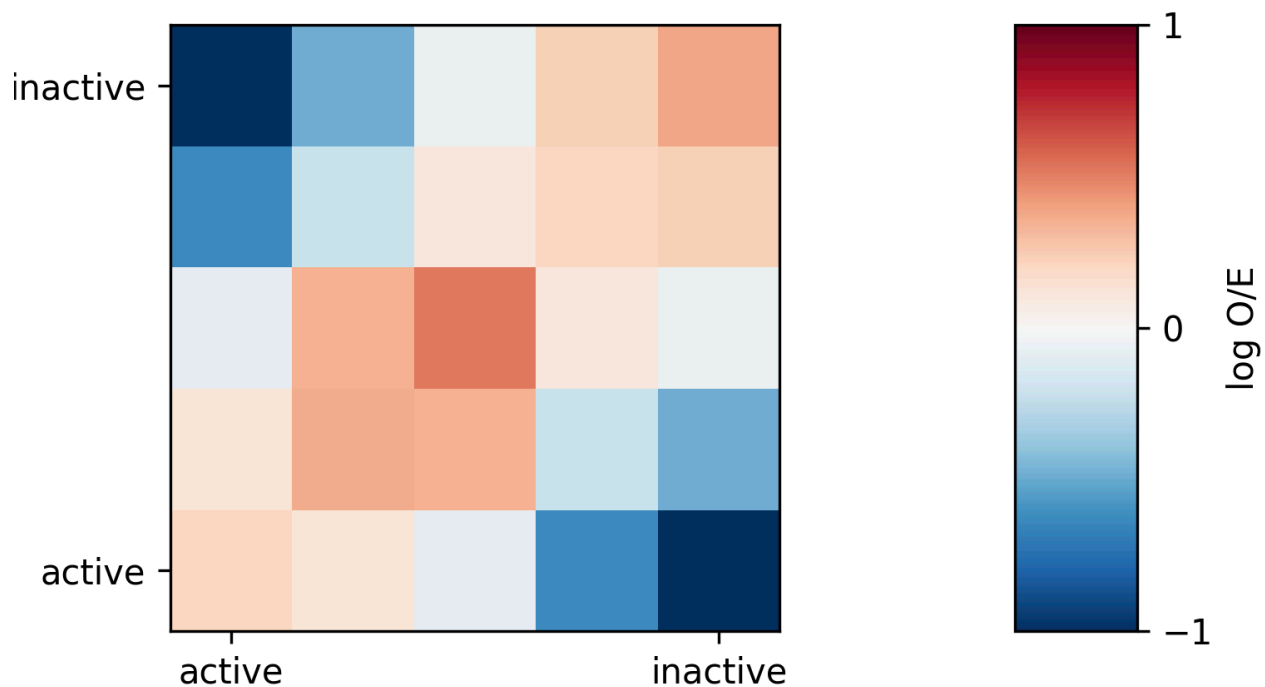
A useful way of displaying genome-wide changes at the compartment-level is to calculate average interaction frequencies between regions of the genome that have different compartmental values. i.e. To answer the question: How many interactions are there between A-A regions? how many interactions are there between B-B regions of the genome? and how many interactions occur across compartments (A-B)? Is there a change of overall compartmentalisation between conditions?

The compartmentalisation of a genome can be visualised in an enrichment profile plot. This will divide bins into percentiles using the associated eigenvector values, and then the average observed/expected (O/E) values of contacts in each pair of percentile bins is calculated and plotted.

(6) Use the `-e <plot_file>` option to generate this plot. You can also use the `-m <matrix_file>` option to output the enrichment matrix values to file for further analysis.

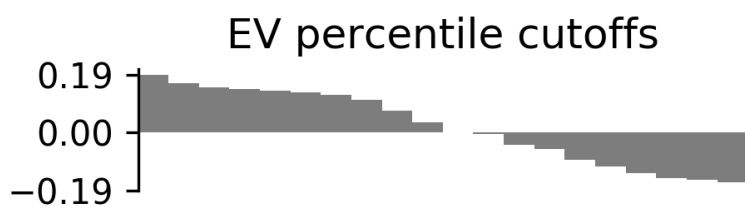
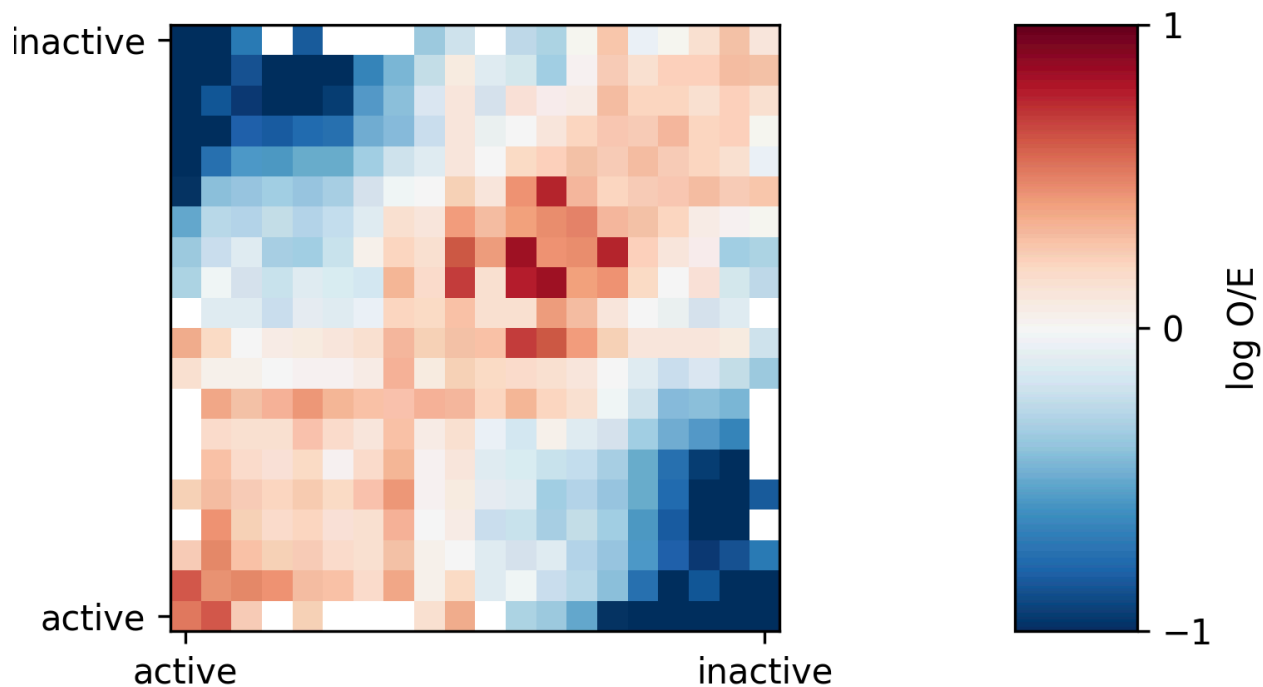
```
fanc compartments -g hg19_chr18_19.fa \
  -e architecture/compartments/fanc_example_1mb.ab_profile.png \
  output/hic/binning/fanc_example_1mb.hic \
  architecture/compartments/fanc_example_1mb.ab
```

This process will need to specify the `.hic` matrix to calculate the enrichment of the contacts over the expected values. It is also advised to use a FASTA genome file to correctly orient the compartments.



