Bioconductor Regulatory Genomics Workflow

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Abstract The identification of therapeutic targets is a critical step in the research and developement of new drugs, with several drug discovery programmes failing because of a weak linkage between target and disease. Genome-wide association studies and large-scale gene expression experiments are providing insights into the biology of several common and complex diseases, but the complexity of transcriptional regulation mechanisms often limit our understanding of how genetic variation can influence changes in gene expression. Several initiatives in the field of regulatory genomics are aiming to close this gap by systematically identifying and cataloguing regulatory elements such as promoters and enhacers across different tissues and cell types. In this Bioconductor workflow, we will explore how different types of regulatory genomic data can be used for the functional interpretation of disease-associated variants and for the prioritisation of gene lists from gene expression experiments.

Keywords

bioconductor; r; rstats; regulatory genomics; functional genomics; genetics; gwas; transcriptomics; integration; multiomics

Introduction

Discovering and bringing new drugs to the market is a long, expensive and inefficient process $^{1;2}$. Increasing the success rates of drug discovery programmes would be transformative to the pharmaceutical industry and significantly improve patients $\hat{a} \in ^{TM}$ access to medicines. Of note, the majority of drug discovery programmes fail for efficacy reasons 3 , with up to 40% of these failures due to lack of a clear link between the target and the disease under investigation 4 .

Target selection, the first step in drug discovery programmes, is thus a critical decision point. It has previously been shown that therapeutic targets with a genetic link to the disease under investigation are more likely to progress through the drug discovery pipeline, suggesting that genetics can be used as a tool to prioritise and validate drug targets in early discovery^{5,6}.

Over the last decade, genome-wide association studies (GWASs) have revolutionised the field of human genetics allowing to survey DNA mutations associated with disease and other complex traits on an unprecedented scale⁷. Similarly, phenome-wide association studies (PheWAS) are emerging as a complementary methodology to decipher the genetic bases of the human phenome⁸. While many of these associations might not actually be relevant for the disease aetiology⁹, these methods hold much promise to guide pharmaceutical scientists towards the next generation of drug targets ¹⁰.

Arguably, one of the biggest challenges in translating findings from GWASs to therapies is that the great majority of single nucleotide polymorphisms (SNPs) associated with disease are found in non-coding regions of the genome and therefore cannot be easily linked to a target gene ¹¹. Many of these SNPs could be regulatory variants, affecting the expression of nearby or distal genes by interfering with the process of transcription (e.g.: binding of transcription factors at promoters or enhancers) ¹².

The most established way to map disease-associated regulatory variants to target genes is probably to use expression quantitative trait loci (eQTLs)¹³, variants that affect the expression of specific genes. Over the last few years, the GTEx consortium assembled a valuable resource by performing large-scale mapping of genome-wide correlations between genetic variants and gene expression across 44 human tissues¹⁴.

However, depending on the power of the study, it might not be possible to detect all existing regulatory variants as eQTLs. An alternative is to use information on the location of promoters and distal enhancers across the genome and link these regulatory elements to their target genes. Large, multi-centre Initiatives such as ENCODE ¹⁵, Roadmap Epigenomics ¹⁶ and BLUEPRINT ^{17;18} mapped regulatory elements in the genome by profiling a number of chromatin features including DNase hypersensitive sites (DHSs), several types of histone marks and binding of chromatin-associated proteins in a large number of cell lines, primary cell types and tissues. Similarly, the FANTOM consortium used cap analysis of gene expression (CAGE) to identify promoters and enhancers across hundreds of cells and tissues ¹⁹.

Knowing that a certain stretch of DNA is an enhancer is however not informative of the target gene(s). One way to infer links between enhancers and promoters *in silico* is to identify significant correlations across a large panel of cell types, an approach that was used for distal and promoter DHSs²⁰ as well as for CAGE-defined promoters and enhancers²¹. Experimental methods to assay interactions between regulatory elements also exist. Chromatin interaction analysis by paired-end tag sequencing (ChIA-PET)^{22;23} couples chromatin immunoprecipitation with DNA ligation and sequencing to identify regions of DNA that are interacting thanks to the binding of a specific protein. Promoter capture Hi-C^{24;25} extends chromatin conformation capture by using "baits" to enrich for promoter interactions and increase resolution.

Overall, linking genetic variants to their candidate target genes is not straightforward, not only because of the complexity of the human genome and transcriptional regulation, but also because of the variety of data types and approaches that can be used. To address this, we developed STOPGAP (systematic target opportunity assessment by genetic association predictions), a database of disease variants mapped to their most likely target gene(s) using different types of regulatory genomic data²⁶. The database is currently undergoing a major overhaul and will eventually be superseded by POSTGAP²⁷. A similar resource and valid alternative is INFERNO (inferring the molecular mechanisms of noncoding variants)²⁸.

Workflow

Overview

In this workflow we will explore how regulatory genomic data can be used to connect the genetic and transcriptional layers by providing a framework for the functional annotation of SNPs from GWASs. We will use eQTL data from GTEx ¹⁴, FANTOM5 correlations between promoters and enhancers ²¹ and promoter capture Hi-C data [Javierre2016].

We start with a common scenario: we run a RNA-seq experiment comparing patients with a disease and healthy individuals, and would like to discover key disease genes and potential therapeutic targets by integrating genetic information in our analysis.

Install required packages

R version 3.4.2 and Bioconductor version 3.6 were used for the analysis. The code below will install all required packages and dependencies from Bioconductor and CRAN:

```
source("https://bioconductor.org/biocLite.R")
# uncomment the following line to install packages
#biocLite(c("DESeq2", "GenomicFeatures", "GenomicRanges", "ggplot2", "gwascat", "recount", "pheatmanger")
```

Gene expression data and differential gene expression analysis

The RNA-seq data we will be using comes from blood of patients with systemic lupus erythematosus (SLE) and healthy controls ²⁹.

We are going to use recount 30 to obtain gene-level counts:

```
library(recount)
# uncomment the following line to download dataset
#download_study("SRP062966")
load(file.path("SRP062966", "rse_gene.RData"))
rse <- scale_counts(rse_gene)
rse

## class: RangedSummarizedExperiment
## dim: 58037 117
## metadata(0):
## assays(1): counts
## rownames(58037): ENSG00000000003.14 ENSG00000000005.5 ...
## ENSG00000283698.1 ENSG00000283699.1
## rowData names(3): gene_id bp_length symbol
## colnames(117): SRR2443263 SRR2443262 ... SRR2443147 SRR2443149
## colData names(21): project sample ... title characteristics</pre>
```

Other Bioconductor packages that can be used to access data from gene expression experiments directly in R are GEOquery ³¹ and ArrayExpress ³².

So, we have 117 samples. This is what the data looks like:

```
assay(rse)[1:10, 1:10]
```

##		SRR2443263	SRR2443262	SRR2443261	SRR2443260	SRR2443259
##	ENSG00000000003.14	19	6	10	10	8
##	ENSG00000000005.5	0	0	0	0	0
##	ENSG00000000419.12	489	238	224	323	281
##	ENSG00000000457.13	594	503	530	670	775
##	ENSG00000000460.16	232	173	166	252	268
##	ENSG00000000938.12	21554	18918	14260	19869	26586
##	ENSG00000000971.15	94	57	45	59	35
##	ENSG00000001036.13	500	397	358	407	500
##	ENSG00000001084.10	373	298	336	367	391
##	ENSG00000001167.14	827	832	837	1091	1013
##		SRR2443258	SRR2443257	SRR2443256	SRR2443255	SRR2443254
##	ENSG0000000003.14	SRR2443258 6	SRR2443257 2	SRR2443256 24	SRR2443255 21	SRR2443254 11
	ENSG00000000003.14 ENSG00000000005.5				21112 110200	21112 110201
##		6	2	24	21	11
## ##	ENSG0000000005.5	6	2	24	21	11 0
## ## ##	ENSG0000000005.5 ENSG00000000419.12	6 0 333	2 0 214	24 0 390	21 0 270	11 0 359
## ## ## ##	ENSG0000000005.5 ENSG00000000419.12 ENSG00000000457.13	6 0 333 712	2 0 214 461	24 0 390 603	21 0 270 613	11 0 359 609
## ## ## ##	ENSG00000000015.5 ENSG00000000419.12 ENSG00000000457.13 ENSG00000000460.16	6 0 333 712 263	2 0 214 461 160	24 0 390 603 228	21 0 270 613 245	11 0 359 609 234
## ## ## ## ##	ENSG00000000005.5 ENSG00000000419.12 ENSG00000000457.13 ENSG00000000460.16 ENSG000000000938.12	6 0 333 712 263 17377	2 0 214 461 160 19981	24 0 390 603 228 15136	21 0 270 613 245 13039	11 0 359 609 234 16994
## ## ## ## ##	ENSG00000000005.5 ENSG00000000419.12 ENSG00000000457.13 ENSG00000000460.16 ENSG00000000938.12 ENSG000000000971.15	6 0 333 712 263 17377 76	2 0 214 461 160 19981 26	24 0 390 603 228 15136 53	21 0 270 613 245 13039 60	11 0 359 609 234 16994 50
## ## ## ## ## ##	ENSG00000000005.5 ENSG00000000419.12 ENSG00000000457.13 ENSG00000000460.16 ENSG00000000938.12 ENSG00000000971.15 ENSG00000001036.13	6 0 333 712 263 17377 76 714	2 0 214 461 160 19981 26 364	24 0 390 603 228 15136 53 575	21 0 270 613 245 13039 60 438	11 0 359 609 234 16994 50 638

We note that genes are annotated using the GENCODE³³ v25 annotation, which will be useful later on. Let's look at the metadata to check how we can split samples between cases and controls:

