Bioinformatics Methods

Development of a CRISPR-based epigenetic screening method

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Chapter 1

Degenerate gRNA libraries

1.1 Finding all gRNA targets in the repeat-masked human genome

The genomic target sites of gRNAs designed for use with the *S.pyogenes* CRISPR/Cas system are of the general form $GN_{20}GG$. The presence of the G at the start of the motif is required for initiation of transcription from the U6 promoter, present on many gRNA expression vectors. This is followed by 19 random bases (N) that determine the gRNA target site - these bases comprise the gRNA protospacer. The protospacer sequence is followed by the protospacer-adjacent motif (PAM), which is not part of the gRNA sequence but is present in the genomic target sequence just 3' of the gRNA target site. For the *S.pyogenes* CRISPR/Cas system, the PAM has to be comprised of the bases NGG.

A gRNA library is a pooled collection of gRNAs. A random library has a complexity of 4¹⁹, which corresponds to 10¹² different sequences. The aim here is to reduce this complexity and maximise binding to the genome. To this end, I first identified the number of occurrences of the sequences GN20GG in the human genome. The Bioconductor (Bioconductor version 2.14) package BSgenome [16] was used to identify all sequences matching this pattern in the repeat-masked human genome in R (version 3.1.1). An R script named find_all_gRNAs.R contains the code and outputs a file containing chromosome coordinates and strand information for all hits, with the last column containing the pattern identifier. The script looks for the GN₂₀GG by default (and outputs a file named GN20GG_masked_allregions.txt), unless another pattern is supplied by the user as follows:

```
./find_all_gRNAs.R -p [pattern, e.g. "GTACN"], -n [pattern-name, e.g. "my-pattern"], -o [outputfilename, e.g. "All_GTACN_hg19_RM.txt"]
```

The regions mapping to chromosomes 1-22 and the sex chromosomes were extracted from the outputfile GN20GG_masked_allregions.txt as follows:

```
grep -v 'random' GN20GG_masked_allregions.txt | grep -v 'hap'
```

```
| grep -v 'chrUn' | grep -v chrM > GN20GG_masked_autoXY.txt
;

#remove the last column
awk '{print $1 "\t" $2 "\t" $3 "\t" $4}' GN20GG_masked_autoXY.
txt > GN20GG_masked_autoXY.bed;
```

1.2 Identifying gRNAs that overlap with known promoters

In order to generate an annotation file containing promoter regions, the annotation file for known transcripts (human GRCh37-70) was downloaded from Ensembl (version70) and parsed through a script supplied by Gareth Wilson. This script can be found here: https://github.com/regmgw1/regmgw1_scripts/blob/master/ensembl_scripts/transcript2promoter.pl). This script extracts coordinates -1000 and +500 bp from the start of the transcript. From this file non-overlapping promoter sequences were derived by strand-specific merging using the Bedtools suite (v2.17.0) [17].

Then, FASTA coordinates were retrieved using twoBitToFa [8], which is available from the UCSC website (http://hgdownload.cse.ucsc.edu/admin/exe/).

```
#Separate according to whether pattern is on plus or
#minus strand
grep '+' GN20GG_masked_autoXY_promoters_merged >
GN20GG_masked_autoXY_promoters_merged_PLUS;
grep '-' GN20GG_masked_autoXY_promoters_merged >
GN20GG_masked_autoXY_promoters_merged_MINUS
#make into a gff file
awk '{print $1 ":" ($2 - 1) "-" $3}'
   GN20GG_masked_autoXY_promoters_PLUS
GN20GG_masked_autoXY_promoters_PLUS.gff;
twoBitToFa yourpath2/human/GRCh37/hg19.2bit
GN20GG_masked_autoXY_promoters_merged_PLUS.fa
-seqList=GN2OGG_masked_autoXY_promoters_merged_PLUS.gff
#repeat for file containing hits on the minus strand
The Python script reverse_complement_fasta.py was used to reverse-complement
the FASTA sequences on the minus strand [4]. The script was invoked as follows:
python reverse_complement_fasta.py
   GN20GG_masked_autoXY_promoters_merged_MINUS.fa >
GN20GG_masked_autoXY_promoters_merged_MINUS_REVERSE_Complement
   .fa
The fasta files on the plus and minus strand were combined using the cat command,
lowercase letters converted to uppercase using the segret tool from EMBOSS [18],
and sequences collapsed into a unique set with fastx_collapser [1].
```

GN20GG_masked_autoXY_promoters_merged_MINUS_REVERSE_Complement

.fa > GN20GG_masked_autoXY_promoters_merged_TOTAL.fa

cat GN20GG_masked_autoXY_promoters_merged_PLUS.fa

#combine the files

```
#convert to uppercase
seqret GN20GG_masked_autoXY_promoters_merged_TOTAL.fa
GN20GG_masked_autoXY_promoters_merged_TOTAL_UPPER.fa -sformat
    fasta -supper Y

#get unique fasta sequences
fastx_collapser <
    GN20GG_masked_autoXY_promoters_merged_TOTAL_UPPER.fa >
GN20GG_masked_autoXY_promoters_merged_PlusMinus_UNIQUE.fa
```

This yields a file containing 4,113,530 sequences.

1.3 Identifying a consensus sequence for gRNAs that fall into promoters

The Bioconductor package Biostrings [15] was used to derive a consensus sequence from the list of FASTA sequences generated above as follows (run in R):

```
>library(Biostrings)
>promMINUS <-readDNAStringSet(
"GN20GG_masked_autoXY_promoter_minus_REVERSECOMPLEMENT.fa",
    format="fasta")
>fm<-consensusMatrix(promMINUS)
minus <-fm[1:4,]
pwm_minus <-t(t(minus)/rowSums(t(minus)))
The following code was used to generate sequence logo plots [6] in R.
>library(ggplot2)
```

```
>berrylogo <- function (pwm, gc_content = 0.5, zero = .0001) {
 backFreq <-list(A=(1-gc_content)/2,C=gc_content/2,G=gc_content
    /2,T=
  (1-gc_content)/2)
  pwm [pwm == 0] < -zero</pre>
bval <-plyr::laply(names(backFreq),function(x){log(pwm[x,])-log
  backFreq[[x]])})
row.names(bval)<-names(backFreq)</pre>
p<-ggplot2::ggplot(reshape2::melt(bval,varnames=c("nt","pos"))
  ggplot2::aes(x=pos,y=value,label=nt))+
    ggplot2::geom_abline(ggplot2::aes(slope=0), colour = "grey
       ", size=2) +
    ggplot2::geom_text(ggplot2::aes(colour=factor(nt)),size=8)
    ggplot2::theme(legend.position="none")+
    ggplot2::scale_x_continuous(name="Position",breaks=1:ncol(
       bval))+
    ggplot2::scale_y_continuous(name="Log relative frequency")
  return(p)
}
#invoke the function with:
berrylogo(pwm_minus, gc_content=0.5, zero=.0001)
```

1.4 Reducing complexity by identifying the most significant clusters

I reasoned that it might be possible to reduce the complexity of a random library by clustering. I defined regions of interest as the 4,671,728 genomic hits of the form

GN20GG that fall into or next to known promoter sequences in the human genome with a minimum of 1 bp overlap. I used LCS-HIT (Version 0.5.2) [14] to cluster those regions of interest on the basis of sequence similarity. Clusters were ranked in descending order by the number of members and the top 15 clusters extracted.

```
#cluster sequences based on sequence similarity threshold
# of 0.2 and use the "exact algorithm"
lcs_hit-0.5.21/lcs_hit -i GN20GG_masked_autoXY_promoters
_merged_PlusMinus_UNIQUE.fa -O LCSHIT_OUTPUT20G1 -c 0.2 -g 1 &
```

LCS-HIT outputs the FASTA-identifiers only, therefore FASTA sequences were extracted using the script RetrieveFasta.pl, downloaded from reference [3]. This step is exemplified here for the top cluster (Cluster190).

```
./RetrieveFasta.pl Cluster190
    GN20GG_masked_autoXY_promoters_merged
_PlusMinus_UNIQUE.fa > Cluster190.fa;
```

For each of the top 15 clusters consensus sequences were computed using the Bioconductor package Biostrings (run in R).

