# Build and run Saturn DREAM-ENCODE project

```
Clone the repository
git clone git@github.com:JohnReid/Saturn.git
Create and active a python virtual environment
cd Saturn/python
bin/create-virtualenv-py2
. ./activate-py2
pip install -r reqd-pkgs.txt
pip install -r reqd-pkgs-py2.txt
Create a data directory
cd ..
mkdir Data
Download the challenge data into the 'Data directory
annotations/hg19.genome.fa.gz
annotations/ladder regions.blacklistfiltered.bed.gz
annotations/ladder_regions.blacklistfiltered.merged.bed
annotations/ladder_regions.processed.bed
annotations/test_regions.blacklistfiltered.bed.gz
annotations/test regions.blacklistfiltered.merged.bed
annotations/train_regions.blacklistfiltered.bed.gz
annotations/train_regions.blacklistfiltered.merged.bed
DNASE/peaks/conservative/DNASE.A549.conservative.narrowPeak.gz
DNASE/peaks/conservative/DNASE.GM12878.conservative.narrowPeak.gz
DNASE/peaks/conservative/DNASE.H1-hESC.conservative.bed
DNASE/peaks/relaxed/DNASE.A549.relaxed.narrowPeak.gz
DNASE/peaks/relaxed/DNASE.GM12878.relaxed.narrowPeak.gz
DNASE/peaks/relaxed/DNASE.H1-hESC.relaxed.bed
DNASE/bams/DNASE.H1-hESC.biorep1.techrep1.bam
```

. . .

```
ChIPseq/labels/ARID3A.train.labels.tsv.gz
ChIPseq/labels/ATF2.train.labels.tsv.gz
ChIPseq/labels/ATF3.train.labels.tsv.gz
```

#### **DNase**

Create DNase features from the p-values of the relaxed peaks for each cell type by

```
scripts/feature-DNase.R <cell type>
Merge the DNase sequencing reads for each cell type by
merge-dnase <cell type>
Install Wellington.
```

pip install pyDNase

Run wellington on the relaxed peaks for each cell type by

run-wellington <cell type> relaxed

### Known motifs

Download the necessary known motif databases from the MEME suite website into the following locations

```
/etc/STEME/MEME-dbs/EUKARYOTE/jolma2010.meme
/etc/STEME/MEME-dbs/EUKARYOTE/jolma2013.meme
/etc/STEME/MEME-dbs/EUKARYOTE/SwissRegulon_human_and_mouse.meme
/etc/STEME/MEME-dbs/EUKARYOTE/zhao2011.meme
/etc/STEME/MEME-dbs/JASPAR/JASPAR_CORE_2016_vertebrates.meme
/etc/STEME/MEME-dbs/CIS-BP/Homo_sapiens.meme
```

Filter those motifs we are interested in.

KNOWNMOTIFSDIR=\$SATURNDIR/Data/Motifs/Known get-motifs >\$KNOWNMOTIFSDIR/tf-motifs.meme Install STEME<sup>1</sup>.

Scan the genome for the known motifs using STEME.

<sup>&</sup>lt;sup>1</sup>https://github.com/JohnReid/STEME

```
STEME_USE_GENOME_INDEX=1 steme-pwm-scan \
    --prediction-Z-threshold=.7 \
    --lambda=.001 \
    --cache-index \
    -o $KNOWNMOTIFSDIR \
    $SATURNDIR/Data/Motifs/Known/tf-motifs.meme \
$SATURNDIR/Data/annotations/hg19.chrs.fa
```

Create a set of features from the scan.

#### scripts/feature-scan.R \$KNOWNMOTIFSDIR Known

Create a set of features from the scan that are filtered by the Wellington footprints for each cell type.

```
scripts/feature-scan.R \
    --cell=<cell type 1> \
    --cell=<cell type2> \
    ... \
    --wellington \
$KNOWNMOTIFSDIR KnownWell
```

#### De novo motifs

Install DREME<sup>2</sup>.

Generate positive and negative sequences for each TF that DREME can discriminate between

```
fasta-get <TF>
```

Run DREME on the sequences for each TF (this can take up to 8 days on my machine)

```
tf-dreme <TF>
```

Scan the genome for each TF

\$SATURNDIR/python/bin/scan-genome \$SATURNDIR/Data/ChIPseq/seqs/DREME-<TF>

Create a set of features from the scans.

```
scripts/feature-scan.R --tf=<TF> $SATURNDIR/Data/ChIPseq/seqs/DREME-<TF> DREME
```

Create a set of features from the scan that are filtered by the Wellington footprints for each cell type relevant for each TF.

```
scripts/feature-scan.R \
   --tf=<TF> \
   --cell=<cell type 1> \
```

<sup>&</sup>lt;sup>2</sup>http://meme-suite.org/doc/download.html?man\_type=web

```
--cell=<cell type 2> \
... \
--wellington \
$SATURNDIR/Data/ChIPseq/seqs/DREME-<TF>
DREMEWell
```

Now all the features should be ready for prediction

## Predict

Run the prediction script on each TF/cell type combination of interest

```
scripts/predict.R \
  -f DNase \
  -f Known \
  -f KnownWell \
  -f DREME \
  -f DREMEWell \
  <TF> \
  <cell type>
```

Smooth the predictions using the TF-specific parameters given in \$SATURNDIR/R/smooth.R. Note we do not smooth the predictions for each TF.

```
scripts/predictions-smooth.R \
    --logodds \
    --width=<width> \
    --length-scale=<length scale> \
    $SATURNDIR/Data/Predictions/predictions-....tsv \
    <output file>
```