colData(rse)

```
## DataFrame with 117 rows and 21 columns
                  project
                                sample experiment
                                                            run
##
              <character> <character> <character> <character>
## SRR2443263
                SRP062966 SRS1048033 SRX1168388 SRR2443263
## SRR2443262
                SRP062966 SRS1048034 SRX1168387
                                                     SRR2443262
## SRR2443261
                SRP062966 SRS1048035
                                        SRX1168386
                                                     SRR2443261
## SRR2443260
                SRP062966 SRS1048036 SRX1168385
                                                     SRR2443260
## SRR2443259
                SRP062966
                            SRS1048037
                                        SRX1168384
                                                     SRR2443259
##
                       . . .
                                   . . .
## SRR2443151
                SRP062966
                            SRS1048145
                                        SRX1168276
                                                     SRR2443151
## SRR2443150
                SRP062966
                            SRS1048146
                                        SRX1168275
                                                     SRR2443150
## SRR2443148
                SRP062966
                            SRS1048147
                                        SRX1168273
                                                     SRR2443148
## SRR2443147
                SRP062966 SRS1048148 SRX1168272
                                                     SRR2443147
## SRR2443149
                SRP062966 SRS1048149 SRX1168274
                                                     SRR2443149
##
              read_count_as_reported_by_sra reads_downloaded
##
                                   <integer>
                                                     <integer>
                                   103977424
## SRR2443263
                                                     103977424
## SRR2443262
                                   125900891
                                                     125900891
## SRR2443261
                                   129803063
                                                     129803063
## SRR2443260
                                   105335395
                                                     105335395
## SRR2443259
                                   101692332
                                                     101692332
##
## SRR2443151
                                    87315854
                                                      87315854
## SRR2443150
                                    96825506
                                                      96825506
## SRR2443148
                                                     121365435
                                   121365435
## SRR2443147
                                   104038425
                                                     104038425
## SRR2443149
                                   113083096
                                                     113083096
              proportion_of_reads_reported_by_sra_downloaded paired_end
##
##
                                                     <numeric>
                                                                <logical>
## SRR2443263
                                                                     FALSE
                                                             1
## SRR2443262
                                                             1
                                                                     FALSE
## SRR2443261
                                                                     FALSE
                                                             1
## SRR2443260
                                                                     FALSE
                                                             1
## SRR2443259
                                                                     FALSE
                                                             1
## ...
## SRR2443151
                                                                     FALSE
                                                             1
## SRR2443150
                                                                     FALSE
                                                             1
## SRR2443148
                                                             1
                                                                     FALSE
## SRR2443147
                                                                     FALSE
                                                             1
## SRR2443149
                                                                     FALSE
##
              sra_misreported_paired_end mapped_read_count
##
                                <logical>
                                                   <integer> <numeric>
## SRR2443263
                                                   103499268 5149333280
                                    FALSE
## SRR2443262
                                    FALSE
                                                   125499809 6244059473
## SRR2443261
                                    FALSE
                                                   125043355 6201504759
## SRR2443260
                                    FALSE
                                                   104872856 5211910530
## SRR2443259
                                                   101258496 5033612693
                                    FALSE
##
  . . .
                                       . . .
## SRR2443151
                                    FALSE
                                                    86874384 4319264868
                                                    96316303 4787601223
## SRR2443150
                                    FALSE
## SRR2443148
                                                   120819733 6009515064
                                    FALSE
## SRR2443147
                                    FALSE
                                                   103588909 5153702232
                                                   112640054 5598306153
## SRR2443149
                                    FALSE
##
              sharq_beta_tissue sharq_beta_cell_type
##
                     <character>
                                          <character>
## SRR2443263
                              NA
## SRR2443262
                              NA
                                                    NΑ
## SRR2443261
                              NA
                                                    NA
## SRR2443260
                              NΑ
                                                    NΑ
## SRR2443259
                              NΑ
                                                    NΑ
## ...
                             . . .
                                                   . . .
## SRR2443151
                              NΑ
                                                    NΑ
## SRR2443150
                              NΑ
                                                    NA
```

```
## SRR2443148
                             NA
                                                  NA
## SRR2443147
                             NΑ
                                                  NΑ
## SRR2443149
                             NΑ
                                                  NΑ
##
              biosample_submission_date biosample_publication_date
                            <character>
                                                       <character>
## SRR2443263
                2015-08-28T16:41:29.000
                                           2015-09-16T01:24:17.350
## SRR2443262
               2015-08-28T16:41:28.000
                                         2015-09-16T01:24:16.410
## SRR2443261
               2015-08-28T16:41:27.000
                                         2015-09-16T01:24:14.823
## SRR2443260
                2015-08-28T16:41:35.000
                                           2015-09-16T01:24:13.450
## SRR2443259
               2015-08-28T16:41:33.000
                                           2015-09-16T01:24:12.433
## ...
## SRR2443151
               2015-08-28T16:42:24.000
                                          2015-09-16T01:19:06.787
## SRR2443150
               2015-08-28T16:42:23.000
                                         2015-09-16T01:19:05.557
               2015-08-28T16:42:21.000
## SRR2443148
                                          2015-09-16T01:20:16.080
                2015-08-28T16:42:19.000
                                           2015-09-16T01:20:14.923
## SRR2443147
## SRR2443149
                2015-08-28T16:42:22.000
                                           2015-09-16T01:19:04.583
##
                biosample_update_date avg_read_length geo_accession
##
                          <character>
                                            <integer>
                                                        <character>
## SRR2443263 2015-09-16T01:28:05.297
                                                   50
                                                         GSM1863749
## SRR2443262 2015-09-16T01:28:05.027
                                                   50
                                                         GSM1863748
## SRR2443261 2015-09-16T01:28:04.803
                                                         GSM1863747
                                                   50
## SRR2443260 2015-09-16T01:28:04.587
                                                   50
                                                         GSM1863746
## SRR2443259 2015-09-16T01:28:04.347
                                                   50
                                                         GSM1863745
## ...
## SRR2443151 2015-09-16T01:23:41.897
                                                   50
                                                         GSM1863637
## SRR2443150 2015-09-16T01:23:41.453
                                                   50
                                                         GSM1863636
## SRR2443148 2015-09-16T01:23:41.093
                                                   50
                                                         GSM1863634
## SRR2443147 2015-09-16T01:23:40.840
                                                   50
                                                         GSM1863633
## SRR2443149 2015-09-16T01:23:40.597
                                                   50
                                                         GSM1863635
##
               bigwig_file
                                  title
##
                <character> <character>
## SRR2443263 SRR2443263.bw control18
## SRR2443262 SRR2443262.bw
                             control17
                             control16
## SRR2443261 SRR2443261.bw
## SRR2443260 SRR2443260.bw
                             control15
## SRR2443259 SRR2443259.bw
                             control14
## ...
                                    . . .
## SRR2443151 SRR2443151.bw
                                   SLE5
## SRR2443150 SRR2443150.bw
                                   SLE4
## SRR2443148 SRR2443148.bw
                                   SLE2
## SRR2443147 SRR2443147.bw
                                   SLE1
## SRR2443149 SRR2443149.bw
                                   SLE3
##
                                                                                        characterist:
##
                                                                                        <CharacterLi
## SRR2443263
                                      disease status: healthy, tissue: whole blood, anti-ro: control,
## SRR2443262
                                      disease status: healthy, tissue: whole blood, anti-ro: control,
## SRR2443261
                                      disease status: healthy, tissue: whole blood, anti-ro: control,
## SRR2443260
                                      disease status: healthy, tissue: whole blood, anti-ro: control,
## SRR2443259
                                      disease status: healthy, tissue: whole blood, anti-ro: control,
## ...
## SRR2443151 disease status: systemic lupus erythematosus (SLE),tissue: whole blood,anti-ro: med,
## SRR2443150 disease status: systemic lupus erythematosus (SLE), tissue: whole blood, anti-ro: high,
## SRR2443148 disease status: systemic lupus erythematosus (SLE),tissue: whole blood,anti-ro: high,
## SRR2443147 disease status: systemic lupus erythematosus (SLE), tissue: whole blood, anti-ro: high,
## SRR2443149 disease status: systemic lupus erythematosus (SLE), tissue: whole blood, anti-ro: high,
```

The most interesting part of the metadata is contained in the characteristics column, which is a CharacterList object:

```
## CharacterList of length 117
## [[1]] disease status: healthy tissue: whole blood anti-ro: control ism: control
## [[2]] disease status: healthy tissue: whole blood anti-ro: control ism: control
## [[3]] disease status: healthy tissue: whole blood anti-ro: control ism: control
```

```
## [[4]] disease status: healthy tissue: whole blood anti-ro: control ism: control
## [[5]] disease status: healthy tissue: whole blood anti-ro: control ism: control
## [[6]] disease status: healthy tissue: whole blood anti-ro: control ism: control
## [[7]] disease status: healthy tissue: whole blood anti-ro: control ism: control
## [[8]] disease status: healthy tissue: whole blood anti-ro: control ism: control
## [[9]] disease status: healthy tissue: whole blood anti-ro: control ism: control
## [[10]] disease status: healthy tissue: whole blood anti-ro: control ism: control
## ...
## <107 more elements>
```

Let's create some new columns with this information that can be used for the differential expression analysis. We will also make sure that they are encoded as factors and that the correct reference layer is used:

```
# disease status
colData(rse)$disease_status <- sapply(colData(rse)$characteristics, "[", 1)</pre>
colData(rse) $disease_status <- sub("disease status: ", "", colData(rse) $disease_status)
colData(rse)$disease_status <- sub("systemic lupus erythematosus \\(SLE\\)", "SLE", colData(rse)$dis
colData(rse) $disease_status <- factor(colData(rse) $disease_status, levels = c("healthy", "SLE"))
# tissue
colData(rse)$tissue <- sapply(colData(rse)$characteristics, "[", 2)</pre>
colData(rse)$tissue <- sub("tissue: ", "", colData(rse)$tissue)</pre>
colData(rse)$tissue <- factor(colData(rse)$tissue)</pre>
# anti-ro
colData(rse)$anti_ro <- sapply(colData(rse)$characteristics, "[", 3)</pre>
colData(rse)$anti_ro <- sub("anti-ro: ", "", colData(rse)$anti_ro)</pre>
colData(rse)$anti_ro <- factor(colData(rse)$anti_ro)</pre>
colData(rse)$ism <- sapply(colData(rse)$characteristics, "[", 4)</pre>
colData(rse)$ism <-sub("ism: ", "", colData(rse)$ism)</pre>
colData(rse)$ism <- factor(colData(rse)$ism)</pre>
```

We can have a look at the new format:

```
colData(rse)[c("disease_status", "tissue", "anti_ro", "ism")]
```

```
## DataFrame with 117 rows and 4 columns
             disease_status
                              tissue anti_ro
                                                     ism
##
                   <factor>
                              <factor> <factor> <factor>
## SRR2443263
                    healthy whole blood control control
## SRR2443262
                    healthy whole blood control control
## SRR2443261
                    healthy whole blood control control
## SRR2443260
                    healthy whole blood control control
## SRR2443259
                    healthy whole blood control control
                                            . . .
                        SLE whole blood
## SRR2443151
                                            med ISM_low
## SRR2443150
                        SLE whole blood
                                          high ISM_low
## SRR2443148
                        SLE whole blood
                                           high ISM_high
## SRR2443147
                        SLE whole blood
                                           high ISM_high
## SRR2443149
                        SLE whole blood
                                           high ISM_high
```

It looks more readable. Let's now check how many samples we have in each group:

```
table(colData(rse) $disease_status)
```

```
## ## healthy SLE ## 18 99
```

To speed up code execution we will limit the number of SLE samples. For simplicity, we select the first 18 (healthy) and the last 18 (SLE) samples from the original RangedSummarizedExperiment object:

```
rse <- rse[, c(1:18, 82:99)]
```

Now we are ready to perform a simple differential gene expression analysis with DESeq234:

colData names(27): project sample ... sizeFactor replaceable

```
library(DESeq2)
dds <- DESeqDataSet(rse, ~ disease_status)
dds <- DESeq(dds)
dds

## class: DESeqDataSet
## dim: 58037 36
## metadata(1): version
## assays(5): counts mu cooks replaceCounts replaceCooks
## rownames(58037): ENSG00000000003.14 ENSG00000000005.5 ...
## ENSG00000283698.1 ENSG00000283699.1
## rowData names(25): gene_id bp_length ... maxCooks replace
## colnames(36): SRR2443263 SRR2443262 ... SRR2443166 SRR2443165</pre>
```

Note that we used an extremely simple model; in the real world you will probably need to account for covariables, potential confounders and interactions between them. edgeR ³⁵ and limma ³⁶ are good alternatives to DESEq2 for performing differential expression analyses.

We can now look at the data in more detail. We use the variance stabilising transformation (VST) ³⁷ for visualisation purposes:

```
vsd <- vst(dds, blind = FALSE)</pre>
```

First, let's look at distances between samples to see if we can recover a separation between SLE and healthy samples:

```
sampleDists <- as.matrix(dist(t(assay(vsd))))
rownames(sampleDists) <- vsd$disease_status
sampleDists[c(1, 18, 19, 36), c(1, 18, 19, 36)]

## SRR2443263 SRR2443248 SRR2443182 SRR2443165
## healthy 0.00000 106.6933 93.30292 99.84061
## healthy 106.69330 0.0000 115.87958 127.27997
## SLE 93.30292 115.8796 0.00000 115.06568
## SLE 99.84061 127.2800 115.06568 0.00000</pre>
```

We will use the pheatmap ³⁸ and RColorBrewer ³⁹ packages for drawing the heatmap (Figure 1):

```
library(pheatmap)
library(RColorBrewer)
colors <- colorRampPalette(rev(brewer.pal(9, "Blues")))(255)
pheatmap(sampleDists, col = colors)</pre>
```

Similarly, we can perform a principal component analysis (PCA) on the most variable 500 genes (Figure 2):

```
plotPCA(vsd, intgroup = "disease_status")
```

This looks better, we can see some separation of healthy and SLE samples along both PC1 and PC2, though some SLE samples appear very similar to the healthy ones. Next, we select genes that are differentially expressed below a 0.05 adjusted p-value threshold:

```
res <- results(dds, alpha = 0.05)
res

## log2 fold change (MLE): disease status SLE vs healthy
## Wald test p-value: disease status SLE vs healthy
## DataFrame with 58037 rows and 6 columns
## baseMean log2FoldChange lfcSE stat
## <numeric> <numeric> <numeric> <numeric></numeric>
```

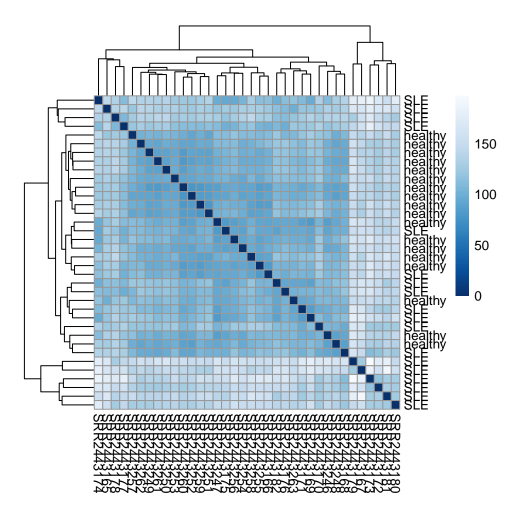


Figure 1. Clustered heatmap showing distances between samples.