The threshold option allows the user to define the percentage threshold at which a given nucleotide will be incorporated into the consensus sequence at a given position. The threshold was varied between 0.14 and 0.25 so as to yield consensus sequences representing roughly 10^4 , 10^5 and 10^6 different sequences respectively. The complexity, or number of sequences represented by a cluster, was computed using a C script downloaded from [5] and named AllSequencesFromConsensus.c.

```
#compile with
```

```
gcc -o AllSequencesFromConsensus AllSequencesFromConsensus.c
```

```
#run the script and pipe the output to line-count (as
# shown for the consensus sequence of Cluster190_T20)
./AllSequencesFromConsensus RRRGRGRRRRGRRGRRGV | wc -1
```

Next, the "GenomeSearch" function of the Biostrings package (see Section 1.1) was used to run each of the consensus sequences for each of the top 15 clusters against the masked human genome.

Because gRNAs are known to tolerate mismatches in their target sites [12], I reasoned that counting only exact matches probably underestimates the true number of target sites of any given sequence. Thus, the analysis was repeated allowing for an arbitrary number of up to 3 mismatches (gRNAs are known to tolerate more mismatches, especially towards the 5' end of the protospacer sequence. However, position of the mismatch in the gRNA was not taken into account here).

```
#allowing for a maximum of three mismatches
#change relevant parameter in the "GenomeSearch" function
#of the code
>plus_matches <- matchPattern(pattern, subject, max.mismatch
   =3, min.mismatch=0, fixed=c(pattern=FALSE, subject=TRUE))
>runAnalysis_masked_3mismatch(dict0, outfile="
   Top15Clusters_masked_allregions_3mismatch.txt")
Output files from both scripts were processed as follows and the results stored in
Table 3.1 on p. 96 of the PhD thesis.
# retrieve hits for autosomes and sex chromosomes only
grep "Cluster190_T20" Top15Clusters_masked_allregions.txt
   grep -v 'random' | grep -v 'hap' | grep -v 'chrUn' | grep -
   v chrM | awk '{print $1 "\t" $2 "\t" $3}' >
   Cluster190_T20_autoXY
#count number of hits that fall into promoter regions
intersectBed -a Cluster190_T20_autoXY -b annotation_files/
   promoters_merged.bed -wa -wb | sortBed | wc -1;
#count the number of unique promoter regions hit
intersectBed -a Cluster190_T20_autoXY -b annotation_files/
   promoters_merged.bed -wb | cut -f 5-8 | sortBed | uniq | wc
    -1;
```

I defined the targeting efficiency for each of the possible consensus sequences of a given cluster by dividing the number of promoter hits by the total number of unique sequences (complexity) of the consensus. For each cluster the consensus sequences with the highest targeting sequence was chosen. The 15 top clusters were then ranked by targeting efficiency and the top 6 clusters chosen for library preparation.

Chapter 2

Data analysis: sequencing of the degenerate and MNase digest libraries

2.1 Read trimming and quality filtering

Addition of sequencing adapters was non-directional, which means the reads contain gRNA cassettes both in forward and reverse direction.

First, the gRNAs that were sequenced in the forward direction were extracted using cutadapt [13] to trim away the plasmid sequence (shown here for the fastq sequencing data file generated from the Random N19 library).

```
cutadapt -g
   AAGTATTTCGATTTCTTGGCTTTATATATCTTGTGGAAAGGACGAAACACC -0 41 -
   m 2 --untrimmed-output=Untrimmed255_R1_50F.fastq.gz -0 255-
   N19_R1_F50_trimmed.fastq.gz ../255-N19_S5_L001_R1_001.fastq.gz;

cutadapt -a GTTTTAGAGCTAGAAATAGCAAGTTAAAATAAGGCTAGTCCG -0 33 -
   m 2 --untrimmed-output=Untrimmed255_R1_19R.fastq.gz -0 255-
   N19_R1_F50+19R_trimmed.fastq.gz 255-N19_R1_F50_trimmed.
   fastq.gz;
```

Next, the reads stored in the untrimmed output, which will contain reads of the cassette sequenced in the reverse direction were concatenated, sorted and gRNA sequences extracted as follows (again showing example of reads from the N19 Random library):

```
cat Untrimmed255_R1_50F.fastq.gz Untrimmed255_R1_19R.fastq.gz
> Untrimmed255_R1_50F+19R.fastq.gz;

zcat Untrimmed255_R1_50F+19R.fastq.gz | paste - - - - | sort -
k1,1 -t " " | tr "\t" "\n" > Untrimmed255_R1_50F+19R_sorted
.fastq;

cutadapt -a
   GGTGTTTCGTCCTTTCCACAAGATATATAAAGCCAAGAAATCGAAATACTT -O 41 -
```

```
m 2 --untrimmed-output=Untrimmed255_50FRC.fastq.gz -o 255
   _R1_50FRC_trimmed.fastq.gz Untrimmed255_R1_50F+19R_sorted.
   fastq;
cutadapt -g CGGACTAGCCTTATTTTAACTTGCTATTTCTAGCTCTAAAAC -0 33 -
   m 2 --untrimmed-output=Untrimmed255_19RRC.fastq.gz -o 255
   _R1_50FRC+19RRC_trimmed.fastq.gz 255_R1_50FRC_trimmed.fastq
   .gz;
This results in extraction of gRNA sequences sequenced in the reverse direction.
### Double untrimmed reads:
cat Untrimmed255_50FRC.fastq.gz Untrimmed255_19RRC.fastq.gz >
   Untrimmed255_50FRC+19RRC.fastq.gz;
zcat Untrimmed255_50FRC+19RRC.fastq.gz | paste - - - | sort
   -k1,1 -t " " | tr "\t" \n > Untrimmed255_50FRC+19
   RRC_sorted.fastq;
### Successfully trimmed reads:
cat 255-N19_R1_F50+19R_trimmed.fastq.gz 255_R1_50FRC+19
   RRC_trimmed.fastq.gz > 255_R1_4waytrimmed.fastq.gz;
zcat 255_R1_4waytrimmed.fastq.gz | paste - - - - | sort -k1,1
   -t " " | tr "\t" \n" > 255_R1_4waytrimmed_sorted.fastq;
### Extracted gRNA sequences sequenced in reverse direction
   need to be reverse-complemented
zcat 255_R1_50FRC+19RRC_trimmed.fastq.gz |
```

fastx_reverse_complement -Q33 -z > 255_R1_50FRC+19

RRC_trimmed_onedirection.fastq.gz;

```
cat 255-N19_R1_F50+19R_trimmed.fastq.gz 255_R1_50FRC+19
    RRC_trimmed_onedirection.fastq.gz > 255
    _R1_4waytrimmed_onedirection.fastq.gz;

zcat 255_R1_4waytrimmed_onedirection.fastq.gz | paste - - - -
    | sort -k1,1 -t " " | tr "\t" "\n" > 255
    _R1_4waytrimmed_sorted_onedirection.fastq;
```

Next, read length distribution between 15 and 40 bp were plotted for each of the libraries (again shown here for the file containing reads from the N19 Random library):

Per base sequence quality after trimming was assessed using FASTQC [2]

```
fastqc 255_R1_4waytrimmed_sorted_smaller40.fastq --outdir=../
    FASTQC_Trimmedreads
```