- (7) You can customise the enrichment analysis using additional parameters. By default, the percentiles for eigenvector binning are chosen at 20, 40, 60, 80, and 100. To choose a finer binning, for example, you can use `-p 10 20 30 40 50 60 70 80 90 100`. The "0" percentile is always added automatically. The eigenvector values are not necessarily centred around 0, which can lead to some unexpected enrichment plots. To force the central enrichment matrix bin to 0, and perform separate percentile calculations for values  $< 0$  and  $\geq 0$ , use the `-s 0` option. Note, however, that this will lead to differences in the number of bins plotted on the left and right side of the matrix.





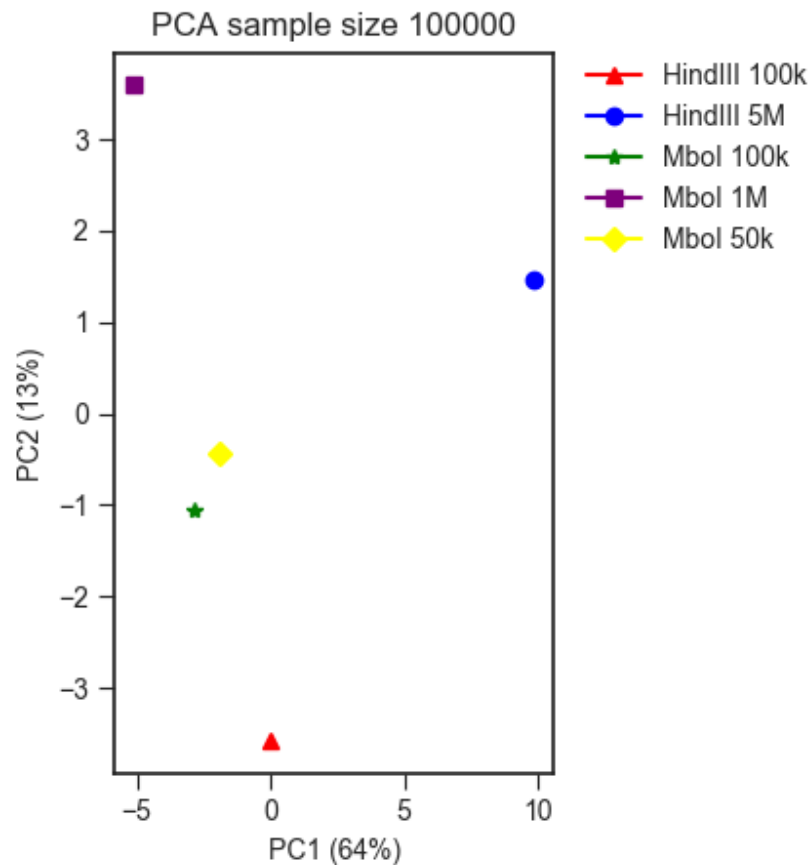
- (8) Compartment strength as defined by Flyamer, Gassler, and Imakaev et. al 2017 can be calculated using `--compartment-strength <filename>`. It is defined in their supplement as follows “[In a 5x5 compartment enrichment map,] “to calculate the strength of compartment signal, we took the natural logarithm of the  $AA * BB / AB^2$ ”.

### Practical 3: PCA Analysis

When working with multiple Hi-C libraries, it is often useful to assess the variability between replicates and samples from different conditions. This can provide valuable information about potential experimental biases and whether samples from different replicates can be safely merged.

As an example, we ran a PCA analysis on the 1mb resolution mESC Hi-C matrices from our Low-C paper (Díaz et al., Nat. Comms. 2018) using different restriction enzymes (MboI and HindIII), as well as different input cell numbers.

```
fanc pca -n "HindIII 100k" "HindIII 5M" \
  "MboI 100k" "MboI 1M" "MboI 50k" \
  -Z -s 100000 -r chr19 -p architecture/pca/lowc.pca.png \
  architecture/other-hic/lowc_hindiii_100k_1mb.hic \
  architecture/other-hic/lowc_hindiii_5M_1mb.hic \
  architecture/other-hic/lowc_mboi_100k_1mb.hic \
  architecture/other-hic/lowc_mboi_1M_1mb.hic \
  architecture/other-hic/lowc_mboi_50k_1mb.hic \
  architecture/pca/lowc.pca
```



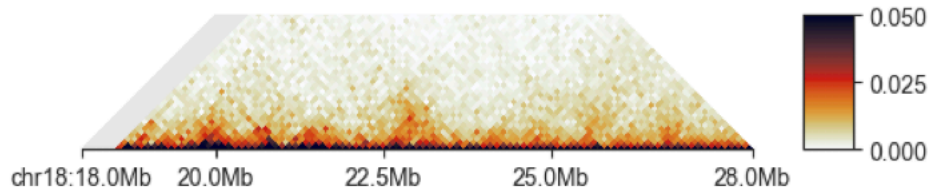
By default, PCA is run on pixels across the whole genome. In the example above, we have restricted the analysis to chromosome 19 using the `-r chr19` argument. `-Z` instructs `fanc pca` to use only non-zero matrix entries for the PCA - this can help mitigate the effect of very weak contacts on the variability.

Is run by default on the `s` most variable pixels in the Hi-C matrix. You can change which pixels get used for the PCA by changing `--strategy fold-change` to choose contacts with the largest fold-change or `--strategy passthrough` to make no prior selection of contacts. If you only want to include contacts up to (or above a) a certain distance, you can specify that distance using the `--max-distance` (or `--min-distance`) option.

## Practical 4: TAD boundaries and TADs

Like compartments, topologically associating domains, or TADs, for a fundamental level of genome organisation.

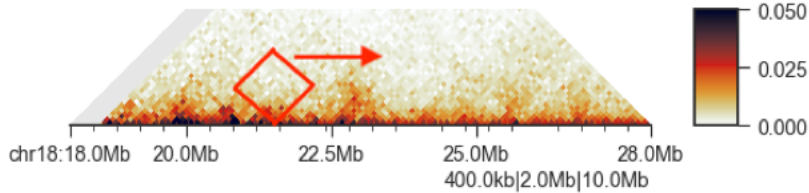
```
fancplot \
-o architecture/domains/fanc_example_100kb_tads.png \
chr18:18mb-28mb \
-p triangular output/hic/binned/fanc_example_100kb.hic \
-m 4000000 -vmin 0 -vmax 0.05
```



FAN-C provides multiple “scores” that are designed to find the boundaries between domains.