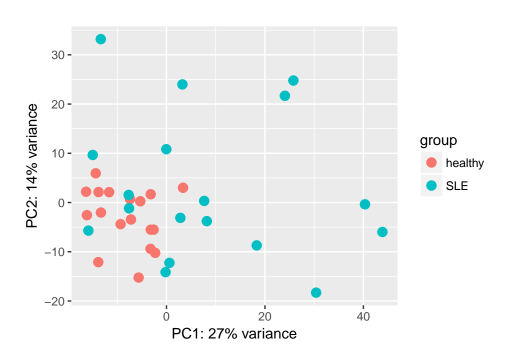


Figure 2. Principal component analysis with samples coloured according to their disease status.

```
## ENSG0000000003.14 10.4189981
                                    -0.20051804 0.24868451 -0.80631496
## ENSG0000000005.5
                                    0.03330732 2.96442394
                       0.0317823
                                                           0.01123568
## ENSG0000000419.12 389.9025130
                                    0.66288230 0.11427371
                                                            5.80082925
## ENSG00000000457.13 636.6928414
                                    0.17336365 0.08062862
                                                           2.15015047
## ENSG00000000460.16 234.6479796
                                    0.20589404 0.07445624
                                                            2.76530274
## ...
## ENSG00000283695.1
                      0.0000000
                                            NΑ
                                                       NΑ
                                                                   NΑ
## ENSG00000283696.1
                     19.1311904
                                   0.252144173 0.1545613
                                                          1.631353425
## ENSG00000283697.1
                     14.9180870
                                   0.179070242 0.1522931 1.175826692
## ENSG00000283698.1
                       0.2289885
                                   0.021962044 1.1315739 0.019408404
## ENSG00000283699.1
                       0.5398951
                                  ##
                           pvalue
                                         padj
##
                        <numeric>
                                    <numeric>
## ENSG0000000003.14 4.200613e-01 6.706002e-01
## ENSG0000000005.5 9.910354e-01
## ENSG00000000419.12 6.598777e-09 3.058479e-06
## ENSG00000000457.13 3.154331e-02 1.463634e-01
## ENSG0000000460.16 5.686999e-03 4.643041e-02
## ...
                              . . .
## ENSG00000283695.1
                              ΝA
                                           ΝA
## ENSG00000283696.1
                                    0.3075119
                        0.1028158
## ENSG00000283697.1
                        0.2396641
                                    0.4987872
## ENSG00000283698.1
                        0.9845153
                                           NΑ
## ENSG00000283699.1
                        0.9967822
                                           NΑ
```

We can look at a summary of the results:

```
summary(res)
```

```
##
## out of 43005 with nonzero total read count
## adjusted p-value < 0.05
## LFC > 0 (up) : 2526, 5.9%
## LFC < 0 (down) : 1069, 2.5%
## outliers [1] : 0, 0%
## low counts [2] : 14735, 34%
## (mean count < 1)
## [1] see 'cooksCutoff' argument of ?results
## [2] see 'independentFiltering' argument of ?results</pre>
```

We can also visualise the log fold changes using an MA plot (Figure 3):

```
plotMA(res, ylim = c(-5,5))
```

For convenience, we will save our differentially expressed genes (DEGs) in another object:

```
degs <- subset(res, padj < 0.05)
degs <- as.data.frame(degs)
head(degs)</pre>
```

```
##
                       baseMean log2FoldChange
                                                     lfcSE
## ENSG0000000419.12
                      389.90251
                                     0.6628823 0.11427371
                                                           5.800829
## ENSG0000000460.16
                                     0.2058940 0.07445624
                      234.64798
                                                           2.765303
## ENSG00000002549.12 1970.95648
                                     0.8657769 0.25181202 3.438187
## ENSG0000003096.13
                       11.18475
                                    -0.7894018 0.25613621 -3.081961
## ENSG0000003147.17
                       71.79432
                                     0.6113739 0.15162606 4.032116
## ENSG00000003249.13 119.18587
                                    -0.8520562 0.27061961 -3.148538
##
                           pvalue
                                           padi
## ENSG0000000419.12 6.598777e-09 3.058479e-06
## ENSG00000000460.16 5.686999e-03 4.643041e-02
## ENSG00000002549.12 5.856225e-04 9.776328e-03
## ENSG00000003096.13 2.056419e-03 2.291728e-02
## ENSG00000003147.17 5.527679e-05 1.927054e-03
## ENSG00000003249.13 1.640893e-03 1.955034e-02
```

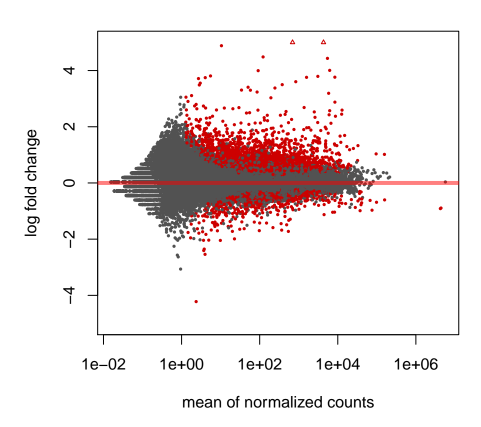


Figure 3. MA plot showing genes differentially expressed in SLE patients compared to healthy patients.

We also map the GENCODE gene IDs to gene symbols using the annotation in the original RangedSummarizedExperiment object, which is going to be convenient later on:

```
## DataFrame with 58037 rows and 3 columns
                    gene_id bp_length
                                                symbol
##
                <character> <integer> <CharacterList>
## 1
         ENSG0000000003.14
                                 4535
                                                TSPAN6
## 2
         ENSG00000000005.5
                                 1610
                                                  TNMD
         ENSG00000000419.12
## 3
                                                  DPM1
                                 1207
         ENSG00000000457.13
## 4
                                 6883
                                                 SCYL3
## 5
         ENSG0000000460.16
                                 5967
                                              C1orf112
## ...
                                   . . .
## 58033 ENSG00000283695.1
                                   61
                                                    NΑ
## 58034 ENSG00000283696.1
                                  997
                                                    NΑ
## 58035
         ENSG00000283697.1
                                  1184
                                          LOC101928917
## 58036
         ENSG00000283698.1
                                  940
                                                    NΑ
## 58037
         ENSG00000283699.1
                                   60
                                               MIR4481
degs <- merge(rowData(rse), degs, by.x = "gene_id", by.y = "row.names", all = FALSE)
tail(degs)
## DataFrame with 6 rows and 9 columns
```

```
##
                    gene_id bp_length symbol
                                               baseMean log2FoldChange
##
                <character> <integer> <list>
                                              <numeric>
                                                             <numeric>
## [3590,] ENSG00000283444.1
                                  831
                                          NΑ
                                               2.756993
                                                             1.3404014
## [3591,] ENSG00000283479.1
                                  420
                                          NΑ
                                              1.928773
                                                             1.9512651
                                        ASPH 277.956104
## [3592,] ENSG00000283485.1
                                 2190
                                                             1.3415229
## [3593,] ENSG00000283571.1
                                 306
                                          NA 1.791920
                                                             1.8502738
## [3594,] ENSG00000283602.1
                                 2089
                                          NA 130.233552
                                                             0.5752086
## [3595,] ENSG00000283623.1
                                        ATG5 107.731105
                                                             0.4144398
                                    pvalue
              lfcSE
                         stat
                                                  padi
##
          <numeric> <numeric>
                                 <numeric>
                                             <numeric>
## [3590,] 0.4729127 2.834353 0.0045918633 0.040127193
## [3591,] 0.5681341 3.434515 0.0005936154 0.009822205
## [3592,] 0.3694185 3.631445 0.0002818390 0.005898176
## [3593,] 0.6557494 2.821617 0.0047782147 0.041137170
## [3594,] 0.2047652 2.809112 0.0049678327 0.042178839
## [3595,] 0.1066472 3.886081 0.0001018754 0.002951150
```

Accessing GWAS data

Genome: GRCh38

rowData(rse)

We have more than 3500 genes of interest at this stage. Since we know that therapeutic targets with genetic evidence are more likely to progress through the drug discovery pipeline ⁶, one way to prioritise them could be to check which of these can be genetically linked to SLE. To get hold of relevant GWAS data, we will be using the gwascat Bioconductor package ⁴⁰, which provides an interface to the GWAS catalog ⁴¹. An alternative is to use the GRASP ⁴² database with the grasp2db ⁴³ package.

```
library(gwascat)
# uncomment the following line to download file and build the gwasloc object all in one step
#snps <- makeCurrentGwascat()
# uncomment the following line to download file
#download.file("http://www.ebi.ac.uk/gwas/api/search/downloads/alternative", destfile = "gwas_catalos_snps <- read.delim("gwas_catalos_v1.0.1-associations_e90_r2017-12-04.tsv", check.names = FALSE, strs.snps <- gwascat:::gwdf2GRanges(snps, extractDate = "2017-12-04")
genome(snps) <- "GRCh38"
snps</pre>
## gwasloc instance with 61107 records and 37 attributes per record.
## Extracted: 2017-12-04
```

```
## Excerpt:
```

```
## GRanges object with 5 ranges and 3 metadata columns:
##
         segnames
                                  ranges strand | DISEASE/TRAIT
                                                                       SNPS
##
            <Rle>
                               <IRanges> <Rle>
                                                    <character> <character>
             chr1 [203186754, 203186754]
                                             * | YKL-40 levels rs4950928
##
     [1]
                                              *
##
     [2]
            chr13 [ 39776775, 39776775]
                                                      Psoriasis
                                                                  rs7993214
            chr15 [ 78513681, 78513681]
##
     [3]
                                              *
                                                    Lung cancer
                                                                  rs8034191
             chr1 [159711078, 159711078]
##
     [4]
                                              *
                                                    Lung cancer
                                                                  rs2808630
##
     [5]
             chr3 [190632672, 190632672]
                                              *
                                                    Lung cancer
                                                                  rs7626795
##
           P-VALUE
##
         <numeric>
     [1]
##
             1e-13
##
             2e-06
     [2]
##
             3e-18
     [3]
             7e-06
##
     [4]
##
             8e-06
     [5]
##
     seginfo: 23 sequences from GRCh38 genome; no seglengths
```

SNPs is a gwasloc object which is simply a wrapper around a GRanges object, the standard way to express genomic ranges in Bioconductor. We are interested in SNPs associated with SLE:

```
snps <- subsetByTraits(snps, tr = "Systemic lupus erythematosus")</pre>
snps
## gwasloc instance with 402 records and 37 attributes per record.
## Extracted: 2017-12-04
## Genome: GRCh38
## Excerpt:
\#\# GRanges object with 5 ranges and 3 metadata columns:
##
          segnames
                                      ranges strand
##
             <Rle>
                                   <IRanges> <Rle>
##
     Γ17
             chr16 [ 31301932, 31301932]
             chr11 [ 589564,
##
     [2]
                                     589564]
##
     [3]
              chr3 [ 58384450, 58384450]
              chr1 [173340574, 173340574]
##
     [4]
##
     Г5 Т
              chr8 [ 11491677, 11491677]
                                                    *
##
                           DISEASE/TRAIT
                                                   SNPS
                                                           P-VALUE
##
                             <character> <character> <numeric>
##
     [1] Systemic lupus erythematosus
                                           rs9888739
                                                              2e-23
     [2] Systemic lupus erythematosus rs4963128
[3] Systemic lupus erythematosus rs6445975
[4] Systemic lupus erythematosus rs10798269
                                                              3e-10
##
##
                                                              7e-09
##
                                                              1e-07
##
     [5] Systemic lupus erythematosus rs13277113
                                                              1e-10
##
```

seqinfo: 23 sequences from GRCh38 genome; no seqlengths

##

We can visualise these as a Manhattan plot to look at the distribution of GWAS p-values over chromosomes on a negative log scale (Figure 4). Note that p-values lower than 1e-25 are truncated in the figure and that we have to load ggplot2⁴⁴ to modify the look of the plot:

```
library(ggplot2)
traitsManh(gwr = snps, sel = snps, traits = "Systemic lupus erythematosus") +
  theme(legend.position = "none",
        axis.title.x = element_blank(),
        axis.text.x = element_blank())
```

We note here that genotyping arrays typically include a very small fraction of all possible SNPs in the human genome, and there is no guarantee that the *tag* SNPs on the array are the true casual SNPs ⁴⁵. The alleles of other SNPs can be imputed from tag SNPs thanks to the structure of linkage disequilibrium (LD) blocks present in chromosomes. Thus, when linking variants to target genes in a real-world setting, it is important to take into consideration neighbouring SNPs that are in high LD and inherited with the tag SNPs. For simplicity, we will skip this LD expansion step and refer the reader to the Ensembl REST API ^{46;47}, the Ensembl Linkage Disequilibrium Calculator ⁴⁸ and the Bioconductor packages trio ⁴⁹ and ldblock ⁵⁰ to perform this task.