2.2 Histogram of gRNA lengths

This code was used to generate the plots in Figure 3.12 on page 100 of the PhD thesis. Histograms of read lengths were generated in R:

```
par(mar=c(5,6+2,4,2)+1)
hist(reads[,1], breaks=seq(0,40,by=1), col="grey", main="
    Method 1 - Random N19", xlab="Insert length (bp)", ylim=c
    (0, 500000), las=1, ylab="", cex.main=1.3, cex.axis=1.3,
    cex.lab=1.3)
title(ylab = "Frequency", line = 6, cex.lab=1.3)
```

2.3 Histogram of read frequencies

This code was used to generate the plots in Figure 3.11 on page 99 of the PhD thesis.

```
awk '{if(NR%4==2) print}' 255
   _R1_4waytrimmed_sorted_onedirection_smaller40.fastq | sort
   | uniq -c | sort | sed 's/^* *// > 255
   _R1_4waytrimmed_sorted_onedirection_smaller40_sortedbyfrequency
#Histograms of read frequency were generated in R:
reads <-read.table("255
   _R1_4waytrimmed_sorted_onedirection_smaller40_sortedbyfrequency
   ", sep=" ", head=F)
colnames(reads)<-c("counts", "sequence")</pre>
reads_upto5 <-reads[reads$counts <=5,]
reads_greater5 <-reads[reads$counts>=5,]
par(oma=c(0,0,0,0))
par(mar=c(6,6,8,3))
par(fig=c(0.1,1,0,0.75))
plot(rownames(reads), reads$counts, type="p", xlab="gRNA
   ranked by counts", xlim = c(0, 600000), ylim = c(5,350), ylab
   ="", las=1, col="orange", lwd=4, cex.axis=1.7, cex.lab=1.9)
title(ylab = "Read counts", line = 4, cex.lab=1.9)
```

2.4 Alignment to the reference genome

Trimmed reads were aligned to the reference genome, either human (GRCh37/hg19) or mouse (NCBI37/mm9) as appropriate, using BWA [9], again shown using the file containing reads from the N19 Random library as an example.

```
#use bwa to align reads to human genome without allowing
   mismatches and printing all alignments, seed length is set
   to 19
```

```
bwa aln -n 0 -o 0 -l 19 -N /mnt/store2/local_data/genomic_data /human/GRCh37/human_GRCh37.tmp 255_R1_4waytrimmed_sorted. fastq > 255_R1_4waytrimmed.sai
```

```
#generate the sam file
bwa samse -n 10000 /path2/human_GRCh37.tmp 255_R1_4waytrimmed.
    sai 255_R1_4waytrimmed_sorted.fastq > 255_R1_4waytrimmed.
    sam
```

Uniquely mapped reads were counted using Samtools [10].

```
samtools view -S -q1 255_R1_4waytrimmed.sam | wc -l
```

The remainder of the analysis (parts of which are shown in Fig. 3.18 on page 111 of the thesis) was conducted by my collaborator Karolina Worf at the Helmholtz Insti-

tute in Munich. The documentation for this analysis is available at hmgubox (https://hmgubox.helmholtz-muenchen.de:8001/d/6c6e75236e/; password: Coralina).

Chapter 3

Design of the EMT5000 library

3.1 Defining regions of interest

A list of genes known to be involved in the regulation of epithelial-to-mesenchymal transition (EMT) was taken from DeCraene et al. [7]. Regions of interest within the promoter of these genes were identified manually by inspection of chromatin marks in UCSC genome browser (table 3.1).

Chromosome	Start	Stop	Gene
chr1	170626538	170637878	PRRX1
chr2	145272896	145282545	${ m ZEB2}$
chr2	145310788	145311630	ZEB2
chr6	166578775	166584033	Brachyury (T gene)
chr6	166586466	166588249	Brachyury (T gene)
chr7	19155427	19162115	TWIST1
chr8	49831094	49838789	SLUG
chr10	31549929	31552360	ZEB1
chr10	31603262	31611019	ZEB1
chr14	61113258	61126351	SIX1
chr14	95235188	95236645	GSC (Goosecoid)
chr16	68765472	68768900	Cdh1
chr16	68770501	68779468	Cdh1
chr16	86596822	86601033	FOXC2
chr18	25616470	25616815	$\mathrm{Cdh2}$
chr18	25753143	25759002	$\mathrm{Cdh2}$
chr18	25763319	25764152	$\mathrm{Cdh2}$
chr18	25783828	25784775	$\mathrm{Cdh2}$
chr18	52966479	52970132	TCF4
chr18	52983584	52991765	TCF4
chr18	52994740	52997201	TCF4
chr18	53067559	53071402	TCF4
chr18	53072594	53073776	TCF4
chr18	53087724	53090359	TCF4
chr18	53176467	53178784	TCF4
chr18	53252816	53257791	TCF4
chr18	53259747	53260381	TCF4
chr18	53301547	53303603	TCF4
chr19	1631468	1633670	E47/TCF3
chr19	1646514	1653855	E47/TCF3
chr19	1655335	1656204	E47/TCF3
chr19	1660918	1661677	E47/TCF3
chr20	48592707	48600991	SNAIL1
chrX	56258091	56260688	KLF8

Table 3.1: Regions of interest around the promoter regions of 15 genes known to be involved in the regulation of EMT [7]

3.2 Identifying all gRNAs that fall into regions of interest

The genomic regions listed in **table 3.1** where chosen as target sites for the design of gRNAs and saved in the file EMT_genepromoter_comprehensive.gff. All gRNAs falling into these regions were then found using the Bedtools [17] intersect function on the file containing all gRNAs with an NGG PAM found in the genome (see section 1.1 for how this file was generated).

```
intersectBed -a GN20GG_masked_autoXY.gff -b
EMT_genepromoter_comprehensive.gff -f 1 -wa -wb >
GN20GG_masked_autoXY_EMT_genepromoter_comprehensive.gff
```

3.3 Subsetting only gRNAs that map uniquely to the human genome

Guide RNAs were then aligned to the genome in order to identify those that map uniquely. Before alignment, files had to be converted into the correct format as follows:

```
##for alignment need to first convert file into fasta format
##separate + and - strand
cut -f 1,2,3,4
    GN20GG_masked_autoXY_EMT_genepromoter_comprehensive.gff |
    grep '+' | awk '{print "chr" $1 ":" ($2-1) "-" $3}' >
    GN20GG_masked_autoXY_EMT_genepromoter_comprehensive_PLUS.2
    bit;

cut -f 1,2,3,4
    GN20GG_masked_autoXY_EMT_genepromoter_comprehensive.gff |
    grep -v '+' | awk '{print "chr" $1 ":" $2 "-" $3}' >
```

```
GN20GG_masked_autoXY_EMT_genepromoter_comprehensive_MINUS.2
  bit;
##convert to FASTA format using twoBitToFa
twoBitToFa /path2/human/GRCh37/hg19.2bit
   GN20GG_masked_autoXY_EMT_genepromoter_comprehensive_PLUS.fa
    seqList=
   GN20GG_masked_autoXY_EMT_genepromoter_comprehensive_PLUS.2
   bit:
twoBitToFa /path2/human/GRCh37/hg19.2bit
   {\tt GN20GG\_masked\_autoXY\_EMT\_genepromoter\_comprehensive\_MINUS}.
   fa seqList=
   GN20GG_masked_autoXY_EMT_genepromoter_comprehensive_MINUS.2
   bit;
##create reverse complement of guide RNAs on the minus
##strand using the script reverse_complement_fasta.py
python reverse_complement_fasta.py
   GN20GG_masked_autoXY_EMT_genepromoter_comprehensive_MINUS.
   GN20GG_masked_autoXY_EMT_genepromoter_comprehensive_MINUS_RC
   .fa;
##combine the files for the + and - strand
cat GN20GG_masked_autoXY_EMT_genepromoter_comprehensive_PLUS.
   fa
   GN20GG_masked_autoXY_EMT_genepromoter_comprehensive_MINUS_RC
   .fa > GN20GG_masked_autoXY_EMT_genepromoter_comprehensive.
   fa;
```