## Insulation Score

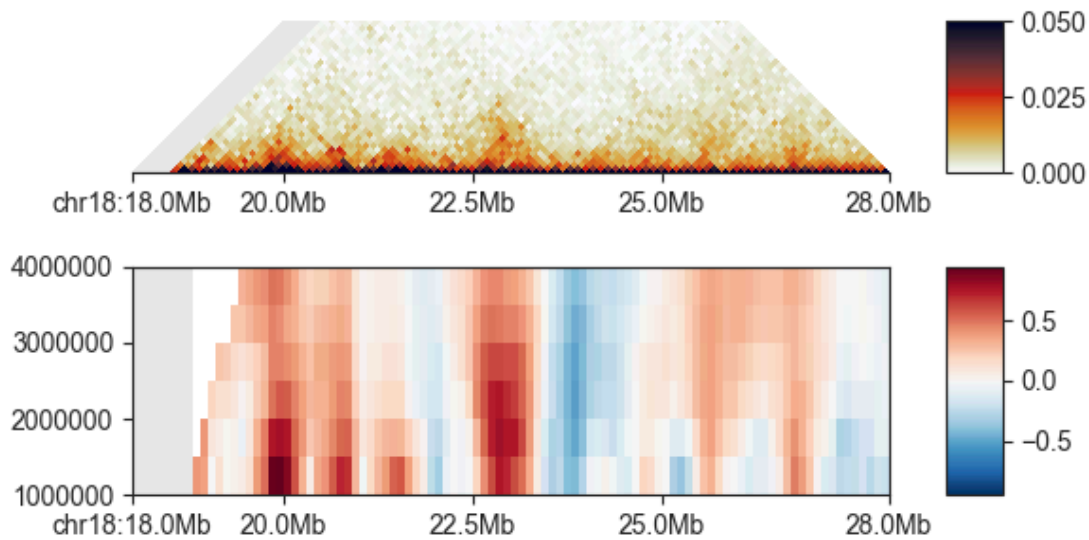
The insulation score (Crane et al. 2015) adds up contacts in a sliding diamond window along the Hi-C matrix diagonal.



Regions with low score are “insulating”, i.e. regions between domains. Regions with high scores are most likely found inside domains.

- (1) `fanc insulation` is typically used to calculate insulation scores with multiple window sizes at the same time, as a single window size might be prone to local matrix differences:

```
fanc insulation output/hic/binned/fanc_example_100kb.hic \
architecture/domains/fanc_example_100kb.insulation \
-w 1000000 1500000 2000000 2500000 3000000 3500000 4000000
```



(2) We can easily plot all insulation scores at the same time using `fancplot`:

```
fancplot \
-o architecture/domains/fanc_example_50kb_tads_insulation.png \
chr18:18mb-28mb \
-p triangular output/hic/binned/fanc_example_100kb.hic \
-m 4000000 -vmin 0 -vmax 0.05 \
-p scores architecture/domains/fanc_example_100kb.insulation
```

(3) We can additionally obtain `.bed` files, `.gff` files or `.bigwig` files for display in other genome browsers:

```
fanc insulation output/hic/binned/fanc_example_100kb.hic \
architecture/domains/fanc_example_100kb.insulation \
-w 1000000 1500000 2000000 2500000 3000000 3500000 4000000 \
-o bed
```

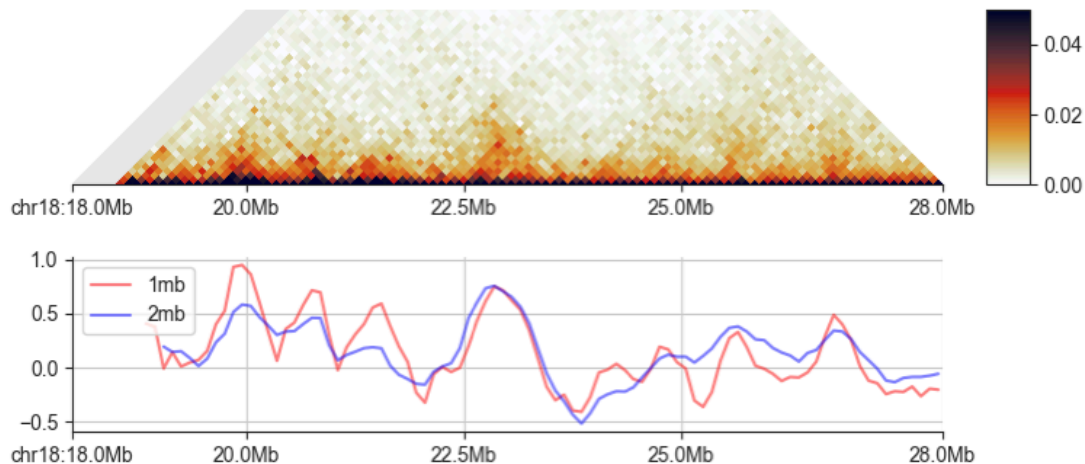
Which will generate multiple bed files:

```
ls architecture/domains/*.bed
architecture/domains/fanc_example_100kb.insulation_1.5mb.bed
architecture/domains/fanc_example_100kb.insulation_1mb.bed
architecture/domains/fanc_example_100kb.insulation_2.5mb.bed
architecture/domains/fanc_example_100kb.insulation_2mb.bed
architecture/domains/fanc_example_100kb.insulation_3.5mb.bed
architecture/domains/fanc_example_100kb.insulation_3mb.bed
architecture/domains/fanc_example_100kb.insulation_4mb.bed
```

(4) You can then plot scores from one or more window sizes using the line plot in `fancplot`:

```
fancplot --width 6 \
-o architecture/domains/fanc_example_100kb_tads_insulation_1mb.png \
chr18:18mb-28mb \
```

```
-p triangular output/hic/binned/fanc_example_100kb.hic \
-m 4000000 \
-vmin 0 -vmax 0.05 \
-p line architecture/domains/fanc_example_100kb.insulation_1mb.bed \
  architecture/domains/fanc_example_100kb.insulation_2mb.bed \
-l "1mb" "2mb"
```



By default, insulation scores are normalised to the chromosomal average and then log-transformed. There are plenty of normalisation and imputation strategies for insulation index calculations. Please refer to our Normalisation help for further details.

## Insulating boundaries (TAD boundaries)

Regions in the genome where the insulation score reaches a local minimum represent the region between two self-interacting domains, or TADs. You can use fanc boundaries to identify these regions.

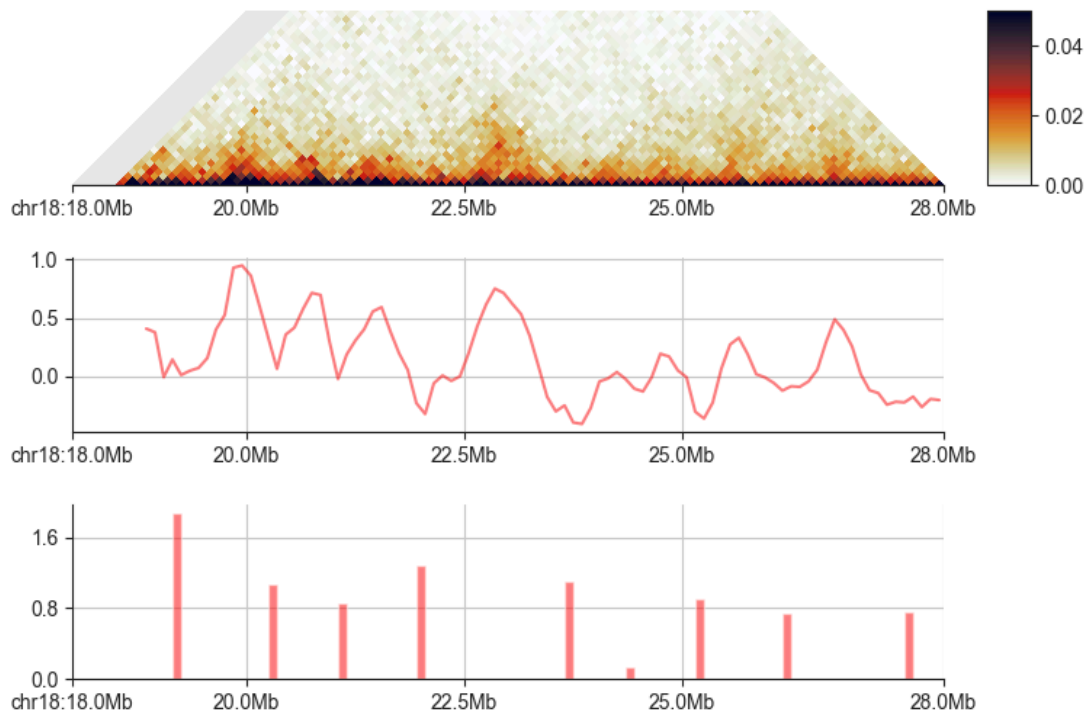
(5) When we run fanc boundaries on the above example using 1mb and 2mb as the window sizes:

```
fanc boundaries \
  architecture/domains/fanc_example_100kb.insulation \
  architecture/domains/fanc_example_100kb.insulation_boundaries \
-w 1mb 2mb
```