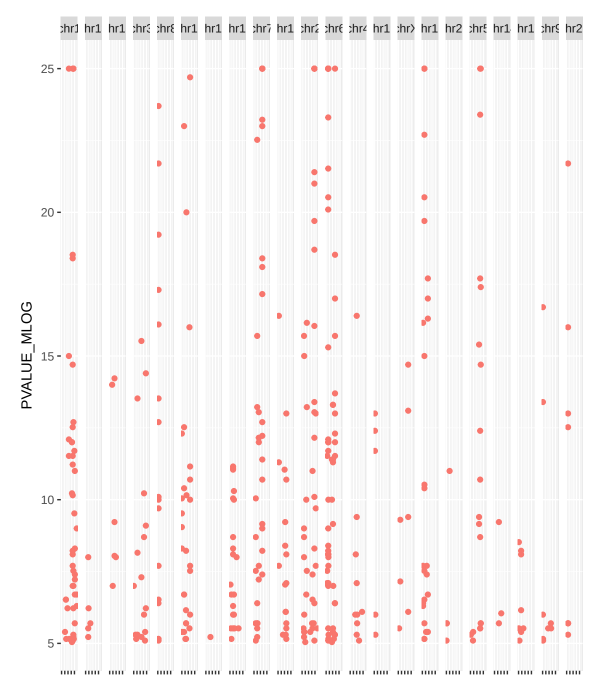


Figure 4. Manhattan plot showing variants significantly associated with SLE.

Annotation of coding and proximal SNPs to target genes

In order to annotate these variants, we need a a TxDb object, a reference of where transcripts are located on the genome. We can build this using the GenomicFeatutres ⁵¹ package and the GENCODE v25 gene annotation:

```
library(GenomicFeatures)
# uncomment the following line to download file
#download.file("ftp://ftp.sanger.ac.uk/pub/gencode/Gencode_human/release_25/gencode.v25.annotation.p
txdb <- makeTxDbFromGFF("gencode.v25.annotation.gff3.gz")</pre>
txdb <- keepStandardChromosomes(txdb)</pre>
txdb
## TxDb object:
## # Db type: TxDb
## # Supporting package: GenomicFeatures
## # Data source: gencode.v25.annotation.gff3.gz
## # Organism: NA
## # Taxonomy ID: NA
## # miRBase build ID: NA
## # Genome: NA
## # transcript_nrow: 198093
## # exon_nrow: 1182765
## # cds_nrow: 704859
## # Db created by: GenomicFeatures package from Bioconductor
## # Creation time: 2018-01-10 11:57:23 +0000 (Wed, 10 Jan 2018)
## # GenomicFeatures version at creation time: 1.30.0
## # RSQLite version at creation time: 2.0
## # DBSCHEMAVERSION: 1.2
We also have to convert the gwasloc object into a standard GRanges object:
snps <- GRanges(snps)</pre>
Let's check if the gwasloc and TxDb object use the same notation for chromosomes:
seqlevelsStyle(snps)
## [1] "UCSC"
seqlevels(snps)
## [1] "chr1" "chr13" "chr15" "chr3" "chr8" "chr11" "chr18" "chr10"
## [9] "chr7" "chr12" "chr2" "chr6" "chr4" "chr19" "chrX" "chr16"
## [17] "chr20" "chr5" "chr14" "chr17" "chr21" "chr9" "chr22"
seqlevelsStyle(txdb)
## [1] "UCSC"
seqlevels(txdb)
## [1] "chr1" "chr2" "chr3" "chr4" "chr5" "chr6" "chr7" "chr8"
   [9] "chr9" "chr10" "chr11" "chr12" "chr13" "chr14" "chr15" "chr16"
## [17] "chr17" "chr18" "chr19" "chr20" "chr21" "chr22" "chrX" "chrY"
```

OK, they do. Now we can annotate our SNPs to genes using the VariantAnnotation 52 package:

```
library(VariantAnnotation)
snps_anno <- locateVariants(snps, txdb, AllVariants())
snps_anno <- unique(snps_anno)
snps_anno</pre>
```

```
## GRanges object with 299 ranges and 9 metadata columns:
##
                                     ranges strand | LOCATION LOCSTART
           seqnames
##
              <Rle>
                                  <IRanges> <Rle> |
                                                         <factor> <integer>
##
       Γ17
              chr16 [ 31301932, 31301932] + |
                                                                       40161
                                                          intron
              chr11 [ 589564, 589564]
chr3 [ 58384450, 58384450]
chr1 [173340574, 173340574]
chr8 [ 11491677, 11491677]
       [2]
                                                  +
##
                                                           intron
                                                                       12531
##
       [3]
                                                  +
                                                                       51074
                                                           intron
##
       [4]
                                                  * | intergenic
                                                                       <NA>
##
       [5]
                                                  * | intergenic
                                                                        < NA >
##
                . . .
                                                . . . .
                                                             . . .
##
     Г295]
               chr6 [137874014, 137874014]
                                                +
                                                                       6162
                                                          intron
##
     [296]
               chr6 [ 32619077, 32619077]
                                                  * | intergenic
                                                                       <NA>
               chr6 [137685367, 137685367]
##
     [297]
                                                  +
                                                          intron
                                                                     11552
##
     [298]
               chrX [153924366, 153924366]
                                                  -
                                                           intron
                                                                       1770
##
     [299]
               chr5 [160459613, 160459613]
                                                  * | intergenic
                                                                        <NA>
##
              LOCEND
                        QUERYID TXID
                                                   CDSID
                                                                         GENEID
##
           <integer> <integer> <character> <IntegerList>
                                                                    <character>
                           1 143788
       [1]
               40161
                                                            ENSG00000169896.16
##
##
               12531
                             2
                                      99581
                                                            ENSG00000070047.11
       [2]
       [3]
               51074
                                       34101
                                                            ENSG00000168297.15
##
                            3
##
       [4]
                <NA>
                             4
                                       <NA>
                                                                           < NA >
                            5
##
       [5]
                <NA>
                                       <NA>
                                                                           <NA>
##
                 . . .
                                        . . .
       . . .
                            . . .
##
     [295]
                6162
                            393
                                      64150
                                                            ENSG00000118503.14
##
     [296]
                < NA >
                            397
                                       <NA>
                                                             ENSG00000230533.2
##
     [297]
               11552
                            398
                                       64145
                                                            ENSG00000089820.15
##
     [298]
                1770
                            399
                                     196900
##
     [299]
                <NA>
                            400
                                        < NA >
                                                                           <NA>
##
                                                                 PRECEDETD
##
                                                           <CharacterList>
##
       Γ17
##
       [2]
##
       [3]
##
             ENSG00000076321.10, ENSG00000117592.8, ENSG00000117593.9, ...
##
       [5] ENSG00000079459.12, ENSG00000136573.12, ENSG00000136574.17, ...
##
##
     [295]
     [296] ENSG00000030110.12, ENSG00000112473.17, ENSG00000112511.17, ...
##
##
     [297]
##
     [298]
     [299] ENSG00000118322.12, ENSG00000145864.12, ENSG00000253417.5, ...
##
##
                                                                   FOLLOWID
##
                                                           <CharacterList>
##
       [1]
       [2]
##
##
       [3]
##
       [4]
            ENSG00000094975.13, ENSG00000117560.7, ENSG00000117586.10, ...
##
            ENSG00000104643.9, ENSG00000154316.15, ENSG00000154319.14, ...
       [5]
##
       . . .
     [295]
##
##
     [296] ENSG00000166278.14,ENSG00000168477.17,ENSG00000196126.10,...
##
     [297]
##
     [298]
##
     [299] ENSG00000113312.10, ENSG00000135083.14, ENSG00000145861.7, ...
##
##
     seqinfo: 23 sequences from GRCh38 genome; no seqlengths
```

We lost all the metadata from the original snps object, but we can recover it using the QUERYID column in snps_anno. We will only keep the SNP IDs and GWAS p-values:

```
snps_metadata <- snps[snps_anno$QUERYID]</pre>
mcols(snps_anno) <- cbind(mcols(snps_metadata)[c("SNPS", "P-VALUE")], mcols(snps_anno))</pre>
snps_anno
## GRanges object with 299 ranges and 11 metadata columns:
##
           seqnames
                                     ranges strand
                                                             SNPS P.VALUE
                                  <IRanges> <Rle> | <character> <numeric>
##
              <Rle>
##
       [1]
              chr16 [ 31301932, 31301932]
                                                + | rs9888739
                                                                       2e-23
##
       [2]
              chr11 [ 589564, 589564]
                                                  + | rs4963128
                                                                       3e - 10
               chr3 [ 58384450, 58384450]
chr1 [173340574, 173340574]
       Гз٦
                                                       rs6445975
                                                                       7e-09
##
                                                 +
##
       Γ47
                                                 * | rs10798269
                                                                       1e-07
               chr8 [ 11491677, 11491677]
##
       [5]
                                                  * | rs13277113
                                                                       1e-10
##
                                                . . . .
                                        . . .
                                                              . . .
       . . .
                                                                         . . .
               chr6 [137874014, 137874014]
                                                        rs5029937
##
     [295]
                                                 + |
                                                                       5e-13
               chr6 [ 32619077, 32619077]
##
     [296]
                                                  *
                                                        rs9271366
                                                                       1e-07
               chr6 [137685367, 137685367]
##
     [297]
                                                 +
                                                        rs6920220
                                                                       4e-07
               chrX [153924366, 153924366]
##
     [298]
                                                  - |
                                                        rs2269368
                                                                       8e-07
                                                 *
               chr5 [160459613, 160459613]
##
     [299]
                                                        rs2431099
                                                                       2e-06
##
             LOCATION LOCSTART LOCEND QUERYID
                                                        TXID
                                                                           CDSID
             <factor> <integer> <integer> <integer> <character> <IntegerList>
##
##
       [1]
                           40161
                                     40161
                                                          143788
               intron
                                                  1
##
                                                    2
       [2]
               intron
                          12531
                                     12531
                                                            99581
##
       [3]
                          51074
                                     51074
                                                    3
                                                            34101
               intron
##
       [4] intergenic
                           <NA>
                                      <NA>
                                                   4
                                                             <NA>
                                                   5
##
       [5] intergenic
                           < N A >
                                      <NA>
                                                             < NA >
##
                                  6162
<NA>
##
                          6162
                                                            64150
     [295]
               intron
                                                  393
                           < N A >
                                                  397
##
     [296] intergenic
                                                             <NA>
##
     Γ297 ]
               intron
                          11552
                                     11552
                                                  398
                                                            64145
                          1770
                                                            196900
##
     [298]
               intron
                                     1770
                                                  399
     [299] intergenic
                            < N A >
                                      <NA>
                                                  400
                                                              <NA>
##
##
                        GENEID
##
                   <character>
       [1] ENSG00000169896.16
##
##
       [2] ENSG00000070047.11
##
       [3] ENSG00000168297.15
       [4]
##
                          < NA >
##
       Г5 Т
                          < NA >
##
       . . .
##
     [295] ENSG00000118503.14
##
     [296]
##
     [297] ENSG00000230533.2
     [298] ENSG00000089820.15
##
##
     [299]
##
                                                                 PRECEDEID
##
                                                           <CharacterList>
##
       Γ17
##
       [2]
       Г31
##
##
       [4]
             ENSG00000076321.10, ENSG00000117592.8, ENSG00000117593.9, ...
##
       [5] ENSG00000079459.12, ENSG00000136573.12, ENSG00000136574.17, ...
##
##
     [295]
##
     [296] ENSG00000030110.12, ENSG00000112473.17, ENSG00000112511.17, ...
##
     [297]
##
     [298]
##
     [299]
            ENSG00000118322.12, ENSG00000145864.12, ENSG00000253417.5, ...
##
                                                                  FOLLOWID
                                                           <CharacterList>
##
##
       [1]
```

ENSG00000094975.13, ENSG00000117560.7, ENSG00000117586.10, ...

ENSG00000104643.9, ENSG00000154316.15, ENSG00000154319.14, ...

##

##

##

##

##

[2]

[3]

[4]

[5]

. . .