The PAM sequence was removed using trimming.py and alignment to the genome

```
without the PAM (i.e. as GN19 instead of GN20GG).
```

```
bwa aln -n 0 -o 0 -l 10 -N -I ~/path_to/GRCh37/human_GRCh37.
    tmp

GN20GG_masked_autoXY_EMT_genepromoter_comprehensive_noPAM.fa >
GN20GG_masked_autoXY_EMT_genepromoter_comprehensive_noPAM.sai;

bwa samse -n 10000 ~/path_to/GRCh37/human_GRCh37.tmp

GN20GG_masked_autoXY_EMT_genepromoter_comprehensive_noPAM.sai
GN20GG_masked_autoXY_EMT_genepromoter_comprehensive_noPAM.fa >
GN20GG_masked_autoXY_EMT_genepromoter_comprehensive_noPAM.sam;

samtools view -S -q1 GN20GG_masked_autoXY_EMT_genepromoter_
comprehensive_complete_noPAM.sam | cut -f 1 | awk -F ':' '{
    print $1 "\t" $2}' | awk -F '-' '{print $1 "\t" $2 "\t" $3
    }' | sortBed | uniq > GN20GG_masked_autoXY_EMT_
genepromoter_comprehensive_complete_noPAM_unique.bed
```

This file contains 5086 sequences, i.e. 5086 gRNA sequences of the form GN19 that align uniquely to the genome and are followed by an NGG PAM and are found in the regions of interest.

3.4 Addition of plasmid sequences for use in Gibson cloning

For cloning into the gRNA vector pgRNA-pLKO.1 (see Methods) by Gibson cloning, vector-derived sequences were attached to either end of the gRNA sequence. To this end the FASTA files for the final set of 5086 unique gRNAs were retrieved and sequences appended as follow:

```
#split according to + and - strand
```

```
grep '+'
   GN20GG_masked_autoXY_EMT_genepromoter_comprehensive_complete_
    noPAM_unique_strand.bed \mid awk '{print $1 ":" $2 "-" $3}' >
   GN20GG_masked_autoXY_EMT_genepromoter_comprehensive_complete_
    noPAM_unique_strand_PLUS.2bit
grep -v '+'
   GN20GG_masked_autoXY_EMT_genepromoter_comprehensive_complete_
    noPAM_unique_strand.bed | awk '{print $1 ":" $2 "-" $3}' >
   GN20GG_masked_autoXY_EMT_genepromoter_comprehensive_complete_
    noPAM_unique_strand_MINUS.2bit
#convert to FASTA format and reverse-complement sequences on
   the
#minus strand
twoBitToFa /path2/human/GRCh37/hg19.2bit
   GN20GG_masked_autoXY_EMT_genepromoter_comprehensive_complete_
    noPAM_unique_strand_PLUS.fa seqList=
   GN20GG_masked_autoXY_EMT_genepromoter_comprehensive_complete_
    noPAM_unique_strand_PLUS.2bit;
twoBitToFa /path2/human/GRCh37/hg19.2bit
   GN20GG_masked_autoXY_EMT_genepromoter_comprehensive_complete_
    noPAM_unique_strand_MINUS.fa seqList=
   GN20GG_masked_autoXY_EMT_genepromoter_comprehensive_complete_
    noPAM_unique_strand_MINUS.2bit
python reverse_complement_fasta.py
   GN20GG_masked_autoXY_EMT_genepromoter_comprehensive_complete_
    noPAM_unique_strand_MINUS.fa >
```

```
GN20GG_masked_autoXY_EMT_genepromoter_comprehensive_complete_
   noPAM_unique_strand_MINUS_RC.fa
#merge the two files
cat
  GN20GG_masked_autoXY_EMT_genepromoter_comprehensive_complete_
noPAM_unique_strand_PLUS.fa
  GN20GG_masked_autoXY_EMT_genepromoter_comprehensive_complete_
noPAM_unique_strand_MINUS_RC.fa >
  GN20GG_masked_autoXY_EMT_genepromoter_comprehensive_complete_
noPAM_unique_strand.fa
# use trimming.py to remove PAM (because coordinates extracted
# from SAM file contained PAM, while alignment was run without
# PAM) and paste the vector sequences for Gibson cloning on
# either side of the gRNA sequence
sed 'n; s/$/GTTTTAGAGCTAGAAATAGCAAGTTAAAATAAGGCT/'
  GN20GG_masked_autoXY_EMT_genepromoter_
  comprehensive_complete_noPAM_unique.fa | sed 'n; s/^/
  TCTTGTGGAAAGGACGAAACACC/g' | paste - - | awk '{print $2 "\t
  " $1}' > EMT_guides_Custom_Array.txt
```

A pool of sequences of the form 5'-TCTTGTGGAAAGGACGAAACACC-GN19-GTTTTAGAGCT AGAAATAGCAAGTTAAAATAAGGCT-3', where GN19 donates the 5086 different guide sequences was then ordered from Custom Array Inc.

Chapter 4

Data analysis: Screens with the EMT5000 library and stable cell lines expressing the dCas9-SET7 or dCas9-p300 chromatin modifier

4.1 Read trimming and Quality Filtering

gRNA sequences integrated into the genome of FACS-sorted cells were amplified using PCR to add Illumina Nextera adapters. Libraries were sequenced on the Illumina HiSeq instrument. The resulting sequencing reads have the following general structure:

NNNNNNAAGTATTTCGATTTCTTGGCTTTATATATCTTGTGGAAAGGACGAAACACCG-N19-GTTTT
AGAGCTAGAAATAGCAAGTTAAAATAAGGCTAGTCCGNNNNNNN

The first 7 N bases are the 5' barcode, followed by the plasmid 'stuffer'. GN19 denotes the gRNA sequence from the EMT5000 library, which is followed by another plasmid sequence and the last 7 N bases are the 3' barcode. The 5' and 3' barcodes serve as unique molecular identifiers (UMIs), allowing counting of original gRNA sequences extracted from lentivirus-infected cells by removing PCR-amplification bias.

Sequences from Lane1 and Lane2 of the flow cell were combined using the unix 'cat' command. Next, the 5' barcode was extracted from the reads using cutadapt (version 1.2.1) [13], requiring a minimum overlap of 35 bp between the plasmid stuffer sequence and the read with a maximum error of 10% and a minimum barcode length of 7 bp.

```
Command line parameters:

-a AAGTATTTCGATTTCTTGGCTTTATATATCTTGTGGAAAGGACGAAACACC -0 35 -
    m 7

# example bash loop to run cutadapt over all fastq files
# in the directory
for i in *.fastq.gz
do cutadapt -a
    AAGTATTTCGATTTCTTGGCTTTATATATCTTGTGGAAAGGACGAAACACC -0 35 -
    m 7 --untrimmed-output="${i%.fastq.gz}"_5bc_untrimmed.fastq
    .gz -o "${i%.fastq.gz}"_5bc.fastq.gz "$i";
done
```

The 3' barcode was retrieved from the read in an analogous way:

```
Command line parameters:
-g GTTTTAGAGCTAGAAATAGCAAGTTAAAATAAGGCTAGTCCG -O 28 -m 7
# example bash loop to run cutadapt over all fastq files
# in the directory
for i in *.fastq.gz
do cutadapt -g GTTTTAGAGCTAGAAATAGCAAGTTAAAATAAGGCTAGTCCG -O
   28 -m 7 --untrimmed-output="${i%.fastq.gz}"_3bc_untrimmed.
   fastq.gz -o "${i%.fastq.gz}"_3bc.fastq.gz "$i";
done
Finally, the gRNA sequence was extracted from the read, requiring a minimum length
of 2 bp (to discard reads that contain no gRNAs and are derived from primer dimers):
Command line parameters:
cutadapt -g
   AAGTATTTCGATTTCTTGGCTTTATATATCTTGTGGAAAGGACGAAACACC -a
   GTTTTAGAGCTAGAAATAGCAAGTTAAAATAAGGCTAGTCCG -n 2 -m 2 -0 10
# example bash loop to run cutadapt over all fastq files
# in the directory
for i in *.fastq.gz
do cutadapt -g
   AAGTATTTCGATTTCTTGGCTTTATATATCTTGTGGAAAGGACGAAACACC -a
   GTTTTAGAGCTAGAAATAGCAAGTTAAAATAAGGCTAGTCCG -n 2 -m 2 -0 10
   --untrimmed-output="${i%.fastq.gz}"_gRNA_untrimmed.fastq.gz
    -o "${i%.fastq.gz}"_gRNA.fastq.gz "$i";
done
```