We get two output files with all insulation score minima and associated scores (the depth of the minimum compared to the two neighbouring maxima).

(6) Let's plot the boundaries from the 1mb scores:

```
fancplot --width 6 \
-o architecture/domains/fanc_example_100kb_insulation_1mb_boundaries.png \
chr18:18mb-28mb \
-p triangular output/hic/binned/fanc_example_100kb.hic \
-m 4000000 -vmin 0 -vmax 0.05 \
-p line architecture/domains/fanc_example_100kb.insulation_1mb.bed \
-l "1mb" \
-p bar architecture/domains/fanc_example_100kb.insulation_boundaries_1mb.bed
```



- (7) As you can see, lower minima get higher scores. By default, fanc boundaries outputs all minima, but you may set a threshold using `--min-score <s>` to report only boundaries with scores greater than `s`.

```
fanc boundaries \
  architecture/domains/fanc_example_100kb.insulation \
  architecture/domains/fanc_example_100kb.insulation_boundaries_score0.7 \
  -w 1mb 2mb --min-score 0.7
```

Note that the output of this analysis depends on the window size and thresholds that you decide to use. We recommend trying a few combinations and only use a set of ‘reproducible’ boundaries to avoid technical issues. In general, we also recommend annotating the boundaries with the respective insulation score to be able to compare quantitatively the scores called with the same window size and thresholds across conditions.

## Directionality Index

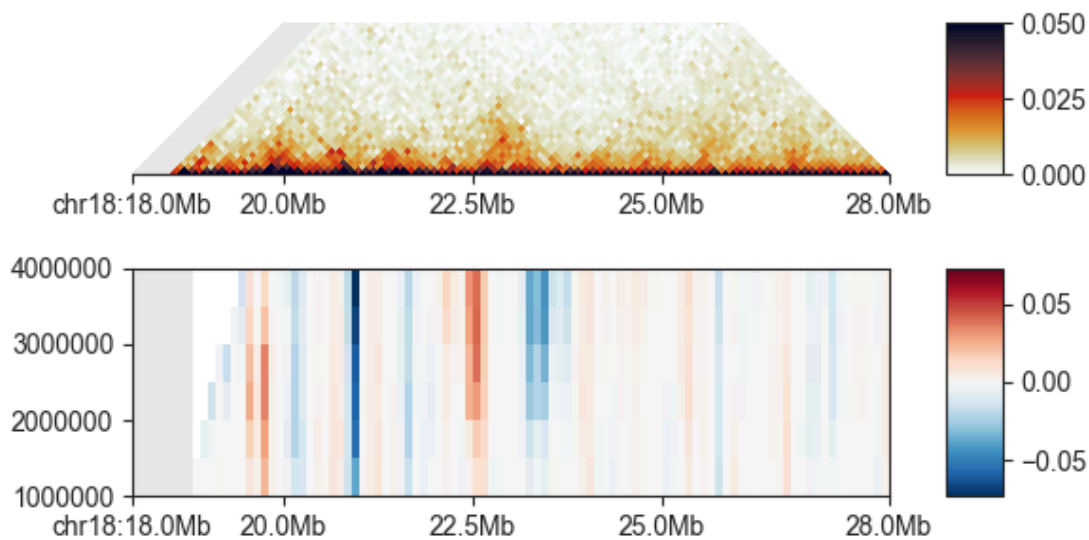
The directionality index (Dixon et al. 2012) measures the bias in contact frequency up- and downstream of an Hi-C region. When inside TADs, this measure tends towards zero, as interactions in either direction are equally frequent. However, when approaching a TAD boundary this measure changes drastically, as one direction will remain inside the TAD, where there is a high contact intensity, whereas the other direction will lie in a low intensity region outside the TAD.

- (8) **fanc directionality** is very similar in syntax to **fanc insulation**. It is typically used to calculate directionality indexes with multiple window sizes at the same time, as a single window size might be prone to local matrix differences:

```
fanc directionality \
  output/hic/binned/fanc_example_100kb.hic \
  architecture/domains/fanc_example_100kb.directionality \
  -w 1000000 1500000 2000000 2500000 3000000 3500000 4000000
```

(9) We can easily plot all directionality indexes at the same time using fancplot:

```
fancplot \
  -o architecture/domains/fanc_example_100kb_tads_directionality.png \
  chr18:18mb-28mb \
  -p triangular output/hic/binned/fanc_example_100kb.hic \
  -m 4000000 -vmin 0 -vmax 0.05 \
  -p scores architecture/domains/fanc_example_100kb.directionality
```



## TAD calling

FAN-C does not provide a TAD calling functionality itself as there are plenty of tools available that specialise on this topic. We ourselves have developed a tool to call TADs called *tadtool* with an interactive display for users to have the opportunity to decide on their parameters based on what looks appropriate visually.

However, we would like to point out that TAD calling algorithms often depend critically on their input parameters, and different TAD callers can lead to very different results. So, we encourage the use of multiple TAD calls or TADs called with multiple parameter combinations with a quantitative annotation of TAD strength.