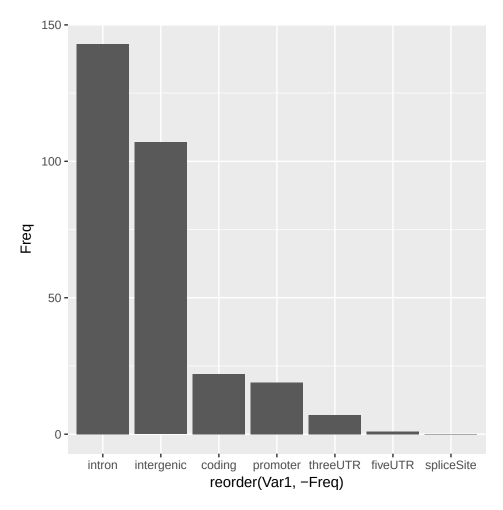


Figure 5. Barplot showing genomic locations associated with SLE variants.

```
##
     [295]
##
     [296] ENSG00000166278.14, ENSG00000168477.17, ENSG00000196126.10, ...
##
     [297]
##
     [298]
     [299] ENSG00000113312.10, ENSG00000135083.14, ENSG00000145861.7, ...
##
##
     seqinfo: 23 sequences from GRCh38 genome; no seqlengths
```

We can visualise where these SNPs are located with ggplot2 44 (Figure 5):

4

5

```
loc <- data.frame(table(snps_anno$LOCATION))</pre>
ggplot(data = loc, aes(x = reorder(Var1, -Freq), y = Freq)) +
 geom_bar(stat="identity")
```

As expected 11, the great majority of SNPs are located within introns and in intergenic regions. For the moment, we will focus on SNPs that are either coding or in promoter and UTR regions, as these can be assigned to target genes rather unambiguously:

```
snps_easy <- subset(snps_anno, LOCATION == "coding" | LOCATION == "promoter" | LOCATION == "threeUTA"</pre>
snps_easy <- as.data.frame(snps_easy)</pre>
head(snps_easy)
##
     seqnames
                    start
                                  end width strand
                                                             SNPS P. VALUE LOCATION
## 1
        chr4 101829919 101829919 1 + rs10516487
                                                                     4e-10
                                                                             coding
## 2
          chr7 128954129 128954129
                                          1
                                                   - rs10488631
                                                                     2e-11 promoter
        chr11 55368743 55368743 1 + rs7927370
chr6 137874929 137874929 1 + rs2230926
chr11 118702810 118702810 1 + rs4639966
## 3
                                                                     7e-06 coding
```

1e-17

coding

1e-16 promoter

```
## 6
        chr16 30624338 30624338
                                               - rs7186852
                                                               3e-07 promoter
##
     LOCSTART LOCEND QUERYID
                                TXID
                                             CDSID
                                                                GENEID PRECEDEID
## 1
                               46105 170258, .... ENSG00000153064.11
          137
                 137
                            7
## 2
                               77786
           NΑ
                  NA
                           23
                                                    ENSG00000275106.1
## 3
          860
                 860
                           45 101610
                                            370677 ENSG00000181958.3
## 4
          380
                 380
                           57 64150 232398, .... ENSG00000118503.14
## 5
           NA
                  NΑ
                           63 104974
                                                    ENSG00000255422.1
## 6
           NA
                   NA
                           68 148763
                                                   ENSG00000156853.12
     FOLLOWID
##
## 1
## 2
## 3
## 4
## 5
## 6
```

Now we can check if any of the genes we found to be differentially expressed in SLE is also genetically associated with the disease:

```
snps_easy_in_degs <- merge(degs, snps_easy, by.x = "gene_id", by.y = "GENEID", all = FALSE)
snps_easy_in_degs</pre>
```

```
## DataFrame with 7 rows and 24 columns
##
                              gene_id bp_length symbol
                                                           baseMean
##
                          <character> <integer>
                                                  t>
                                                          <numeric>
## ENSG00000096968 ENSG00000096968.13
                                            6170
                                                    JAK2 1279.47795
## ENSG00000099834 ENSG00000099834.18
                                            3873
                                                   CDHR5
                                                           10.20177
## ENSG00000115267
                   ENSG00000115267.5
                                            4528
                                                   IFIH1 1415.91330
## ENSG0000120280
                    ENSG00000120280.5
                                            1855 CXorf21 637.78094
## ENSG00000185507 ENSG00000185507.19
                                            2628
                                                    IRF7 4883.20891
## ENSG00000204366
                    ENSG00000204366.3
                                            1875
                                                  ZBTB12
                                                           22.99200
                   ENSG00000275106.1
                                             790
## ENSG00000275106
                                                      NΑ
                                                            10.32171
##
                   log2FoldChange
                                       lfcSE
                                                  stat
                                                             pvalue
##
                        <numeric> <numeric> <numeric>
                                                           <numeric>
## ENSG00000096968
                        0.4854343 \ 0.1553513 \ 3.124753 \ 1.779545e-03
## ENSG00000099834
                        0.8539586 0.2666557
                                              3.202476 1.362516e-03
## ENSG0000115267
                        1.1494945 0.2729847 4.210838 2.544247e-05
## ENSG00000120280
                        0.7819504 0.1541707 5.071977 3.937038e-07
## ENSG0000185507
                        1.4062704 0.2992536 4.699260 2.611057e-06
## ENSG00000204366
                       -0.3892298 0.1348705 -2.885952 3.902318e-03
## ENSG00000275106
                        0.7344844 0.2305300 3.186068 1.442206e-03
##
                           padj seqnames
                                              start
##
                      <numeric> <factor> <integer> <integer> <integer>
## ENSG00000096968 2.068794e-02
                                            4984530
                                                      4984530
                                    chr9
                                                                       1
                                             625085
## ENSG00000099834 1.732902e-02
                                                       625085
                                                                       1
                                    chr11
## ENSG00000115267 1.120363e-03
                                    chr2 162267541 162267541
                                                                       1
## ENSG00000120280 6.047898e-05
                                    chrX 30559729 30559729
                                                                       1
## ENSG00000185507 2.298336e-04
                                    chr11
                                             614318
                                                       614318
                                                                       1
## ENSG00000204366 3.584479e-02
                                     chr6
                                          31902549 31902549
                                                                       1
## ENSG00000275106 1.797861e-02
                                     chr7 128954129 128954129
                                                                       1
##
                                    SNPS
                                          P. VALUE LOCATION LOCSTART
                     strand
##
                   <factor> <character> <numeric> <factor> <integer>
## ENSG00000096968
                          +
                              rs1887428
                                             1e-06 fiveUTR
                                                                  141
## ENSG00000099834
                             rs58688157
                                                                    NΑ
                                             5e-13 promoter
## ENSG0000115267
                                                                  2836
                              rs1990760
                                             4e - 08
                                                     coding
## ENSG00000120280
                                             5e-10
                                                                  627
                               rs887369
                                                     coding
## ENSG0000185507
                                             9e-11
                                                                   217
                              rs1061502
                                                     coding
## ENSG00000204366
                               rs558702
                                             8e-21 promoter
                                                                   ΝA
## ENSG00000275106
                             rs10488631
                                             2e-11 promoter
                                                                    NA
##
                      LOCEND
                                QUERYID
                                               TXID
                                                                        CDSID
##
                   <integer> <integer> <character>
                                                                       <list>
## ENSG00000096968
                         141
                                    329
                                              86536
## ENSG00000099834
                          NΑ
                                    208
                                             105793
                        2836
                                    233
## ENSG00000115267
                                              29219
                                                                       106867
## ENSG00000120280
                                    192
                                                                       692823
                         627
                                             194672
```

```
## ENSG0000185507
                         217
                                   317
                                            105777 385431,385427,385428,...
## ENSG00000204366
                          ΝA
                                   116
                                             65993
## ENSG00000275106
                          NΑ
                                    23
                                             77786
                   PRECEDEID FOLLOWID
##
##
                      t>
                             t>
## ENSG00000096968
## ENSG00000099834
## ENSG00000115267
## ENSG00000120280
## ENSG0000185507
## ENSG00000204366
## ENSG00000275106
```

So, we have 7 genes showing differential expression in SLE that are also genetically associated with the disease. While this is an interesting result, these hits are likely to be already well-known as potential SLE targets given their clear genetic association.

We will store essential information about these hits in a results data.frame:

```
prioritised_hits <- unique(data.frame(
    snp_id = snps_easy_in_degs$SNPS,
    snp_pvalue = snps_easy_in_degs$P.VALUE,
    snp_location = snps_easy_in_degs$LOCATION,
    gene_id = snps_easy_in_degs$gene_id,
    gene_symbol = snps_easy_in_degs$symbol,
    gene_pvalue = snps_easy_in_degs$padj,
    gene_log2foldchange = snps_easy_in_degs$log2FoldChange))
prioritised_hits</pre>
```

```
snp_id snp_pvalue snp_location
                                                                gene_id
## ENSG00000096968 rs1887428
                                  1e-06
                                            fiveUTR ENSG00000096968.13
## ENSG00000099834 rs58688157
                                  5e-13
                                            promoter ENSG00000099834.18
## ENSG00000115267 rs1990760
                                   4e-08
                                            coding ENSG00000115267.5
## ENSG0000120280
                   rs887369
                                  5e-10
                                              coding ENSG00000120280.5
## ENSG00000185507 rs1061502
                                  9e-11
                                              coding ENSG00000185507.19
## ENSG00000204366
                                  8e-21
                                            promoter ENSG00000204366.3
                   rs558702
## ENSG00000275106 rs10488631
                                  2e-11
                                            promoter ENSG00000275106.1
##
                  gene_symbol gene_pvalue gene_log2foldchange
## ENSG0000096968
                         JAK2 2.068794e-02
                                                     0.4854343
## ENSG00000099834
                        CDHR5 1.732902e-02
                                                     0.8539586
## ENSG00000115267
                        IFIH1 1.120363e-03
                                                     1.1494945
## ENSG0000120280
                      CXorf21 6.047898e-05
                                                     0.7819504
## ENSG0000185507
                         IRF7 2.298336e-04
                                                     1.4062704
## ENSG00000204366
                        ZBTB12 3.584479e-02
                                                     -0.3892298
## ENSG00000275106
                           NA 1.797861e-02
                                                     0.7344844
```

Use of regulatory genomic data to map intronic and intergenic SNPs to target genes

But what about all the SNPs in introns and intergenic regions? Some of those might be regulatory SNPs affecting the expression level of their target gene(s) through a distal enhancer. Let's create a dataset of candidate regulatory SNPs that are either intronic or intergenic and remove the annotation obtained with VariantAnnotation:

```
snps_hard <- subset(snps_anno, LOCATION == "intron" | LOCATION == "intergenic", select = c("SNPS", snps_hard</pre>
```

```
## GRanges object with 250 ranges and 3 metadata columns:
##
           seqnames
                                    ranges strand
                                                           SNPS
                                                                  P.VALUE
##
              <Rle>
                                 <IRanges> <Rle> | <character> <numeric>
##
       [1]
              chr16 [ 31301932,
                                 31301932]
                                                +
                                                      rs9888739
                                                                    2e-23
##
              chr11 [ 589564,
       Γ2]
                                   589564]
                                                +
                                                      rs4963128
                                                                    3e-10
              chr3 [ 58384450, 58384450]
                                                                    7e-09
##
       [3]
                                                + |
                                                      rs6445975
                                                                    1e-07
##
       Γ47
               chr1 [173340574, 173340574]
                                                *
                                                     rs10798269
##
       [5]
               chr8 [ 11491677, 11491677]
                                                     rs13277113
                                                                    1e-10
```

```
##
                . . .
                                               . . . .
     [246]
               chr6 [137874014, 137874014]
##
                                                 +
                                                        rs5029937
                                                                      5e-13
##
               chr6 [ 32619077, 32619077]
                                                 *
                                                                      1e-07
     [247]
                                                        rs9271366
               chr6 [137685367, 137685367]
##
                                                                      4e-07
     Γ248]
                                                 +
                                                        rs6920220
               chrX [153924366, 153924366]
                                                                      8e-07
##
     Γ2491
                                                 - |
                                                        rs2269368
##
     [250]
               chr5 [160459613, 160459613]
                                                 *
                                                        rs2431099
                                                                      2e-06
##
             LOCATION
##
             <factor>
##
       [1]
               intron
##
       [2]
               intron
##
       [3]
               intron
##
       [4] intergenic
##
       [5] intergenic
##
     Γ246]
##
               intron
##
     [247] intergenic
##
     [248]
              intron
##
     [249]
               intron
##
     [250] intergenic
##
     seqinfo: 23 sequences from GRCh38 genome; no seqlengths
##
```

eQTL data

A well-established way to gain insights into target genes of regulatory SNPs is to use eQTL data, where correlations between genetic variants and expression of genes are computed across different tissues or cell types ¹³. We will use blood eQTL data from the GTEx consortium ¹⁴. To get the data, you will have to register and download the file GTEx_Analysis_v7_eQTL.tar.gz from the GTEx portal website ⁵³ to the current working directory:

```
# uncomment the following line to extract the gzipped archive file
#untar("GTEx_Analysis_v7_eQTL.tar.gz")
gtex_blood <- read.delim(gzfile("GTEx_Analysis_v7_eQTL/Whole_Blood.v7.signif_variant_gene_pairs.txt
head(gtex_blood)</pre>
```

```
##
                                 gene_id tss_distance ma_samples ma_count
            variant_id
## 1 1_231153_CTT_C_b37 ENSG00000223972.4
                                              219284
                                                             13
       1_61920_G_A_b37 ENSG00000238009.2
                                              -67303
                                                             18
                                                                      20
       1_64649_A_C_b37 ENSG00000238009.2
## 3
                                              -64574
                                                             16
                                                                      16
      1_115746_C_T_b37 ENSG00000238009.2
## 4
                                               -13477
                                                             45
                                                                      45
## 5
      1_135203_G_A_b37 ENSG00000238009.2
                                                 5980
                                                             51
                                                                      51
## 6
      1_988016_T_C_b37 ENSG00000268903.1
                                               852121
                                                             21
                                                                      23
##
          maf pval_nominal
                             slope slope_se pval_nominal_threshold
## 1 0.0191740 3.69025e-08 1.319720 0.233538
                                                       1.35366e-04
## 2 0.0281690 7.00836e-07 0.903786 0.178322
                                                       8.26088e-05
## 3 0.0220386 5.72066e-07 1.110040 0.217225
                                                       8.26088e-05
## 4 0.0628492 6.50297e-10 0.858203 0.134436
                                                       8.26088e-05
## 5 0.0698630 6.67194e-10 0.811790 0.127255
                                                       8.26088e-05
## 6 0.0318560 6.35694e-05 0.501916 0.123743
                                                       8.52870e-05
    min_pval_nominal pval_beta
         3.69025e-08 4.67848e-05
## 1
## 2
         6.50297e-10 1.11312e-06
## 3
         6.50297e-10 1.11312e-06
## 4
         6.50297e-10 1.11312e-06
         6.50297e-10 1.11312e-06
## 5
         6.35694e-05 5.44487e-02
```

