final\_paper

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Mapping of global transcriptional regulator MftR binding loci in *Burkholderia thailandensis* using ChIP-seq

Introduction

Bacterial diseases infect individuals throughout the history of mankind with fatal consequences. Finding an efficient treatment against infectious bacterial diseases is a challenging endeavor. Ancient civilizations used antimicrobial agents to overcome the spread of such maladies [1]. It was reported that in ancient Egypt and ancient Greek, certain types of funguses were used to cure infections [2]. In the late 1880, a Nobel prize laureate Paul Ehrlich developed the first dye based synthetic antibiotic which was called later arsphenamine [3]. In 1928, Sir Alexander Fleming discovered the first effective antibiotic which he called the fungal extract which is known now as penicillin [4].

Since the discovery of penicillin, more than 100 antibiotics were discovered, each has a certain mechanism of action [5]. Different antibiotics target certain vital processes that interfere with the bacterial cellular functions. Antibiotic molecular targets include interfering with protein synthesis, cell wall construction, DNA synthesis and folic acid synthesis [6]. However, bacteria develop defensive mechanisms to resist antibiotics and survive. The misuse of antibiotics worldwide accelerated the evolution of bacterial resistance which has been clearly observed in the last couple of decades. In 2015, it was reported that more than 33 thousand persons died from antibiotic resistance bacteria [7].

One defense mechanism that bacteria use to fight antibiotics and other toxins is efflux pumps. Efflux pumps are transmembrane proteins that are localized in the cytoplasmic membranes in all bacterial cells. Some efflux pumps are specific to one antibiotic while others are multidrug efflux pumps [8]. There are five main families of efflux pumps which are ATP binding cassette family, small multidrug resistance family, multidrug and toxin extrusion family, major facilitator superfamily and resistance nodulation cell division family [9]. In Burkholderia thailandensis, a major facilitator transport protein (MFTP) efflux pump was discovered which is regulated by MftR global transcription factor [10]. MftR is a global regulator of secondary metabolite in Burkholderia thailandensis [11]. Also, MftR was reported to regulate approximately 400 genes in Burkholderia thailandensis [11].

In the current study we aim to study the binding sites of global transcription regulator MftR in Burkholderia thailandensis. In order to do that we are going to use ChIP-seq technique. Chromatin immunoprecipitation (ChIP) is a technique which is used to study protein-nucleic acid interaction in a specific matter. In ChIP, first protein of interest is crosslinked with nucleic acid which is followed by sonication to fragment nucleic acid and finally a specific antibody is used to enrich nucleic acid binding fragments [12].

ChIP can be coupled with PCR, qPCR or Next-generation sequencing. In the current study, we are going to couple ChIP with illumina novaseq 6000 next generation sequencing. By using ChIP-seq, we should be able to identify genes under direct regulation of MftR.  
Methods and Materials Currently I am still working on the wet lab experiments. I will use 3 samples of wild type Burkholderia thailandensis and 3 samples of disrupted mftR strain. I have searched through NCBI Gene Expression Omnibus (GEO) database and found quite similar dataset under GEO accession (GSE61334). They performed ChIP-seq for the same bacterial species with different strain. Link to download the data: <https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE61334>