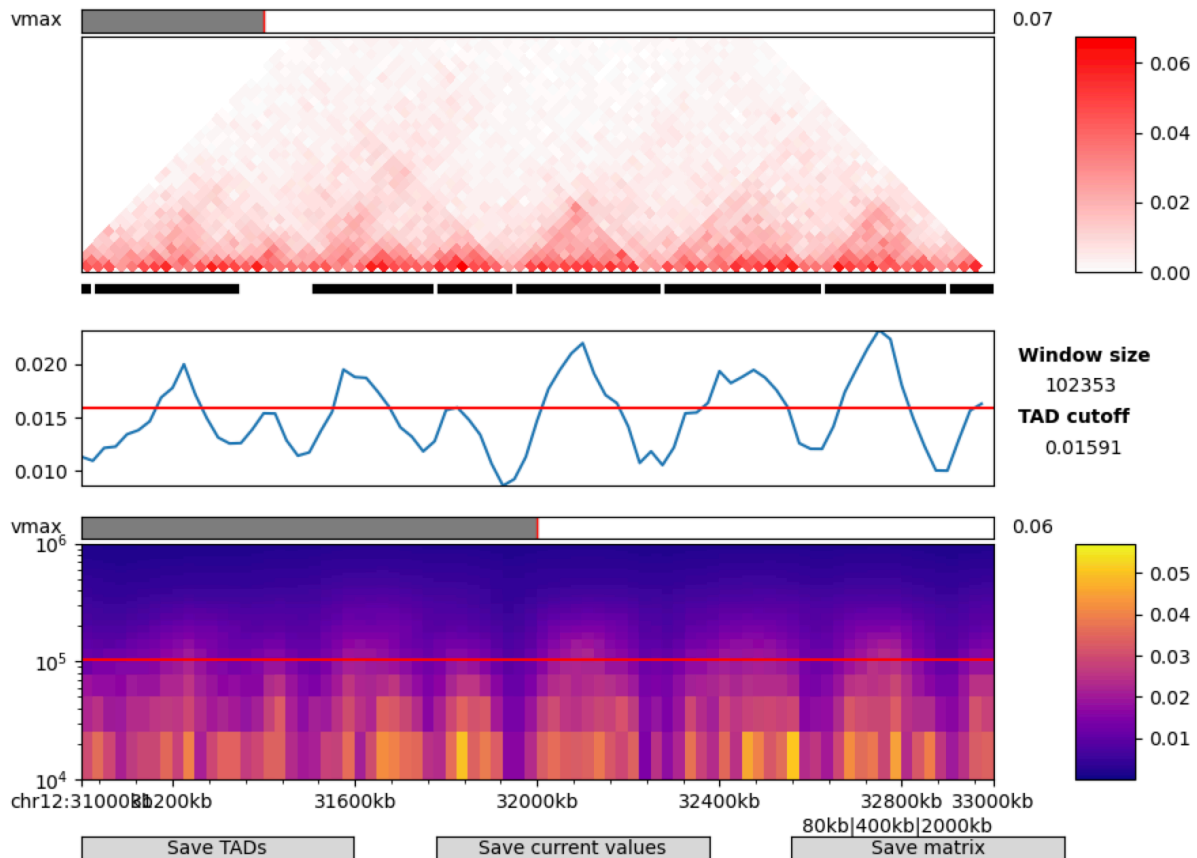
(10) To explore what's the effect of different parameters on TAD calling using *tadtool* we can use this example:

```
# Installing TADtool
git clone https://github.com/vaquerizaslab/tadtool.git
cd tadtool
```

```
python setup.py install
tadtool -help
```

*# Example locus*

```
tadtool plot \
  examples/chr12_20-35Mb.matrix.txt \
  examples/chr12_20-35Mb_regions.bed \
  chr12:31000000-33000000
```

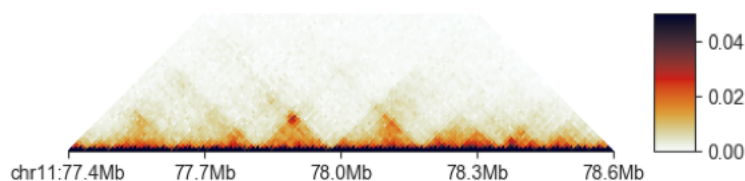


## Practical 5: Loop calling

Loops frequently form between two genomic regions, and are visible in the Hi-C matrix as patches of increased contact intensity:

```
fancplot \
  -o architecture/loops/rao2014.chr11_77400000_78600000.png \
  chr11:77400000-78600000 \
  -p triangular architecture/loops/rao2014.chr11_77400000_78600000.hic \
  -vmin 0.0 -vmax 0.05 -m 600000
```





We can use **fanc loops** to call loops in Hi-C matrices using the HICCUPS algorithm (Rao and Huntley et al., 2014). Please refer to the original paper for details on the algorithm, specifically the different types of local neighbourhoods defined to make loop calling robust.

In FAN-C this process consists of three steps:

- Annotating pixels for loop calling
- Filtering annotated pixels
- Merging unfiltered pixels into loops

## Annotating pixels for loop calling

- (1) The first step in HICCUPS consists of annotating each pixel with various measures related to their loop probability. The most important ones are: Enrichment over expected values in the local neighbourhood  
False Discovery Rate (FDR) of the local enrichment  
Mappability of the local neighbourhood

```
fanc loops \
  architecture/loops/rao2014.chr11_77400000_78600000.hic \
  architecture/loops/rao2014.chr11_77400000_78600000.loops \
  -t 2
```

This is the most computationally expensive step. We recommend large number of threads using the **-t** option. If you have access to a computational cluster running Sun/Oracle Grid Engine, you can take advantage of it by automatic job submission setting the **--sge** flag.

By default, **fanc loops** assumes a loop size of 25kb. This determines the area around a pixel that is not included in the local neighbourhood calculations. If this is chosen too small, the neighbourhood will lie within the peak region, and enrichments are going to be lower. If this is chosen too big, the neighbourhood will no longer be local. If you have reason to believe your loops size differs from the default, you can set it explicitly with **-p**.

Similarly, the width of the neighbourhood is determined as **p + 3** by default. If you want to increase or decrease the neighbourhood width, use the **-w** parameter. You should know, however, that this is just a starting value, and the neighbourhood width might be increased on demand internally.

Finally, you can control the size of the submatrices sent to each thread using the **--batch-size** parameter. The default, 200, should suit most purposes, but if your individual jobs are taking too long, you should reduce this number.

We can now use the output object with annotated pixels for downstream processing.

## Filtering annotated pixels

We need to apply filters to the annotated pixel object that remove all pixels with a low probability of being a loop. These filters typically consist of enrichment filters, FDR filters, and mappability filters. Additionally,

there are filters for minimum distance between regions, and the minimum number of unnormalised valid pairs in a pixel.

You can either set a global enrichment filter that acts on all neighbourhoods using `-e`, or choose individual thresholds for each local neighbourhood with `--enrichment-donut`, `--enrichment-vertical`, `--enrichment-horizontal`, and `--enrichment-lower-left`. You usually want to set at least the `--enrichment-donut` cut-off to something like 2.

For FDR values, also called q-values, you can set a global filter using `-q`. Control the filtering of individual neighbourhoods using `--fdr-donut`, `--fdr-vertical`, `--fdr-horizontal`, and `--fdr-lower-left`. Typical values for each neighbourhood are around 0.1.

Mappability filters act on pixels where a certain fraction of pixels in their local neighbourhoods is unmappable. To set a global mappability cut-off for all neighbourhoods, use the `-m` option. Again, local neighbourhood mappability filters can be fine-tuned using the `--mappability-donut`, `--mappability-vertical`, `--mappability-horizontal`, and `--mappability-lower-left` options.

It is generally a good idea to filter on the minimum distance between regions to consider forming a loop, as a lot of false positive loops will be close to the diagonal. You can use the `-d <b>` parameter to set a threshold on the minimum distance, where `b` is expressed in number of bins.

In addition, we highly recommend applying a filter on the minimum number of valid pairs in a pixel (`-o`), so that false-positive loops due to noise are avoided.

For convenience, we have included the filter applied by Rao and Huntley et al. 2014 in their original HICCUPS algorithm as a convenient pre-set `--rh-filter`. It only retains peaks that are at least 2-fold enriched over either the donut or lower-left neighbourhood, at least 1.5-fold enriched over the horizontal and vertical neighbourhoods, at least 1.75-fold enriched over both the donut and lower-left neighbourhood, and have an FDR  $\leq 0.1$  in every neighbourhood.