We have to extract the genomic locations of the SNPs from the IDs used by GTEx:

```
locs <- strsplit(gtex_blood$variant_id, "_")
gtex_blood$chr <- sapply(locs, "[", 1)
gtex_blood$start <- sapply(locs, "[", 2)
gtex_blood$end <- sapply(locs, "[", 2)
tail(gtex_blood)</pre>
```

```
##
                     variant_id
                                          gene_id tss_distance ma_samples
## 1052537 X_154999134_G_A_b37 ENSG00000168939.6
                                                          1660
## 1052538 X_154999204_TA_T_b37 ENSG00000168939.6
                                                                      219
                                                          1730
## 1052539 X_155004280_A_G_b37 ENSG00000168939.6
                                                          6806
                                                                      186
## 1052540 X_155011926_T_C_b37 ENSG00000168939.6
                                                         14452
                                                                      222
## 1052541 X_155014420_A_G_b37 ENSG00000168939.6
                                                         16946
                                                                      215
## 1052542 X_155186978_G_C_b37 ENSG00000168939.6
                                                        189504
                                                                      250
          ma_count
                        maf pval_nominal
                                              slope slope_se
## 1052537
                259 0.351902 3.19266e-05 -0.162062 0.0383749
## 1052538
                274 0.390313 6.72752e-05 -0.157810 0.0390413
                224 0.303523 1.91420e-08 0.230301 0.0398809
## 1052539
## 1052540
                279 0.379076 3.88977e-05 0.157608 0.0377434
                265 0.360054 4.17781e-05 0.159699 0.0384025
## 1052541
## 1052542
                321 0.436141 1.24355e-04 0.145560 0.0374390
##
           pval_nominal_threshold min_pval_nominal
                                                    pval_beta chr
                                                                       start
## 1052537
                     0.000130368
                                        1.9142e-08 2.75084e-05
                                                                 X 154999134
## 1052538
                      0.000130368
                                        1.9142e-08 2.75084e-05
                                                                 X 154999204
## 1052539
                      0.000130368
                                        1.9142e-08 2.75084e-05
                                                                 X 155004280
## 1052540
                      0.000130368
                                        1.9142e-08 2.75084e-05
                                                                 X 155011926
                                        1.9142e-08 2.75084e-05
## 1052541
                      0.000130368
                                                                 X 155014420
                                        1.9142e-08 2.75084e-05
                                                                 X 155186978
## 1052542
                      0.000130368
##
                 end
## 1052537 154999134
## 1052538 154999204
## 1052539 155004280
## 1052540 155011926
## 1052541 155014420
## 1052542 155186978
```

We can then convert the data.frame into a GRanges object:

```
gtex_blood <- makeGRangesFromDataFrame(gtex_blood, keep.extra.columns = TRUE)
gtex_blood</pre>
```

```
## GRanges object with 1052542 ranges and 12 metadata columns:
##
               seqnames
                                         ranges strand
                                                                    variant_id
##
                  <R.1e>
                                      <IRanges> <Rle> |
                                                                   <character>
##
                                                           1_231153_CTT_C_b37
           [1]
                      1
                               [231153, 231153]
                                                    *
##
           [2]
                              [ 61920, 61920]
                                                     *
                                                              1_61920_G_A_b37
                      1
           [3]
##
                      1
                              [ 64649, 64649]
                                                     *
                                                              1_64649_A_C_b37
##
           [4]
                               [115746, 115746]
                                                     *
                                                             1_115746_C_T_b37
                      1
##
           [5]
                      1
                               [135203, 135203]
                                                     *
                                                             1_135203_G_A_b37
##
##
     [1052538]
                      X [154999204, 154999204]
                                                    * | X_154999204_TA_T_b37
##
     [1052539]
                      X [155004280, 155004280]
                                                     * | X_155004280_A_G_b37
                                                     * | X_155011926_T_C_b37
##
     [1052540]
                      X [155011926, 155011926]
##
     Γ10525417
                      X [155014420, 155014420]
                                                     * | X_155014420_A_G_b37
                      X [155186978, 155186978]
##
     [1052542]
                                                     * | X_155186978_G_C_b37
##
                         gene_id tss_distance ma_samples ma_count
##
                                     <integer> <integer> <integer> <numeric>
                     <character>
           [1] ENSG00000223972.4
##
                                        219284
                                                       13
                                                                 13 0.0191740
           [2] ENSG00000238009.2
                                        -67303
                                                                 20 0.0281690
##
                                                       18
           [3] ENSG00000238009.2
                                        -64574
                                                                 16 0.0220386
##
                                                       16
##
           [4] ENSG00000238009.2
                                        - 13477
                                                       45
                                                                 45 0.0628492
           [5] ENSG00000238009.2
                                                                 51 0.0698630
##
                                          5980
                                                       51
##
                                           . . .
                                                      . . .
##
     [1052538] ENSG00000168939.6
                                          1730
                                                      219
                                                                 274
                                                                     0.390313
##
     [1052539] ENSG00000168939.6
                                          6806
                                                      186
                                                                 224 0.303523
##
     [1052540] ENSG00000168939.6
                                         14452
                                                      222
                                                                 279 0.379076
##
     [1052541] ENSG00000168939.6
                                         16946
                                                      215
                                                                 265 0.360054
##
                                        189504
                                                      250
     [1052542] ENSG00000168939.6
                                                                 321 0.436141
##
               pval_nominal
                                        slope_se pval_nominal_threshold
                                slope
##
                                                              <numeric>
                  <numeric> <numeric> <numeric>
##
                                                            1.35366e-04
           Γ1 ]
                3.69025e-08 1.319720
                                       0.233538
##
           [2] 7.00836e-07 0.903786 0.178322
                                                            8.26088e-05
```

```
[3] 5.72066e-07 1.110040 0.217225
[4] 6.50297e-10 0.858203 0.134436
##
                                                             8.26088e-05
##
                                                             8.26088e-05
           [5] 6.67194e-10 0.811790 0.127255
                                                             8.26088e-05
##
##
     [1052538] 6.72752e-05 -0.157810 0.0390413
                                                             0.000130368
##
##
     [1052539] 1.91420e-08 0.230301 0.0398809
                                                            0.000130368
##
     [1052540] 3.88977e-05 0.157608 0.0377434
                                                            0.000130368
     [1052541] 4.17781e-05 0.159699 0.0384025
##
                                                            0.000130368
     [1052542] 1.24355e-04 0.145560 0.0374390
                                                             0.000130368
##
##
               min_pval_nominal pval_beta
##
                       <numeric>
                                   <numeric>
                    3.69025e-08 4.67848e-05
##
           [1]
                    6.50297e-10 1.11312e-06
##
           [2]
           [3]
                    6.50297e-10 1.11312e-06
##
                    6.50297e-10 1.11312e-06
##
           [4]
##
           [5]
                    6.50297e-10 1.11312e-06
##
           . . .
                             . . .
##
     [1052538]
                     1.9142e-08 2.75084e-05
##
     [1052539]
                     1.9142e-08 2.75084e-05
                     1.9142e-08 2.75084e-05
##
     [1052540]
                     1.9142e-08 2.75084e-05
##
     [1052541]
                     1.9142e-08 2.75084e-05
##
     [1052542]
##
##
     seqinfo: 23 sequences from an unspecified genome; no seqlengths
```

We also need to ensure that the chromosome notation is consistent with the previous objects:

```
seqlevelsStyle(gtex_blood)

## [1] "NCBI" "Ensembl"

seqlevels(gtex_blood)

## [1] "1" "2" "3" "4" "5" "6" "7" "8" "9" "10" "11" "12" "13" "14"

## [15] "15" "16" "17" "18" "19" "20" "21" "22" "X"

seqlevelsStyle(gtex_blood) <- "UCSC"
seqlevels(gtex_blood)

## [1] "chr1" "chr2" "chr3" "chr4" "chr5" "chr6" "chr7" "chr8"

## [9] "chr9" "chr10" "chr11" "chr12" "chr13" "chr14" "chr15" "chr16"

## [17] "chr17" "chr18" "chr19" "chr20" "chr21" "chr22" "chrX"</pre>
```

From the publication ¹⁴, we know the genomic coordinates are mapped to genome reference GRCh37, so we will have to uplift them to GRCh38 using rtracklayer ⁵⁴ and a mapping ("chain") file. The R.utils package ⁵⁵ is required to extract the gzipped file:

```
library(rtracklayer)
library(R.utils)
# uncomment the following line to download file
#download.file("http://hgdownload.cse.ucsc.edu/goldenPath/hg19/liftOver/hg19ToHg38.over.chain.gz", of
# uncomment the following line to extract gzipped file
#gunzip("hg19ToHg38.over.chain.gz")
ch <- import.chain("hg19ToHg38.over.chain")
gtex_blood <- unlist(liftOver(gtex_blood, ch))</pre>
```

We will use the GenomicRanges package⁵¹ to compute the overlap between GWAS SNPs and blood eQTLs:

```
library(GenomicRanges)
hits <- findOverlaps(snps_hard, gtex_blood)
snps_hard_in_gtex_blood = snps_hard[queryHits(hits)]</pre>
```

```
gtex_blood_with_snps_hard = gtex_blood[subjectHits(hits)]
mcols(snps_hard_in_gtex_blood) <- cbind(mcols(snps_hard_in_gtex_blood), mcols(gtex_blood_with_snps_l
snps_hard_in_gtex_blood <- as.data.frame(snps_hard_in_gtex_blood)
head(snps_hard_in_gtex_blood)</pre>
```

```
##
     seqnames
                            end width strand
                                                   SNPS P.VALUE
                                                                  LOCATION
                 start
                                           + rs4963128
## 1
                589564
                         589564
                                                          3e-10
        chr11
                                   1
                                                                    intron
## 2
         chr3 58384450 58384450
                                    1
                                           + rs6445975
                                                          7e-09
                                                                    intron
## 3
         chr8 11491677 11491677
                                    1
                                           * rs13277113
                                                          1e-10 intergenic
## 4
         chr8 11491677 11491677
                                    1
                                           * rs13277113
                                                          1e-10 intergenic
## 5
         chr8 11491677 11491677
                                    1
                                           * rs13277113
                                                          1e-10 intergenic
## 6
         chr8 11491677 11491677
                                           * rs13277113
                                                          1e-10 intergenic
                                    1
##
             variant_id
                                   gene_id tss_distance ma_samples ma_count
## 1 11_589564_T_C_b37 ENSG00000177042.10
                                                                         250
                                                -105969
                                                               212
## 2 3_58370177_G_T_b37 ENSG00000168291.8
                                                 -49407
                                                               205
                                                                         250
## 3 8_11349186_G_A_b37 ENSG00000154319.10
                                                  16962
                                                               157
                                                                         180
## 4 8_11349186_G_A_b37 ENSG00000136573.8
                                                  -2324
                                                               157
                                                                         180
## 5 8_11349186_G_A_b37 ENSG00000255518.1
                                                 -66284
                                                               157
                                                                         180
## 6 8_11349186_G_A_b37 ENSG00000255354.1
                                                 -68343
                                                               157
                                                                         180
         maf pval_nominal
                               slope slope_se pval_nominal_threshold
## 1 0.339674 4.51059e-10 -0.194589 0.0301828
                                                          3.35947e-05
## 2 0.338753 2.05231e-12 0.179408 0.0244587
                                                          6.23219e-05
## 3 0.243902 6.46308e-27 0.778785 0.0656311
                                                          3.79430e-05
## 4 0.243902 5.04687e-18 -0.281643 0.0305280
                                                          3.75653e-05
## 5 0.243902 7.37464e-07 -0.262302 0.0518614
                                                          3.41126e-05
## 6 0.243902 8.41301e-08 -0.243121 0.0442629
                                                          3.66297e-05
                      pval_beta
    min_pval_nominal
## 1
         5.23982e-30 1.63019e-24
## 2
         3.39499e-13 3.97374e-09
## 3
         8.46904e-29 2.22416e-23
## 4
         2.97871e-19 2.22082e-14
## 5
         8.28459e-08 4.81268e-04
## 6
         2.67616e-08 1.37119e-04
```