Next, the barcode and gRNA reads were quality-filtered using fastq_quality_filter from the fastx-toolbox [1]. Reads where any base has a Quality score of less than 20 were discarded (Example code below is for the gRNA reads.)

```
Command line parameters: -Q33 -q 20 -p 1

#loop over all files in directory:

for i in *_gRNA.fastq; do fastq_quality_filter -Q33 -q 20 -p 1

-i "$i" -o "${i%_gRNA.fastq}"_gRNA_Q20.fastq; done
```

Subsequently files were converted from fastq to fasta format using fastq_to_fasta from the fastx-toolbox [1].

```
for i in *_gRNA_Q20.fastq;
do fastq_to_fasta -Q33 -n -i "$i" -o "${i%gRNA_Q20.fastq}"
    gRNA_Q20.fasta;
done
```

4.2 Alignment to the EMT5000 library

Guide RNA reads were next aligned back onto the indexed EMT5000 reference library using bwa (version: 0.6.2-r126) [9].

The indexed EMT5000 library file used as a reference for alignment and was derived from the file GN20GG_masked_autoXY_EMT_genepromoter_comprehensive_complete_noPAM_unique_s (see section 3.1) using bwa index:

```
bwa index GN20GG_masked_autoXY_EMT_genepromoter_comprehensive_
    complete_noPAM_unique_strand_PAMremoved.fa
```

I empirically tested the following alignment parameters:

```
#Command line options 3 mismatches, no indels:
bwa aln -n 3 -o 0 -l 5 -N -I

#Command line options 2 mismatches and default open gaps (1):
bwa aln -n 2 -l 5 -N -I

#Command line options 3 mismatches and default open gaps (1):
do bwa aln -n 3 -l 5 -N -I
```

I found that allowing 2 mismatches and 1 gap gave the best alignment (see also the table of read numbers included in the Appendix of the PhD thesis). The alignment was invoked as follows:

```
for i in ../*_Q20.fasta;
do bwa aln -n 2 -l 5 -N -I EMT5000_library_reference/
    GN20GG_masked_autoXY_EMT_genepromoter_comprehensive_
complete _noPAM_unique_strand_PAMremoved.fa "$i" > "${i%Q20.
    fasta}"Q20_aligned_2mismatches1gap.sai;
done

for i in *Q20_aligned_2mismatches1gap.sai;
do bwa samse -n 10000 EMT5000_library_reference/
    GN20GG_masked_autoXY_EMT_genepromoter_comprehensive_
complete_noPAM_unique_strand_PAMremoved.fa "$i" "${i%
    Q20_aligned_2mismatches1gap.sai}"Q20.fasta > "${i%
    Q20_aligned_2mismatches1gap.sai}"
    Q20_aligned_2mismatches1gap.sai;
done
```

Following alignment allowing 2 mismatches and 1 gap, reads that mapped uniquely to the forward strand were extracted using Samtools [10]:

```
samtools view -F 20 -q 1 -S aligned_gRNA.sam >
gRNA_uniquely_mapped.sam
```

To check how many different gRNAs from the library were sequenced in each sample, use:

```
samtools view -F 20 -S aligned_gRNA.sam | cut -f 3 | sort |
uniq | wc -l
```

4.3 Combining the gRNA and adapter sequences in a single file

For subsequent analysis it was necessary to construct a tab-separated file with 3 columns containing the FASTA identifier (read-ID), gRNA chr:start-end and barcode sequence for each read.

```
# extract the gRNA:
samtools view -F 20 -q 1 -S aligned_gRNA.sam | awk '{print$1
   ".\t" $3}' > gRNA_uniquelymapped

# extract the FASTA identifier:
samtools view -F 20 -q 1 -S aligned_gRNA.sam | awk '{print$1
   "."}' > gRNA_uniquelymapped_readID
```

The files containing the quality-filtered 5' and 3' barcode (UMI sequence) generated above, were modified as follows (shown for 5' barcode only):

Next, only the barcodes associated with gRNAs that aligned uniquely were retrieved using grep:

```
for i in *_5bc_Q20_point.fasta
```

```
do grep -wFf "${i%_5bc_Q20_point.fasta}"
    gRNA_uniquelymapped_readID "$i" > "${i%_5bc_Q20_point.fasta}"
    gRNA_uniquelymapped_readID_with_5bc;
done
```

For each read, identified by its readID, the gRNA sequence and 5' and 3' barcodes were combined into a single file. The three files were joined (finding the union) using the JOIN command, which requires the files to be sorted in the following way:

```
for i in *gRNA_uniquelymapped;
do awk -F "\t" '{print ">" 1 \t" 2' "i \t" sort -k 1b, 1 > "
  ${i%}"_sorted;
done
for i in *gRNA_uniquelymapped_readID_with_3bc;
do sort -k 1b,1 "i" > "i" > "i" sorted;
done
for i in *gRNA_uniquelymapped_readID_with_5bc; do sort -k 1b,1
    "$i" > "${i%}"_sorted;
done
Next, the three files were joined as follows:
for i in *gRNA_uniquelymapped_sorted;
do join "$i" "${i%gRNA_uniquelymapped_sorted}"
   gRNA_uniquelymapped_readID_with_5bc_sorted | join - "${i%
   gRNA_uniquelymapped_sorted}"
   gRNA_uniquelymapped_readID_with_3bc_sorted > "${i%
   gRNA_uniquelymapped_sorted}"
   gRNA_uniquelymapped_readID_gRNA_5bc_3bc_length14;
done
```

This yields a file containing for each uniquely mapped read its readID, the gRNA it mapped to (chr:start-stop) and the UMI found in the read (of length exactly 14 bp).

4.4 Deriving gRNA counts from UMI-barcodes without PCR error correction

The gRNA counts were derived by counting the number of times each gRNA occurs together with each barcode, which acts as a unique molecular identifier (UMI). To do the gRNA counting without any error correction, the script collapse_barcodes.py was run using a maximum edit distance of 0, i.e. not correcting any PCR errors that might have occurred in the barcode. This script can be invoked as follows:

```
python collapse_barcodes.py Inputfilename 0
```

The script accepts a tab-separated file with columns for (1) read-ID, (2) gRNA (chr:start-stop), (3) barcode and outputs two outputfiles with extension _frequency_raw and frequency_no_orphans. In the latter output orphan barcodes, i.e. barcodes that are only present in a single read, were removed prior to gRNA counting. Each output file has two comma-separated columns listing (1) the gRNA (chr:start-stop) and (2) the gRNA count.

The script was run over all files with the extension

```
_uniquelymapped_readID_gRNA_5bc_3bc_length14 as follows:
for i in *length_14;
do python collapse_barcodes.py "$i" 0;
done
```

4.4.1 Assessing PCR error by plotting the number of reads per gRNA against number of different gRNA sequences

To assess whether PCR error drives barcode diversity, I treated the gRNA part of the read like a barcode and plotted the correlation between number of reads and counts (see Figure 4.8. on page 134 of the PhD thesis and also section 4.7 below). This

assumes that the likelihood of introducing an error into the sequence is the same for the UMI barcodes and gRNA portions of the amplicon.