(2) Using the filters defined by Rao and Huntley et al, 2014

```
fanc loops \  
  architecture/loops/rao2014.chr11_77400000_78600000.loops \  
  architecture/loops/rao2014.chr11_77400000_78600000_filtered.loops \  
  --rh-filter -d 5 -o 5
```

## Merging unfiltered pixels into loops

(3) Pixels that pass all filtering steps are good candidates for loops. Often, these pixels appear in clusters, which we merge/join in this step. Pixels that do not form a cluster are generally false-positives, so we filter them using `--remove-singlets`.

```
fanc loops \  
  architecture/loops/rao2014.chr11_77400000_78600000_filtered.loops \  
  architecture/loops/rao2014.chr11_77400000_78600000_merged.loops \  
  -j --remove-singlets
```

(4) Finally, we can export all the merged loops to BEDPE format using `-b`:

```
fanc loops \  
  architecture/loops/rao2014.chr11_77400000_78600000_merged.loops \  
  -b architecture/loops/rao2014.chr11_77400000_78600000_merged.bedpe
```

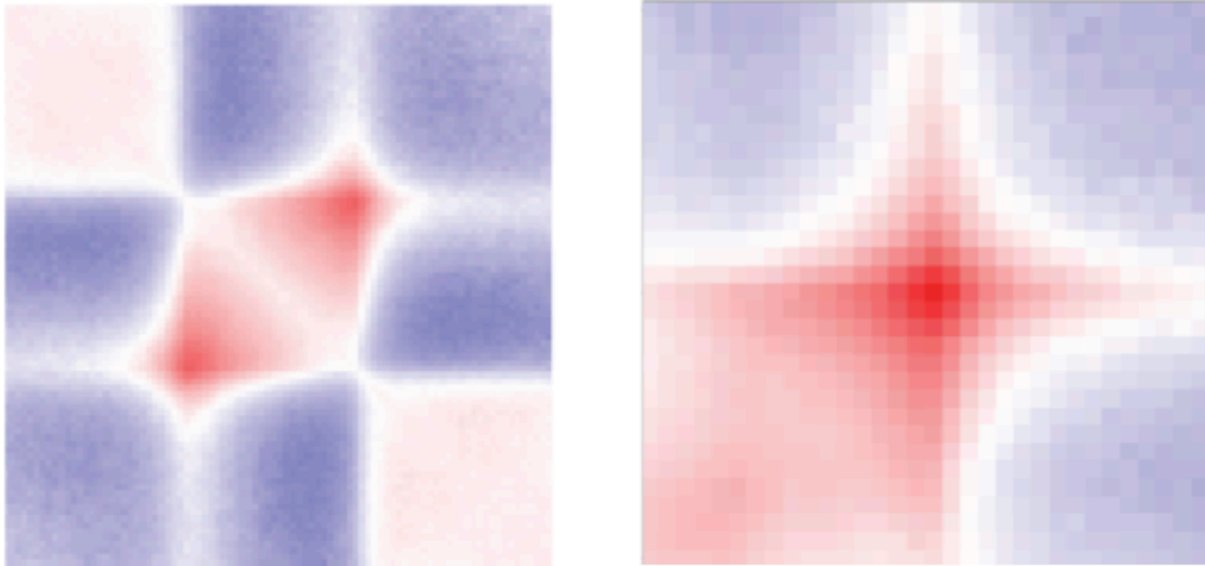
## Practical 8: Hi-C aggregate analysis

Often the resolution of our Hi-C matrix is not high enough to see very obvious changes between conditions when looking at a single locus. However, in a similar way that one can aggregate the ChIP-seq, Cut&Run or

ATAC-seq signal across many features in the genome such as promoters, enhancers, etc. One can aggregate the 2D matrix Hi-C signal over multiple regions of the genome in order to visualise the aggregate change of a series of regions together.

This can be done on paired coordinates such as those stored in BEDPE format, like enhancer-promoter loops, Hi-C peaks, etc. Or it can also be done on lists of 1D regions like those in BED files: TAD boundaries, Enhancers, Promoters, etc. For lists of regions, the aggregate matrix will be located at the Hi-C matrix diagonal. For pairs of regions, matrix subsets can be anywhere in the genome.

Here are examples of TAD and loop aggregate plots from a preprint from the lab (Kruse et al. 2019):



## Aggregate over variable width regions

By default, if you provide fanc aggregate with a list of regions, it will extract the square Hi-C sub-matrices along the diagonal for each region and interpolate them to match the width set by `--pixels` (90 by default). It will then calculate the average value for each pixel, which then form the aggregate matrix.

- (1) Let's try this on TADs called using the arrowhead algorithm (Rao and Huntley et al., 2014). `fanc aggregate` will ignore all regions in the file that are not present in the Hi-C matrix. In our example Hi-C file, that is everything outside of chromosomes 18 and 19:

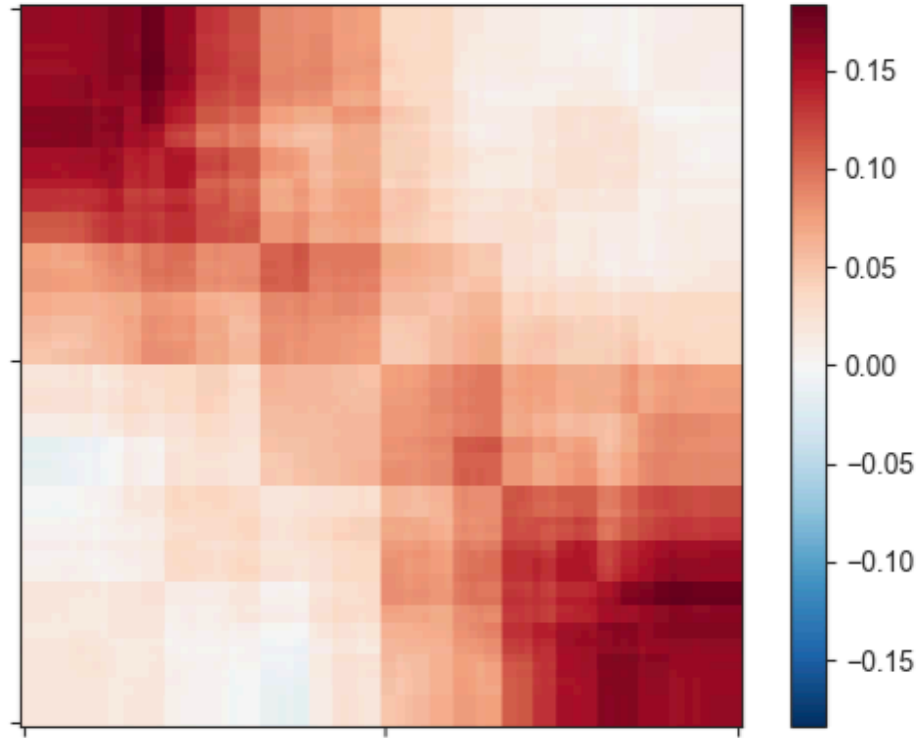
```
fanc aggregate output/hic/bin/binned/fanc_example_100kb.hic \
  architecture/domains/gm12878_tads.bed \
  architecture/aggregate/fanc_example_100kb.agg
```

This command only produces an `AggregateMatrix` file (`fanc_example_100kb.agg`), which is useful for further usage with FAN-C, but not easily readable.