So, we have 59 blood eQTL variants that are associated with SLE. We can now check whether any of the genes differentially expressed in SLE is an *eGene*, a gene whose expression is influenced by an eQTL. We note that gene IDs in GTEx are mapped to GENCODE v19 ¹⁴, while we are using the newer v25 for the DEGs. To match the gene IDs in the two objects, we will simply strip the last bit containing the GENCODE gene version, which effectively gives us Ensembl gene IDs:

```
snps\_hard\_in\_gtex\_blood\$ensembl\_id <- sub("(ENSG[0-9]+)\\.[0-9]+", "\\1", snps\_hard\_in\_gtex\_blood\$gdegs\$ensembl\_id <- sub("(ENSG[0-9]+)\\.[0-9]+", "\\1", degs\$gene\_id) \\ snps\_hard\_in\_gtex\_blood\_in\_degs <- merge(snps\_hard\_in\_gtex\_blood, degs, by = "ensembl\_id", all = FAIsnps\_hard\_in\_gtex\_blood\_in\_degs
```

```
## DataFrame with 6 rows and 30 columns
##
         ensembl_id seqnames
                                  start
                                              end
                                                      width
                                                              strand
##
         <character> <factor> <integer> <integer> <integer> <factor>
## 1 ENSG00000130513
                        chr19
                              18370523
                                        18370523
                                                          1
## 2 ENSG00000140497
                              75018695
                                         75018695
                        chr15
                                                          1
## 3 ENSG00000172890
                        chr11
                              71476633
                                        71476633
                                                          1
## 4 ENSG00000214894
                         chr6
                              31668965
                                        31668965
                                                          1
## 5 ENSG00000214894
                         chr6
                              30973212 30973212
                                                          1
## 6 ENSG00000214894
                         chr6 31753256 31753256
                                                          1
##
           SNPS
                  P.VALUE
                            LOCATION
                                               variant_id
                                                                   gene_id.x
##
     <character> <numeric>
                             <factor>
                                                                 <character>
                                              <character>
## 1
      rs8105429
                     5e-06 intergenic 19_18481333_A_G_b37 ENSG00000130513.6
      rs2289583
                     6e-15
                               intron 15_75311036_C_A_b37 ENSG00000140497.12
## 3
      rs3794060
                               intron 11_71187679_C_T_b37 ENSG00000172890.7
                     1e-20
## 4
      rs9267531
                     8e-08
                               intron 6_31636742_A_G_b37 ENSG00000214894.2
## 5 rs114090659
                     6e-92 intergenic 6_30940989_T_C_b37 ENSG00000214894.2
## 6
      rs3131379
                     2e-52
                               intron 6_31721033_G_A_b37 ENSG00000214894.2
    tss_distance ma_samples ma_count
                                             maf pval_nominal
                                                                  slope
```

```
##
       <integer>
                  <integer> <integer> <numeric>
                                                    <numeric> <numeric>
## 1
                                  189 0.2560980 7.87256e-11 0.350964
           -4208
                        166
## 2
                                   191 0.2588080 7.57250e-06 -0.107460
           145330
                         170
## 3
                        183
                                   231 0.3130080 1.91380e-31 0.407266
           23524
## 4
           838306
                                   54 0.0731707 3.36144e-08 0.479659
                         49
## 5
           142553
                          83
                                   91 0.1233060 7.00411e-11 0.453255
## 6
          922597
                          50
                                   55 0.0745257 2.69451e-08 0.479935
##
     slope_se pval_nominal_threshold min_pval_nominal
                                                        pval_beta
##
     <numeric>
                           <numeric>
                                             <numeric>
                                                         <numeric>
## 1 0.0520458
                          2.52102e-05
                                           1.76820e-11 1.23175e-07
## 2 0.0235858
                          6.38531e-05
                                          2.44784e-27 1.10743e-22
## 3 0.0310305
                                          1.05596e-33 7.87659e-28
                         4.46719e-05
## 4 0.0846154
                          6.02220e-05
                                           3.17673e-13 1.77790e-08
## 5 0.0670210
                          6.02220e-05
                                           3.17673e-13 1.77790e-08
## 6 0.0840440
                          6.02220e-05
                                          3.17673e-13 1.77790e-08
             gene_id.y bp_length symbol
                                             baseMean log2FoldChange
            <character> <integer>
                                     t>
                                             <numeric>
                                                            <numeric>
## 1 ENSG00000130513.6
                            2087
                                     GDF15
                                               6.75448
                                                            0.7883703
## 2 ENSG00000140497.16
                            5000
                                     SCAMP2 3483.03109
                                                           -0.2959934
## 3 ENSG00000172890.11
                                   NADSYN1 4020.56224
                           16263
                                                            0.2619770
## 4 ENSG00000214894.6
                            2171 LINC00243
                                             74.95034
                                                            1.2684089
                            2171 LINC00243
## 5
     ENSG00000214894.6
                                              74.95034
                                                            1.2684089
## 6 ENSG00000214894.6
                            2171 LINC00243
                                              74.95034
                                                            1.2684089
##
         1fcSE
                  stat
                               pvalue
                                               padj
      <numeric> <numeric>
                            <numeric>
                                          <numeric>
## 1 0.28347645 2.781079 5.417861e-03 0.0448154406
## 2 0.08814542 -3.358012 7.850510e-04 0.0119267855
## 3 0.08976429 2.918499 3.517209e-03 0.0333810138
## 4 0.27106143 4.679415 2.876950e-06 0.0002442643
## 5 0.27106143 4.679415 2.876950e-06 0.0002442643
## 6 0.27106143 4.679415 2.876950e-06 0.0002442643
```

We can add these 4 genes to our list:

```
prioritised_hits <- unique(rbind(prioritised_hits, data.frame(
    snp_id = snps_hard_in_gtex_blood_in_degs$SNPS,
    snp_pvalue = snps_hard_in_gtex_blood_in_degs$P.VALUE,
    snp_location = snps_hard_in_gtex_blood_in_degs$LOCATION,
    gene_id = snps_hard_in_gtex_blood_in_degs$gene_id.y,
    gene_symbol = snps_hard_in_gtex_blood_in_degs$symbol,
    gene_pvalue = snps_hard_in_gtex_blood_in_degs$padj,
    gene_log2foldchange = snps_hard_in_gtex_blood_in_degs$log2FoldChange)))
prioritised_hits</pre>
```

```
snp_id snp_pvalue snp_location
##
                                                                   gene_id
## ENSG00000096968
                                    1e-06
                                               fiveUTR ENSG00000096968.13
                     rs1887428
## ENSG00000099834
                   rs58688157
                                    5e-13
                                              promoter ENSG00000099834.18
## ENSG0000115267
                     rs1990760
                                    4e-08
                                                coding ENSG00000115267.5
## ENSG00000120280
                                    5e-10
                                                coding ENSG00000120280.5
                      rs887369
## ENSG0000185507
                     rs1061502
                                    9e-11
                                                coding ENSG00000185507.19
## ENSG00000204366
                      rs558702
                                    8e-21
                                              promoter ENSG00000204366.3
                                              promoter ENSG00000275106.1
## ENSG00000275106
                    rs10488631
                                    2e-11
                                            intergenic ENSG00000130513.6
## 1
                     rs8105429
                                    5e-06
## 2
                                    6e-15
                                                intron ENSG00000140497.16
                     rs2289583
## 3
                     rs3794060
                                    1e-20
                                                intron ENSG00000172890.11
## 4
                                    8e-08
                                                intron ENSG00000214894.6
                     rs9267531
## 5
                   rs114090659
                                    6e-92
                                            intergenic ENSG00000214894.6
## 6
                     rs3131379
                                    2e-52
                                                intron ENSG00000214894.6
##
                   gene_symbol gene_pvalue gene_log2foldchange
## ENSG00000096968
                          JAK2 2.068794e-02
                                                      0.4854343
## ENSG00000099834
                         CDHR5 1.732902e-02
                                                      0.8539586
## ENSG00000115267
                         IFIH1 1.120363e-03
                                                      1.1494945
## ENSG00000120280
                       CXorf21 6.047898e-05
                                                      0.7819504
## ENSG0000185507
                          IRF7 2.298336e-04
                                                      1.4062704
## ENSG00000204366
                       ZBTB12 3.584479e-02
                                                      -0.3892298
```

##	ENSG00000275106	NA	1.797861e-02	0.7344844
##	1	GDF15	4.481544e-02	0.7883703
##	2	SCAMP2	1.192679e-02	-0.2959934
##	3	NADSYN1	3.338101e-02	0.2619770
##	4	LINC00243	2.442643e-04	1.2684089
##	5	LINC00243	2.442643e-04	1.2684089
##	6	LINCO0243	2.442643e-04	1.2684089

FANTOM5 data

The FANTOM consortium profiled gene expression across a large panel of tissues and cell types using CAGE ^{19;21}. This technology allows mapping of transcription start sites (TSSs) and enhancer RNAs (eRNAs) genome-wide. Correlations between these promoter and enhancer elements across a large panel of tissues and cell types can then be calculated to identify significant promoter - enhancer pairs. In turn, we will use these correlations to map distal regulatory SNPs to target genes.

We can read in and have a look at the enhancer - promoter correlation data in this way:

```
# uncomment the following line to download the file
#download.file("http://enhancer.binf.ku.dk/presets/enhancer_tss_associations.bed", destfile = "enhancer_tss_associations.bed", skip = 1, stringsAsFactors = FALSE)
head(fantom)
```

```
##
    X.chrom chromStart chromEnd
## 1
                         861621
       chr1
                858252
## 2
        chr1
                894178
                         956888
## 3
       chr1
                901376
                         956888
## 4
        chr1
                901376 1173762
## 5
       chr1
                935051
                         942164
## 6
                935051 1005621
       chr1
##
## 1
                                     chr1:858256-858648; NM_152486; SAMD11; R:0.404; FDR:0
## 2
                   chr1:956563-956812; NM_015658; NOC2L; R:0.202; FDR:8.01154668254404e-08
                       chr1:956563-956812; NM_001160184, NM_032129; PLEKHN1; R:0.422; FDR:0
## 3
## 4
                     chr1:1173386-1173736; NM_001160184, NM_032129; PLEKHN1; R:0.311; FDR:0
## 5
       chr1:941791-942135; NM_001142467, NM_021170; HES4; R:0.187; FDR:6.32949888009368e-07
## 6 chr1:1005293-1005547;NM_001142467,NM_021170;HES4;R:0.236;FDR:6.28221217150423e-11
##
    score strand thickStart thickEnd itemRgb blockCount blockSizes
                              858453 0,0,0 2
## 1
      404
              .
                     858452
                                                          401,1001
                                     0,0,0
      202
## 2
                     956687
                              956688
                                                      2
                                                           1001,401
                                                     2
## 3
       422
                     956687
                              956688 0,0,0
                                                           1001,401
                                                     2
## 4
       311
                    1173561 1173562 0,0,0
                                                           1001,401
## 5
       187
                     941963
                              941964
                                       0,0,0
                                                      2
                                                           1001,401
## 6
       236
                     1005420 1005421
                                        0,0,0
                                                           1001,401
##
    chromStarts
## 1
         0,2368
## 2
        0,62309
## 3
        0,55111
## 4
       0,271985
## 5
         0,6712
## 6
        0,70169
```

Everything we need is in the fourth column, name: genomic location of the enhancer, gene identifiers, Pearson correlation coefficient and significance. We will use the splitstackshape package 56 to parse it:

```
library(splitstackshape)
fantom <- as.data.frame(cSplit(fantom, splitCols = "name", sep = ";", direction = "wide"))</pre>
head(fantom)
##
     X.chrom chromStart chromEnd score strand thickStart thickEnd itemRgb
## 1
                                                                      0,0,0
        chr1
                 858252
                          861621
                                    404
                                                   858452
                                                            858453
                                            .
## 2
                 894178
                          956888
                                    202
                                                   956687
                                                            956688
                                                                      0,0,0
        chr1
## 3
        chr1
                 901376
                          956888
                                    422
                                                   956687
                                                            956688
                                                                      0,0,0
## 4
                 901376 1173762
        chr1
                                    311
                                                  1173561 1173562
                                                                      0,0,0
```

```
## 5
       chr1
                935051
                        942164
                                  187
                                                 941963
                                                          941964
                                                                   0,0,0
                935051 1005621
## 6
       chr1
                                  236
                                                1005420 1005421
                                                                   0,0,0
##
    blockCount blockSizes chromStarts
                                                    name_1
                                        chr1:858256-858648
## 1
             2
                 401,1001
                            0,2368
## 2
             2
                 1001,401
                             0,62309
                                       chr1:956563-956812
                                       chr1:956563-956812
## 3
             2
                 1001,401
                            0,55111
## 4
             2
                 1001,401
                             0,271985 chr1:1173386-1173736
## 5
             2
                 1001,401
                              0,6712
                                        chr1:941791-942135
## 6
                              0,70169 chr1:1005293-1005547
                 1001,401
##
                    name_2 name_3 name_4
                                                             name 5
## 1
                 NM 152486 SAMD11 R:0.404
                                                              FDR:0
                            NOC2L R:0.202 FDR:8.01154668254404e-08
## 2
                 NM_015658
## 3 NM_001160184, NM_032129 PLEKHN1 R:0.422
                                                              FDR:0
                                                              FDR:0
## 4 NM_001160184, NM_032129 PLEKHN1 R:0.311
## 5 NM_001142467, NM_021170
                              HES4 R:0.187 FDR:6.32949888009368e-07
## 6 NM_001142467, NM_021170
                              HES4 R:0.236 FDR:6.28221217150423e-11
```