The barcode sequence was replaced with the gRNA sequence for each read to generate a tab-separated file with columns (1) readID (2) gRNA chr:start-stop (3) 'pseudo-barcode'(gRNA sequence) as follows:

```
#get gRNA sequence
cat Sample_gRNA_Q20.fasta | paste - - | awk -F ' ' '{print $1
   ".\t" $3}' | sort > Sample_gRNA_Q20_point
#get the read ID
awk -F '\t' '{print $1}' Sample_gRNA_5bc_3bc_length14 | sort >
    Sample_gRNA_5bc_3bc_length14_read_ID
#get gRNA sequence for each readID
grep -wFf Sample_gRNA_5bc_3bc_length14_read_ID
   Sample_gRNA_Q20_point >
   Sample_gRNA_Q20_point_uniquely_mapped
#sort the previously generate file containing [0] readID, [1]
   gRNA chr:start-stop, [2] UMI of 5' and 3' barcode
sort -k 1b,1
   Sample_gRNA_uniquelymapped_readID_gRNA_5bc_3bc_length14 >
   Sample_gRNA_uniquelymapped_
readID_gRNA_5bc_3bc_length14_sorted
#sort the gRNA sequence file
sort -k 1b,1 Sample_gRNA_Q20_point_uniquely_mapped >
   Sample_gRNA_Q20_point_uniquely_mapped_sorted
#join the two files on readID
join Sample_gRNA_Q20_aligned_2mismatches1gap_uniquelymapped_
readID_gRNA_5bc_3bc_length14_sorted
```

```
Sample_gRNA_Q20_point_uniquely_mapped_sorted | awk -F ' '
'{print $1 "\t" $2 "\t" $4}' >
Sample_gRNA_Q20_aligned_2mismatches1gap_uniquelymapped_
readID_gRNA_instead_of_barcode
```

This file was then run through collapse_barcodes.py as above and results were plotted as described in section 4.7.

```
python collapse_barcodes.py
    Sample_gRNA_Q20_aligned_2mismatches1gap_
uniquelymapped_readID_gRNA_instead_of_barcode 0
```

4.5 Deriving gRNA counts from UMI-barcodes with naive PCR error correction

The script collapse_barcodes.py was run using a maximum edit distance of 4. python collapse_barcodes.py Samplefilename 4

This means that before counting how many different barcodes are associated with each gRNA, the barcodes are collapsed into groups. Barcodes are first ranked in decreasing order based on the number of reads harbouring its sequence. The barcode with the most reads forms the first group. If the second-ranked barcode is within 4 edit-distances of this barcode it will be assumed to have originated by PCR error and will be added to the group. If the barcode differs from the group by greater than 4 edits, it will form its own group and so on. The number of groups per gRNA is the count after error correction. This reflects the number of original gRNA-barcode combinations.

The script was run over all files with the extension

_uniquelymapped_readID_gRNA_5bc_3bc_length14 as follows:

```
for i in *length_14;
do python collapse_barcodes.py "$i" 4;
done
```

4.6 Bayesian PCR error correction of barcoded sequencing data

A Bayesian error correction script was written by James E. Barrett to infer gRNA counts from the UMI data. The model takes into account the fact that the 14 bp UMI consists of a 5' and 3' barcode that was attached to the gRNA amplicon during and initial round off PCR amplification during the sequencing library prep. The model infers the most likely number of initial gRNA-barcode data given the barcode sequences observed in the sequencing sample.

This Bayesian model takes as input the number of reads associated with each gRNA-UMI combination (without PCR error correction). I calculated these using the script make-csv-4Bayes.py. The script was run over all samples as follows:

```
for i in *length14; do python ../make_csv_4Bayes.py "$i"; done
```

This script again takes the tab-separated three column file that lists (1) read ID, (2) gRNA (chr:start-end) and (3) barcode consisting of 5' UMI and 3' UMI fused together as input. The output is a csv file with three columns containing (1) gRNA (chr:start-end), (2) barcode consisting of 5' UMI and 3' UMI fused together and (3) number of reads associated with each barcode. The outputfile has the extension _barcode_readcounts.csv and is fed into a Bayesian error correction script described below.

4.6.1 Bayesian PCR error correction of barcoded sequencing count data script (by James E. Barrett)

This script and documentation was kindly contributed by James E. Barrett.

The Bayesian model infers the number of unique original barcoded gRNA molecules from noise-corrupted count data. The model estimates a corrected read count, which may be interpreted as a proxy for the original noise-free number of unique barcodes associated with a particular gRNA.

Model definition

For each gRNA we observe N barcode pairs denoted by $(\mathbf{y}_i^1, \mathbf{y}_i^2)$ where the superscript denotes the first and second barcodes and i = 1, ..., N. Elements of the d-dimensional vector $\mathbf{y}_i^{\eta} \in \{\mathtt{T}, \mathtt{C}, \mathtt{G}, \mathtt{A}\}^d$ where $\eta = [1, 2]$. The number of corresponding sequencing reads is denoted by $\sigma_i \in \mathbb{Z}_+$.

The model assumes that there exist Q latent barcodes $\mathbf{x}_1^{\eta}, \dots, \mathbf{x}_Q^{\eta}$ from which the observed barcodes are generated in a noise corrupting stochastic process (PCR amplification errors and random barcode switching). The model further assumes that for each pair $(\mathbf{y}_i^1, \mathbf{y}_i^2)$ only one of the observed barcodes is written in terms of the latent barcode via

$$\mathbf{y}_{i}^{\eta} = \sum_{q=1}^{Q} w_{iq}^{\eta} \theta(\mathbf{x}_{q}^{\eta}) \quad \text{subject to} \quad w_{iq}^{\eta} \in [0, 1] \quad \text{and} \quad \sum_{q, \eta} w_{iq}^{\eta} = 1. \tag{4.1}$$

There is therefore only one non-zero value of $[\mathbf{w}_i^1, \mathbf{w}_i^2]$ that indicates which latent barcode the observed pair is associated with. The function θ represents a noise corrupting stochastic process where the status of each nucleotide site may be changed randomly with probability $\beta \in [0, 1/2]$. We can therefore write

$$p(y_{i\mu}^{\eta}|x_{q\mu}^{\eta},\beta) = \begin{cases} (1-\beta)\delta_{y_{i\mu}^{\eta}x_{q\mu}^{\eta}} + \beta(1-\delta_{y_{i\mu}^{\eta}x_{q\mu}^{\eta}}) & \text{if } w_{iq}^{\eta} = 1\\ 0 & \text{otherwise} \end{cases}$$
(4.2)

for $\mu = 1, ..., d$. We denote the collections of \mathbf{x}_q^{η} , \mathbf{y}_i^{η} and \mathbf{w}_i^{η} by \mathbf{X} , \mathbf{Y} , and \mathbf{W} respectively. The posterior is

$$p(\mathbf{X}, \mathbf{W}|\mathbf{Y}, \boldsymbol{\sigma}, \beta) \propto p(\mathbf{Y}|\mathbf{X}, \mathbf{W}, \boldsymbol{\sigma}, \beta)p(\mathbf{X})p(\mathbf{W})$$
 (4.3)

with

$$p(\mathbf{Y}|\mathbf{X}, \mathbf{W}, \boldsymbol{\sigma}, \beta) = \prod_{i} \left[\sum_{q,\eta} w_{iq}^{\eta} p(\mathbf{y}_{i}^{\eta} | \mathbf{x}_{q}^{\eta}, \beta) \right]^{\sigma_{i}}.$$
 (4.4)

Maximum entropy priors for \mathbf{X} and \mathbf{W} are uniform distributions so $p(\mathbf{X})$ and $p(\mathbf{W})$ are constant.