- (2) To extract the aggregate matrix in `.txt` format, simply add `-m` and to plot it just use `-p`:

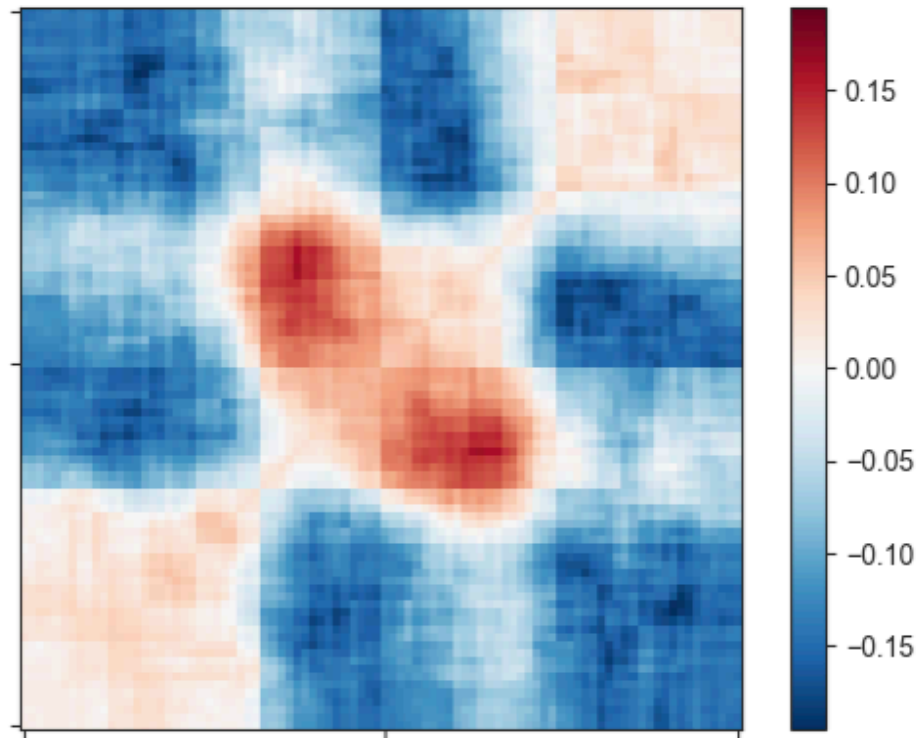
```
fanc aggregate output/hic/bin/binned/fanc_example_100kb.hic \
  architecture/domains/gm12878_tads.bed \
  architecture/aggregate/fanc_example_100kb.agg \
  -p architecture/aggregate/fanc_example_100kb.agg.png \
  -m architecture/aggregate/fanc_example_100kb.agg.txt \
  -e -l
```

Important: For variable sized regions, make sure to use the observed/expected regions by `-e` flag. `-e` works very well with log2-transformed data (-1).



- (3) This still does not look like much of a TAD, but we can add a little more context by expanding the plotting region relative to the region size using `-r`:

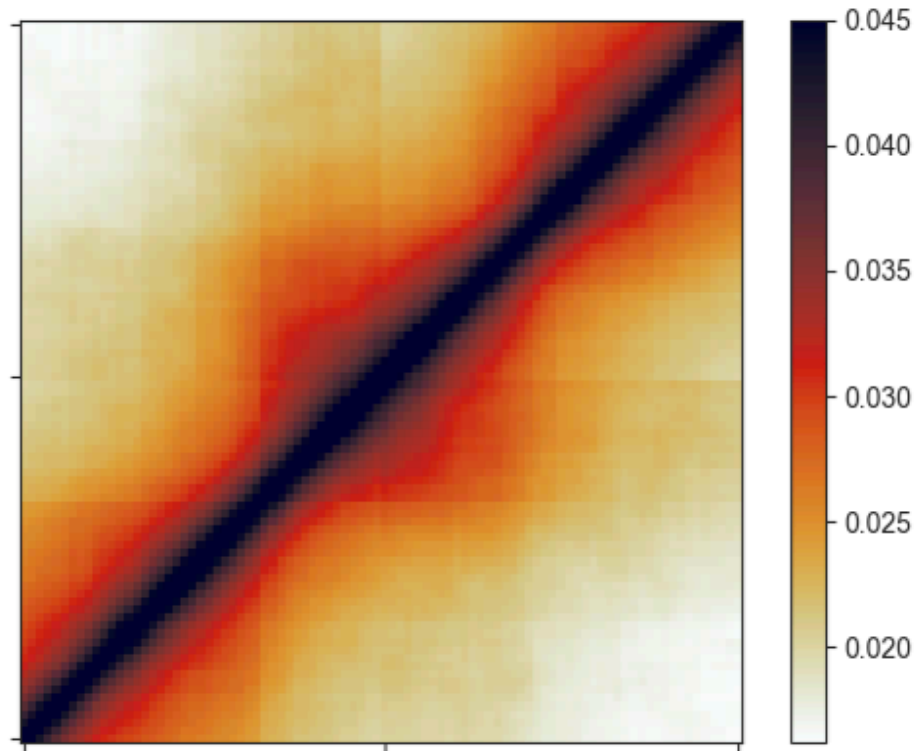
```
fanc aggregate output/hic/bin/binned/fanc_example_100kb.hic \  
  architecture/domains/gm12878_tads.bed \  
  architecture/aggregate/fanc_example_100kb.agg \  
-p architecture/aggregate/fanc_example_100kb_oe_large.agg.png \  
-m architecture/aggregate/fanc_example_100kb_oe_large.agg.txt \  
-e -l -r 1.0
```



That plot depicts a region that is 3x the size of the TAD located in its center and already looks like we would expect: High signal in the center, especially at the TAD corner, where the corner loops are typically located.

- (4) We can further apply an exponential rescaling (`--rescale`) of the data to make this look more like a Hi-C matrix. Here, we are not log-transforming the data and we are setting the saturation of the pixel values at 0.045 using `--vmax`.

```
fanc aggregate output/hic/bin/binned/fanc_example_100kb.hic \
  architecture/domains/gm12878_tads.bed \
  architecture/aggregate/fanc_example_100kb.agg \
  -p architecture/aggregate/fanc_example_100kb_oe_large_res.png \
  -m architecture/aggregate/fanc_example_100kb_oe_large.agg.txt \
  -e -r 1.0 --rescale --vmax 0.045
```