• For this course, I have adapted R pipeline from systemPipeRdata package that can downloaded from <https://github.com/tgirke/systemPipeRdata/blob/master/vignettes/systemPipeChIPseq.Rmd> with minor modifications • The R code can be accessed through RChIP\_seq\_code Results The typical pipeline for ChIP-seq analysis incuded standard steps. First of all, quality control is performed on fastq file which are generated from sequencing machine. After that, over expressed sequences are trimmed from both ends. Then Burkholderia cenocepacia reference genome was downloaded from <https://www.ncbi.nlm.nih.gov/genome/?term=Burkholderia+cenocepacia>. then fastq files were aligned to the efference genome. To identify protein-DNA, peak calling was applied to the aligned reads. Finally, Motif Enrichment Analysis was performed. FASTQ quality report which will be used later to filter DNA sequences.

```{r fastq\_report, eval=FALSE}

fqlist <- seeFastq(fastq=infile1(trim), batchsize=100000, klength=8)

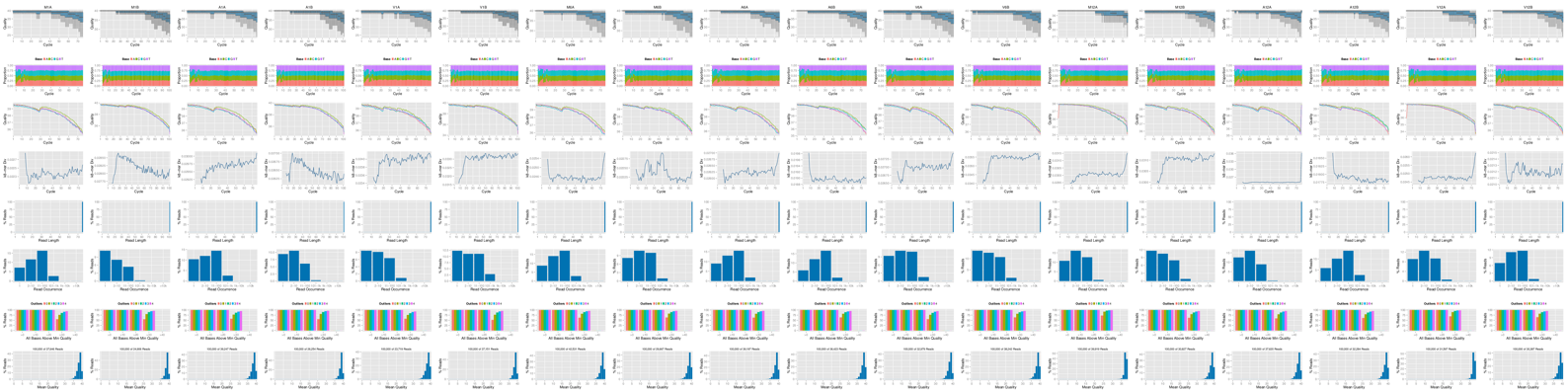
pdf("./results/fastqReport.pdf", height=18, width=4\*length(fqlist))

seeFastqPlot(fqlist)

dev.off()

```

![](results/fastqReport.png)



“FASTQ quality control metrics”

Relative to wildtype B. cenocepacia K56-2, the only genomic region that was found to be enriched by the ChIP-seq analysis of the CzcR-FLAG strain mapped to nucleotide coordinates 787809-798988 on chromosome 2.

## bcf

args <- loadWorkflow(targets = './results/targets\_report\_bcf.txt',

wf\_file = "combine.cwl", input\_file = "varseq.yml", dir\_path = dir\_path)

args <- renderWF(args, inputvars = c(FileName1 = "\_FILE1\_", SampleName = '\_SampleName\_'))

varlist <- sapply(names(subsetWF(args[1:2], slot='output', subset=1, index=1)),

function(x) as.character(read.delim(subsetWF(args[1:2], slot='output', subset=1, index=1)[x])$VARID))

vennset\_bcf <- overLapper(varlist, type="vennsets")

pdf("./results/vennplot\_var.pdf")

vennPlot(list(vennset\_gatk, vennset\_bcf), mymain="", mysub="GATK: red; BCFtools: blue", colmode=2, ccol=c("red", "blue"))

dev.off()

```

![](results/seqlogo.png)

Graphical representation of the sequence conservation of transcription factor binding site

Logo

Description automatically generated

Discussion:

The aim of this study is to identify genes that are under direct control of MftR transcription factor in *Burkholderia thailandensis.* I have tested 4 genes that were differentially expressed in disrupted MftR strain usingChIP-PCR and I identified 2 genes under direct control of MftR “unpublished data”. One gene is called btaR2 which is an ATP-dependent transcription regulator that regulate quorum sensing. The other gene is called BhcM which is a transcription factor which regulates Histone deacetylase inhibitors. This preliminary result gave us confidence to procced with ChIP-seq experiment.

The data we used is for CzcR transcription factor which is essential for heavy metal resistance and virulence in *Burkholderia cenocepacia.*from the results of the ChIP-seq analysis, the only binding site of CzcR transcription factor mapped to nucleotide coordinates 787809-798988 on chromosome 2. That means only one gene is under direct control of CzcR but there might be other genes that are under indirect control of CzcR.

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