Inference of model parameters

The Maximum A Posteriori (MAP) solution of **W** is denoted by **W***. Since only one element of $[\mathbf{w}_i^1, \mathbf{w}_i^2]$ is non-zero the expression (4.4) is maximised by selecting $\arg\max_{q,\eta} p(\mathbf{y}_i^{\eta} | \mathbf{x}_q^{\eta}, \beta)$ as the non-zero element.

To find the MAP solution for nucleotide μ of the latent barcode indexed by (q, η) we consider all observed barcodes that generated from it (as defined by **W**). If we let n_1 and n_0 denote the total number of matches and mismatches respectively between that latent barcode and the associated observed barcodes, then the corresponding data likelihood is $(1-\beta)^{n_{q\mu}^1}\beta^{n_{q\mu}^0}$. This will be maximised if the number of matches is maximised. This is achieved selecting the most common observed nucleotide as the value for the latent nucleotide (while taking into account multiple counts).

If we let N_1 and N_0 denote the total number of matches and mismatches respectively across all of the latent barcodes and observed data then we can write

$$\log p(\mathbf{Y}|\mathbf{X}, \mathbf{W}, \beta) = N_1 \log(1 - \beta) + N_0 \log \beta. \tag{4.5}$$

It is straightforward to show that the MAP estimate for beta is

$$\beta = \frac{N_0}{N_0 + N_1}. (4.6)$$

The optimisation subroutine is initialised as follows:

Cluster into Q groups based on the *Hamming distance* between two barcodes (the Hamming distance is equivalent to the *edit distance*):

$$h(\mathbf{y}_i, \mathbf{y}_j) = \frac{1}{d} \sum_{\mu=1}^{d} \delta_{(1-y_{i\mu})y_{j\mu}}.$$
 (4.7)

The corrected read counts are inferred as follows:

For a given value of Q we denote the value of the likelihood (4.4) at the MAP parameter estimate by

$$L(Q) = p(\mathbf{Y}|\mathbf{X}^*, \mathbf{W}^*, \beta^*). \tag{4.8}$$

The Bayes information criterion (BIC) score is defined by

$$BIC(Q) = -2\log L(Q) + 2dQ\log N \tag{4.9}$$

where 2dQ is the number of free parameters in the model. The *corrected read count* is defined by

$$Q^* = \operatorname{argmin}_Q \mathrm{BIC}(Q). \tag{4.10}$$

Bayesian error correction script: The Code

The analysis is performed in R:

library(reshape2)

Load and prepare a data file

```
# Length of barcode
D <- 7
# Load up one of the data files (needs to be in the current
   directory)
data <- read.csv("Samplename_length14_barcode_readcounts.csv",</pre>
   header=FALSE)
# vector of all the unique gRNA names
gRNA <- unique(data$V1)
# Total number of unique gRNAs
G <- length(gRNA)
### Generate datasets of barcodes
# Preallocate a list structure to hold the barcode datasets
Y <- vector('list',G)
# This loop goes through each gRNA, pulls out all the
   associated barcodes and puts them in a character matrix
for(mu in 1:G){
   ind <- which(data[[1]] == gRNA[mu])</pre>
   N <- length(ind)
   # Converts into character matrix (not the most elegant way
   Y[[mu]] <- matrix(as.vector(melt(lapply(as.character(data
      [[2]][ind]),strsplit,split=""))$value),nrow=N,ncol=2*D,
      byrow=TRUE)
}
### Fit model for each gRNA
# Preallocate a list of model results
res <- vector('list',G)</pre>
```

This analysis calls functions stored in the R scripts fit_model.R, LL.R and hamming.R. A csv file of counts per gRNA for each sample (bayesian_corrected.csv) was then exported.

4.7 Diagnostic plot: Number of UMI-corrected counts versus number of reads per gRNA

4.7.1 Calculating the number of reads per gRNA

To calculate the number of reads per gRNA for each sample, I wrote the script reads_per_gRNA.py. This takes a 3 column tab-separated inputfile with the following columns: (1) read ID (2) gRNA chr:start-stop (3) 14 nt barcode and outputs a csv file with two columns: (1) gRNA chr:start-stop, (2) number of reads. The script can be invoked as follows:

4.7.2 Wrapping gRNA counts of all samples into a table

While the Bayesian model outputs a csv file containing the counts per gRNA for each sample directly, the output of the script collapse-barcodes.py (used to count gRNAs without error correction or to perform a naive PCR error correction) outputs one table per sample. This data can be merged into a single table listing for each gRNA the count in each sample using the script make_table_from_counts.py. This script also adds information about which gene is targeted by each gRNA. The script takes a variable number of inputfiles to wrap into a table:

```
python makeTable_from_counts.py INPUTFILE1 INPUTFILE2 ...
```

This was used to generate the tables Dataframe_allsamples_readcounts.txt

4.7.3 Generating the plots of counts versus number of reads for each gRNA

The number of reads per gRNA were plotted against the counts per gRNA, derived either without error correction, with naive PCR error correction or Bayesian PCR error correction (as described above), using the script plot_counts_vs_number_of_reads.py. Samplesheets can be found in the folder "samplefiles".

```
python plot_counts_vs_number_of_reads.py [Dataframe-Counts] [
    Dataframe-NumberOfReads] [Samplesheet]

# no PCR error correction
plot_counts_vs_number_of_reads.py
    Dataframe_allsamples_no_errors.txt
    Dataframe_allsamples_readcounts.txt samplefile.txt
```

```
# naive PCR error correction (4 mismatches)
plot_counts_vs_number_of_reads.py Dataframe_allsamples.txt
    Dataframe_allsamples_readcounts.txt samplefile.txt

# Bayesian error correction
plot_counts_vs_number_of_reads.py bayesian_corrected.csv
    Dataframe_allsamples_readcounts.txt samplefile.txt
```

This script was used to generate the plots in Figure 4.9 and Figure 4.10 of the PhD thesis.

4.8 Diagnostic plot: Counts versus number of sorted cells

This script accepts three user-supplied arguments, a dataframe of gRNA counts (c), a dataframe of numbers of sorted cells per sample (n), and a samplesheet (s) and returns a scatterplot of sorted cells versus counts with one data point per sample in the samplesheet. It is further hardcoded to color the dots according to whether two or three initial cycles of barcoding PCR were used to attach unique molecular identifiers to gRNA sequences before amplification and sequencing. The samplesheets can be found in the folder samplefiles and the file recording the number of sorted cells is in the folder additional-files. The script was invoked as follows:

```
python cellnumber_vs_counts.py -c bayesian_corrected.csv -n
Number_of_sorted_cells.csv -s samplefile_all.txt
```

This script was used to produce the graph in Figure 4.11 on page 134 of the PhD thesis.

4.9 Enrichment analysis using DESeq2

Enrichment analysis was carried out using the DESeq2 package [11].

4.9.1 Preparing Bayesian error correction output for DE-Seq2

DESeq2 requires a list of counts per gRNA for each experiment to be analysed. The data for each experiment were extracted from the Bayesian analysis output file bayesian_corrected.csv. gRNAs with a lot of missing data where 3/4 of the counts in a given experiment are 0 (or NA) are removed before enrichment analysis using the script make_input_4_DESeq.py The samplesheets can be found in the folder samplefiles.

```
python make_input_4_DESeq.py -i bayesian_corrected.csv -s
    Samplefile_Batch8.txt
```

Using the above example, the script outputs a file named Batch8_p300_counts.csv.

4.9.2 Running DESeq2

This analysis was conducted using the script DESeq2_script.R. An example of how this script is run over the example file for experiment Batch8_p300 is shown below. DESeq2 further requires a file containing experimental information, specifying for each sample whether it belongs to the treatment or control group. These files can be found in the folder additional_files/ColData_forDESeq2.

```
./DESeq2_script.R -c path_to/DeSeq2/Inputfiles/
Batch8_p300_counts.csv -e path_to/additional_files/
ColData_forDESeq2/Batch8_DeSeq2_ColData -o '
Batch8_p300_DESeq2_table.csv'
```

This outputs a csv file with the samplename and the extension _DESeq2_table.csv.