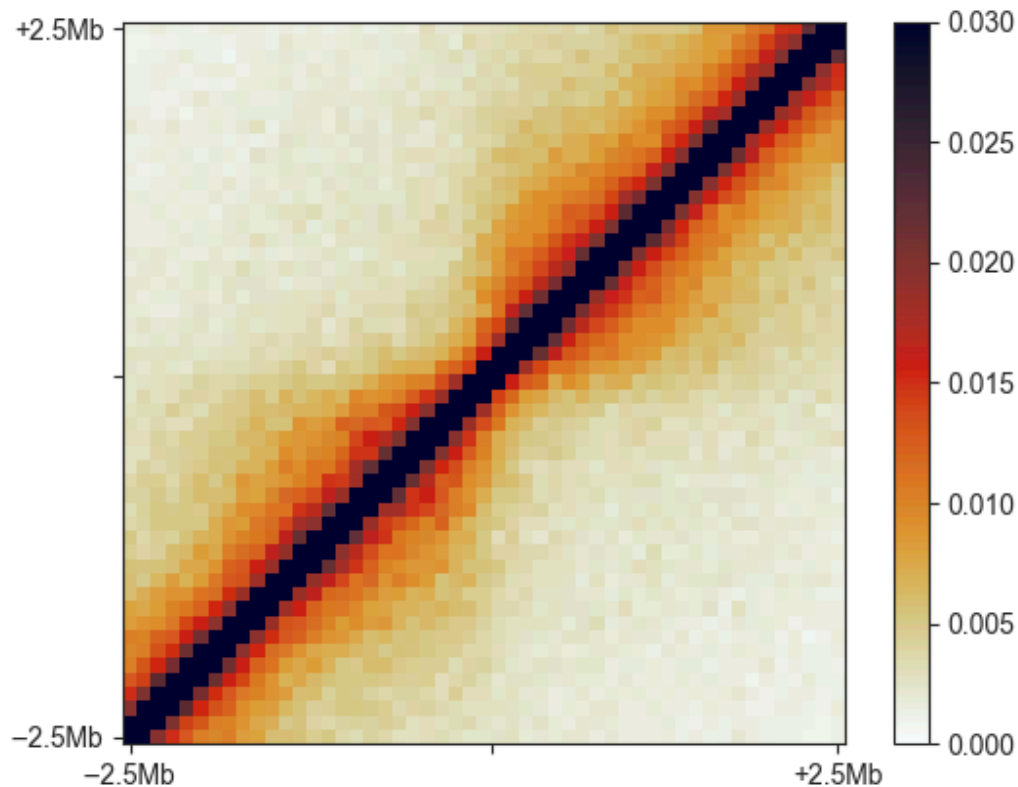
- (5) For both the  $\log_2(O/E)$  and rescaled versions of the aggregate matrices, there are pre-set flags you can use called `--tads` and `--tads-imakaev`, respectively. The latter is named after the first author of the publication that first used rescaled aggregate matrices in this fashion (Flyamer, Gassler, and Imakaev et al., 2017). In the above example, you can simply run:

```
fanc aggregate output/hic/bin/binned/fanc_example_100kb.hic \
  architecture/domains/gm12878_tads.bed \
  architecture/aggregate/fanc_example_100kb.agg \
  -p architecture/aggregate/fanc_example_100kb_oe_large.agg.png \
  -m architecture/aggregate/fanc_example_100kb_oe_large.agg.txt \
  --tads
```

## Aggregate over fixed width regions

- (6) Sometimes, you may want to use a fixed window surrounding a set of features in the aggregate analysis, such as TAD boundaries. `fanc aggregate` provides the `-w` option to plot the aggregate Hi-C matrix in a window of size `w` around the centre of each region in the list provided.

```
fanc aggregate \
  output/hic/bin/binned/fanc_example_100kb.hic \
  architecture/domains/fanc_example_100kb.insulation_boundaries_score0.7_1mb.bed \
  architecture/aggregate/fanc_example_100kb_boundaries.agg \
  -w 5mb -p architecture/aggregate/fanc_example_100kb_boundaries.agg.png \
  -m architecture/aggregate/fanc_example_100kb_boundaries.agg.txt \
  --vmin 0 --vmax 0.03
```



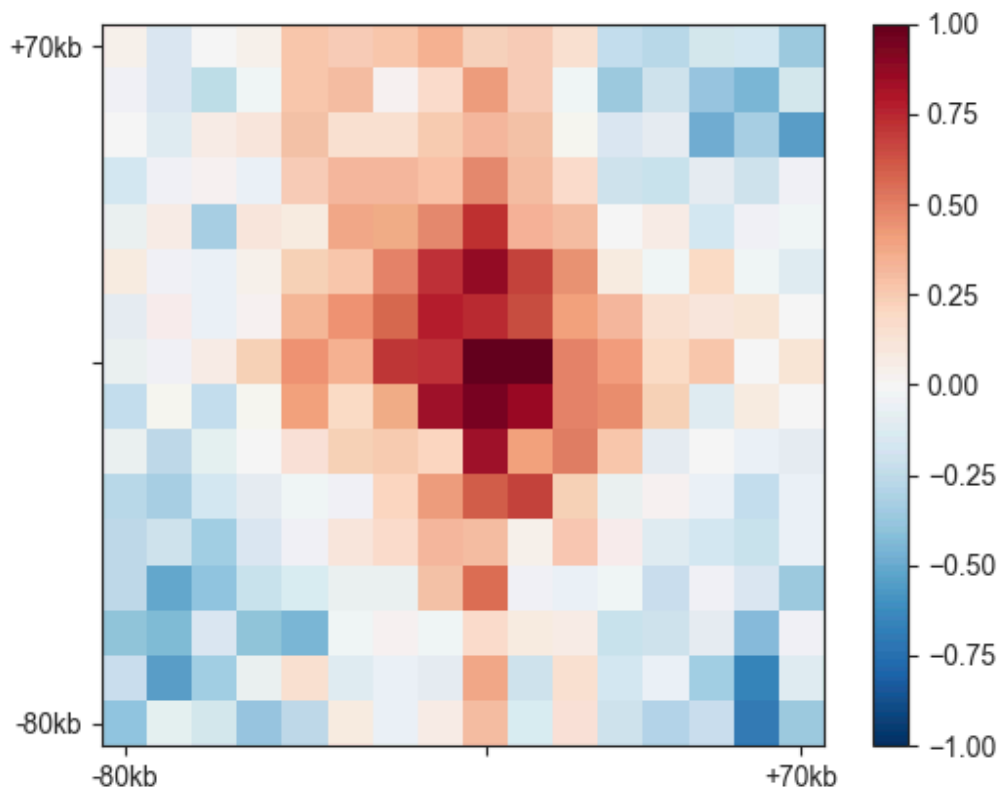
- (7) You can see the relatively faint “average boundary” in the centre of the plot. When using O/E and log2-transformed matrices, this becomes much more obvious:

```
fanc aggregate \
  output/hic/binmed/fanc_example_100kb.hic \
  architecture/domains/fanc_example_100kb.insulation_boundaries_score0.7_1mb.bed \
  architecture/aggregate/fanc_example_100kb_boundaries_oe.agg \
  -w 5mb -p architecture/aggregate/fanc_example_100kb_boundaries_oe.agg.png \
  -m architecture/aggregate/fanc_example_100kb_boundaries_oe.agg.txt \
  -e -1
```

## Loops and other pairwise genomic regions

- (6) When you have loop calls or other pairwise genomic regions in BEDPE format, you can use **fanc aggregate** to make aggregate loop plots. The pre-set for this is **--loops**. Control the size of the plot using the **--pixels** argument.

```
fanc aggregate architecture/loops/rao2014.chr11_77400000_78600000.hic \
  architecture/loops/rao2014.chr11_77400000_78600000.loops_no_singlets.bedpe \
  architecture/loops/rao2014.chr11_77400000_78600000.loops_no_singlets.agg \
  -p architecture/aggregate/rao2014.chr11_77400000_78600000.loops_no_singlets.agg.png \
  --loops
```



## Disclaimer

This tutorial is heavily based on the FAN-C documentation available at:  
<https://fan-c.readthedocs.io/>