4.9.3 Visualizing enrichment

Log2 Fold Change between samples and controls for each gRNA was calculated using DESeq2 as described above. Log2Fold Change values were extracted from the output of DESeq2 (filename is Batch8_p300_DESeq2_table.csv for this particular example) and plotted along the chromosome for each gene in the library. To extract the relevant data and generate the plots I wrote the python script Plot_Log2FC_along_chr.py. This script requires as input the additional files EMT5000_library_regions.bed, and EMT5000_library_TSS.txt in order to plot enrichment for each gRNA with respect to the transcriptional start site of the gene from the library. Both files can can be found in the folder additional_files. The script can be invoked as follows:

```
python Plot_Log2FC_along_chr.py -c Batch8_p300_DESeq2_table.
    csv -l EMT5000_library_regions.bed -t EMT5000_library_TSS.
    txt
```

This script was used to generate the plots in Figure 4.12 on page 136 of the PhD thesis.

4.9.4 Correlation of Log2Fold enrichment scores from DE-Seq2 between experiments

The script plot_LogFC_vs_LogFC.py performs all-by-all comparison of Log2Fold Change per gRNA across an arbitrary number of inputfiles. The script can be invoked as follows:

```
python plot_LogFC_vs_LogFC.py [list of DESeq output
  dataframes to be compared]
```

This script was used to generate the plots in Figure 4.13 on page 138 of the PhD thesis.

4.9.5 Identification of candidate gRNAs using ranking with desirability functions

The Log2Fold Change and Log2Fold Change standard error were extracted from the DESeq2 output in python (using ipython notebook interactively) as follows:

```
import numpy as np
import pandas as pd
import matplotlib.pyplot as plt
import collections as coll
import re
import pylab
import pybedtools
%gui
%matplotlib inline
def load_into_df(arg):
    user_input = pd.DataFrame.from_csv(arg, header=0, sep=',',
        index_col=0)
    return user_input
def set_DESEq2_outliers_to_0_and_wrap_into_df(list_of_dfs,
   names_of_dfs):
    dict_for_df = {}
    for index, df in enumerate(list_of_dfs): #iterate over the
        dataframes
        outliers = df.loc[np.isnan(df['pvalue'])] #make a
           dataframe 'outliers' that holds all rows where
           pvalue is NA, i.e DESeq2 has detected an outlier
```

```
outliers['log2FoldChange']=0 #if pvalue is NA set L2FC
        outliers_removed = df.loc[~np.isnan(df['pvalue'])] #
           make new df only containing rows where pvalue is
           NOT NaN
        df_combined = pd.concat([outliers, outliers_removed])
             #combine the two dataframes
        dict_for_df[names_of_dfs[index] + '_log2FoldChange'] =
            (df_combined["log2FoldChange"])
        dict_for_df[names_of_dfs[index] + '_LfcSE'] = (
           df_combined["lfcSE"]) #add the columns holding L2FC
            of this dataframe to a dict
    my_df=pd.DataFrame(dict_for_df) #take entries from a list
       into a df
    return my_df
Batch8_p300 = load_into_df('path2/DESeq2_output_tables/
   Batch8_p300_DESeq2_table.csv')
Batch4_p300 = load_into_df('path2//DESeq2_output_tables/
   Batch4_p300_DESeq2_table.csv')
BatchSS0209_p300 = load_into_df('path2/DESeq2_output_tables/
   BatchSS0209_p300_DESeq2_table.csv')
BatchSS2608_p300 = load_into_df('path2/DESeq2_output_tables/
   BatchSS2608_p300_DESeq2_table.csv')
BatchSS0209_Set7 = load_into_df('path2/DESeq2_output_tables/
   BatchSS0209_Set7_DESeq2_table.csv')
BatchSS2608_Set7 = load_into_df('path2/DESeq2_output_tables/
   BatchSS2608_Set7_DESeq2_table.csv')
Batch5_Set7 = load_into_df('path2/DESeq2_output_tables/
   Batch5_Set7_DESeq2_table.csv')
Batch3_Set7 = load_into_df('path2/DESeq2_output_tables/
   Batch3_Set7_DESeq2_table.csv')
```

```
list_p300 = [Batch8_p300, Batch4_p300, BatchSS0209_p300,
   BatchSS2608_p300]
names_p300 = ['Batch8_p300', 'Batch4_p300', 'BatchSS0209_p300
   ', 'BatchSS2608_p300']
list_Set7 = [Batch3_Set7, Batch5_Set7, BatchSS0209_Set7,
   BatchSS2608_Set7]
names_Set7 = ['Batch3_Set7', 'Batch5_Set7', 'BatchSS0209_Set7
   ', 'BatchSS2608_Set7']
p300s_L2FC = set_DESEq2_outliers_to_0_and_wrap_into_df(
   list_p300, names_p300)
p300s_L2FC.to_csv('p300s_DESeq_L2FC_NApvalue2zero.csv', sep
  = ', ')
Set7_L2FC = set_DESEq2_outliers_to_0_and_wrap_into_df(
   list_Set7, names_Set7)
Set7_L2FC.to_csv('Set7s_DESeq_L2FC_NApvalue2zero.csv', sep
  = ', ')
```

For each gRNA where DESeq2 had detected outliers (and set the p-value to NA in the output) Log2Fold Change values were set to 0 before the next step, i.e. ranking of gRNAs.

The script Desirability.R was used to rank gRNAs based on large positive Log2FC and small Log2FC standard error. A weighted average is calculated and Log2FC is given four times the weight of its standard error during this ranking:

```
./Desirability.R -f p300s_DESeq_L2FC_NApvalue2zero.csv -w 4 -e 1
```

The script outputs a ranked list of candidate gRNAs, ranked based on the calculated overall desirability (here p300_candidates_4xL2FC_1xLfcSE.csv as well as plots showing Desirabilities across all gRNAs as well as histograms for Log2Fold Changes

and associated standard errors. These are shown in Figure 4.14 on page 139 of the PhD thesis.

The top 10 candidates were extracted from the list of gRNAs ranked by their Desirability score for use in the validation experiments. The sequences were extracted for cloning into the gRNA vector as follows:

```
head -11 p300_candidates.csv | sed 1d | awk -F '",' '{print $1
}' | awk -F '"' '{print $2}' > p300_top10.bed

grep -A1 -wf p300_top10.bed /path2/
   GN20GG_masked_autoXY_EMT_genepromoter_comprehensive_complete_
```

```
noPAM_unique_strand_PAMremoved.fa > p300_top10.fa
```

For the candidate gRNAs the Bayesian-corrected counts in samples and controls were plotted using the custom script Plot_candidates.R. For this, the counts for each gRNA for all samples from a given experiment were first extracted from the output of the Bayesian error correction script (see section 4.6.1) using the script extract_all_counts_per_experiment_from_bayes.py, which was invoked as follows:

```
python extract_all_counts_per_experiment_from_bayes.py
```

This produces, for each experiment a file with extension _all-counts.csv. This file was edited to replace all occurrences of NA with 0 using the script

Print_counts_missing_as_0.R. The file containing the output of the Bayesian error correction script for all experiments using the dCas9-p300 chromatin modifier is shown as an example:

```
./Print_counts_missing_as_0.R -f All_p300_all-counts.csv
```

For this particular example, this script outputs a file named

All_p300_all-counts_missing_as_0.csv, which contains for all experiments using the dCas9-p300 chromatin modifier the counts per gRNA following Bayesian

error correction for all samples and controls with NAs replaced by 0. This file together with the ranked list of candidates (from section 4.9.5) was fed to the script Print_counts_missing_as_0.R as follows:

This produced the plots in Figure 4.15 and 4.16 on page 140 and 141 of the PhD thesis.

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