Now we can extract the genomic locations of the enhancers and the correlation values:

```
locs <- strsplit(as.character(fantom$name_1), "[:-]")</pre>
fantom$chr <- sapply(locs, "[", 1)</pre>
fantom$start <- as.numeric(sapply(locs, "[", 2))</pre>
fantom$end <- as.numeric(sapply(locs, "[", 3))</pre>
fantom$symbol <- fantom$name_3</pre>
fantom$corr <- sub("R:", "", fantom$name_4)</pre>
fantom$fdr <- sub("FDR:", "", fantom$name_5)
head(fantom)
```

```
X.chrom chromStart chromEnd score strand thickStart thickEnd itemRgb
                858252 861621 404 .
## 1
       chr1
                                                858452
                                                         858453
                                                                 0,0,0
## 2
       chr1
                894178
                        956888
                                 202
                                                956687
                                                         956688
                                                                 0,0,0
## 3
       chr1
                901376
                        956888
                                 422
                                                956687
                                                         956688
                                                                 0,0,0
## 4
                901376 1173762
                                 311
       chr1
                                               1173561 1173562
                                                                 0,0,0
## 5
                935051
                        942164
                                 187
                                                941963
                                                         941964
       chr1
                                                                 0,0,0
                935051 1005621
## 6
       chr1
                                 236
                                               1005420 1005421
                                                                 0,0,0
##
    blockCount blockSizes chromStarts
## 1
             2 401,1001
                          0,2368 chr1:858256-858648
## 2
             2
                1001,401
                             0,62309
                                      chr1:956563-956812
                                       chr1:956563-956812
## 3
             2 1001,401
                            0,55111
## 4
            2 1001,401
                            0,271985 chr1:1173386-1173736
## 5
            2 1001,401
                             0,6712
                                      chr1:941791-942135
## 6
            2
                1001,401
                             0,70169 chr1:1005293-1005547
##
                    name_2 name_3 name_4
                                                            name_5 chr
                 NM_152486 SAMD11 R:0.404
## 1
                                                             FDR:0 chr1
## 2
                            NOC2L R:0.202 FDR:8.01154668254404e-08 chr1
                 NM_015658
## 3 NM_001160184, NM_032129 PLEKHN1 R:0.422
                                                             FDR:0 chr1
## 4 NM_001160184, NM_032129 PLEKHN1 R:0.311
                                                             FDR:0 chr1
## 5 NM_001142467,NM_021170
                             HES4 R:0.187 FDR:6.32949888009368e-07 chr1
## 6 NM_001142467, NM_021170
                             HES4 R:0.236 FDR:6.28221217150423e-11 chr1
      start
                end symbol corr
                                                  fdr
     858256 858648 SAMD11 0.404
## 2 956563 956812
                     NOC2L 0.202 8.01154668254404e-08
## 3 956563 956812 PLEKHN1 0.422
                                                    0
## 4 1173386 1173736 PLEKHN1 0.311
                                                    0
## 5 941791 942135
                      HES4 0.187 6.32949888009368e-07
## 6 1005293 1005547
                      HES4 0.236 6.28221217150423e-11
```

We can select only the enhancer - promoter pairs with a decent level of correlation and significance and tidy the data at the same time:

end symbol

chr

start

```
fantom <- unique(subset(fantom, subset = corr >= 0.25 & fdr < 1e-5, select = c("chr", "start", "end
head(fantom)
##
```

```
## 1 chr1 858256 858648 SAMD11
## 3 chr1 956563 956812 PLEKHN1
## 4 chr1 1173386 1173736 PLEKHN1
## 13 chr1 1136075 1136463 ISG15
## 14 chr1 956563 956812 AGRN
## 27 chr1 1060905 1061095 RNF223
```

Now we would like to check whether any of our candidate regulatory SNPs are falling in any of these enhancers. To do this, we have to convert the data.frame into a GRanges object:

```
fantom <- makeGRangesFromDataFrame(fantom, keep.extra.columns = TRUE)
fantom</pre>
```

```
## GRanges object with 33957 ranges and 1 metadata column:
          seqnames
                                   ranges strand
                                                     symbol
##
              <Rle>
                                <IRanges> <Rle> | <factor>
##
              chr1
                       [ 858256, 858648]
                                                     SAMD11
                                 956812]
##
        3
              chr1
                        [ 956563,
                                               *
                                                    PLEKHN1
                       [1173386, 1173736]
                                                    PLEKHN1
##
        4
              chr1
                                               *
                       [1136075, 1136463]
                                                      ISG15
##
       13
              chr1
                                               *
##
                       [ 956563, 956812]
                                               *
       14
                                                      AGRN
              chr1
##
              chrX [154256125, 154256514]
##
    66929
                                                       F8A2
              chrY [ 2871660, 2871926]
##
    66932
                                               *
                                                        ZFY
##
    66933
              chrY [ 2872046,
                                 2872325]
                                               *
                                                        ZFY
##
    66940
              chrY [ 21664138, 21664302]
                                                      KDM5D
##
    66941
              chrY [ 22735456, 22735677]
                                                     EIF1AY
##
##
     seqinfo: 24 sequences from an unspecified genome; no seqlengths
```

Similar to the GTEx data, the FANTOM5 data is also mapped to GRCh37¹⁹, so we will have to uplift the GRCh37 coordinates to GRCh38:

```
fantom <- unlist(liftOver(fantom, ch))
fantom</pre>
```

```
## GRanges object with 34160 ranges and 1 metadata column:
##
           seqnames
                                    ranges strand
                                                      symbol
##
             <Rle>
                                 <IRanges> <Rle> | <factor>
##
                        [ 922876, 923268]
              chr1
                                                *
                                                      SAMD11
##
        3
                        [1021183, 1021432]
                                                     PLEKHN1
              chr1
##
        4
              chr1
                        [1238006, 1238356]
                                                *
                                                     PLEKHN1
##
       13
                        [1200695, 1201083]
              chr1
                                                       ISG15
##
       14
              chr1
                        [1021183, 1021432]
                                                        AGRN
                                                *
##
       . . .
               . . .
                                                         . . .
##
    66929
              chrX [155027850, 155028239]
                                                        F8A2
##
    66932
              chrY [ 3003619, 3003885]
                                                *
                                                         ZFY
##
    66933
              chrY [ 3004005,
                                  3004284]
                                                         ZFY
    66940
               chrY [ 19502252,
                                19502416]
                                                       KDM5D
##
              chrY [ 20573570, 20573791]
##
    66941
                                                      EIF1AY
##
     seqinfo: 24 sequences from an unspecified genome; no seqlengths
##
```

We can now compute the overlap between SNPs and enhancers:

```
hits <- findOverlaps(snps_hard, fantom)
snps_hard_in_fantom = snps_hard[queryHits(hits)]
fantom_with_snps_hard = fantom[subjectHits(hits)]
mcols(snps_hard_in_fantom) <- cbind(mcols(snps_hard_in_fantom), mcols(fantom_with_snps_hard))
snps_hard_in_fantom <- as.data.frame(snps_hard_in_fantom)
snps_hard_in_fantom</pre>
```

```
##
      seqnames
                               end width strand
                                                      SNPS P.VALUE
                                                                     LOCATION
                  start
## 1
          chr2 191099907 191099907
                                     1
                                                rs7574865
                                                             9e-14
                                                                       intron
## 2
         chr2 191099907 191099907
                                                             9e-14
                                      1
                                                rs7574865
                                                                       intron
## 3
                                                             6e-29
              32082981 32082981
         chr6
                                      1
                                                rs1150754
                                                                       intron
## 4
                                                             6e-29
               32082981
                         32082981
                                      1
                                                rs1150754
         chr6
                                                                       intron
## 5
               32082981 32082981
                                      1
                                             - rs1150754
                                                             6e-29
         chr6
                                                                       intron
                                                             4e-09 intergenic
## 6
         chr6
               32689659
                         32689659
                                      1
                                              * rs3129716
## 7
                         32689659
         chr6
               32689659
                                      1
                                              * rs3129716
                                                             4e-09 intergenic
## 8
               32689659
                         32689659
         chr6
                                      1
                                              * rs3129716
                                                             4e-09 intergenic
## 9
               32689659 32689659
                                      1
                                              * rs3129716
                                                             4e-09 intergenic
         chr6
                         32689659
                                      1
## 10
          chr6
               32689659
                                              * rs3129716
                                                             4e-09 intergenic
         chr6 32689659 32689659
## 11
                                      1
                                             * rs3129716
                                                             4e-09 intergenic
## 12
         chr1 235876577 235876577
                                      1
                                             - rs9782955
                                                             1e-09
                                                                       intron
## 13
         chr7 50267214 50267214
                                      1
                                              * rs11185603
                                                             4e-07 intergenic
## 14
               73152652 73152652
                                      1
                                              * rs11235667
        chr11
                                                             7e-11 intergenic
##
        symbol
## 1
         NAB1
## 2
        STAT4
## 3
       LY6G6C
## 4
         TNXB
## 5
         PPT2
## 6
     HLA-DQB1
      HLA-DOB
## 7
## 8
      HLA-DMA
## 9
      HLA-DOA
## 10 HLA-DPA1
## 11 HLA-DPB1
## 12
         LYST
## 13
         IKZF1
## 14
       FCHSD2
```

We note that some of the SNPs are assigned to more than one gene. This is because enhancers are promiscuous and can regulate multiple genes.

We can now check if any of these genes is differentially expressed in our RNA-seq data:

```
snps_hard_in_fantom_in_degs <- merge(snps_hard_in_fantom, degs, by = "symbol", all = FALSE)</pre>
snps_hard_in_fantom_in_degs
## DataFrame with 2 rows and 18 columns
##
       symbol seqnames
                           start
                                        end
                                                width
                                                         strand
                                                                       SNPS
##
     <factor> <factor> <integer> <integer> <integer> <factor> <character>
## 1 HLA-DOA
                  chr6
                       32689659
                                  32689659
                                                                  rs3129716
                                                    1
## 2
        IKZF1
                  chr7
                        50267214 50267214
                                                    1
                                                                 rs11185603
                                      gene_id bp_length baseMean
##
       P.VALUE
                 LOCATION
##
     <numeric>
                 <factor>
                                  <character> <integer> <numeric>
                                                   4012 962.7578
## 1
         4e-09 intergenic ENSG00000204252.13
                                                   9784 7183.7639
## 2
         4e-07 intergenic ENSG00000185811.16
##
     log2FoldChange
                         lf cSE
                                     stat
                                                pvalue
                                                              padj
##
          <numeric>
                     <numeric> <numeric>
                                             <numeric>
                                                         <numeric>
## 1
         -0.4424595 0.15882236 -2.785877 0.0053383163 0.04431304
## 2
         -0.2575717 0.07647486 -3.368057 0.0007569983 0.01162554
##
          ensembl id
##
         <character>
## 1 ENSG00000204252
## 2 ENSG00000185811
```

We have identified 2 genes whose putative enhancers contain SLE GWAS SNPs. Let's add these to our list:

```
prioritised_hits <- unique(rbind(prioritised_hits, data.frame(
    snp_id = snps_hard_in_fantom_in_degs$SNPS,
    snp_pvalue = snps_hard_in_fantom_in_degs$P.VALUE,
    snp_location = snps_hard_in_fantom_in_degs$LOCATION,
    gene_id = snps_hard_in_fantom_in_degs$gene_id,
    gene_symbol = snps_hard_in_fantom_in_degs$symbol,</pre>
```

```
gene_pvalue = snps_hard_in_fantom_in_degs$padj,
gene_log2foldchange = snps_hard_in_fantom_in_degs$log2FoldChange)))
prioritised_hits
```

```
##
                        snp_id snp_pvalue snp_location
                                                                   gene_id
                                    1e-06
## ENSG00000096968
                     rs1887428
                                               fiveUTR ENSG00000096968.13
                    rs58688157
                                              promoter ENSG00000099834.18
## ENSG00000099834
                                    5e-13
## ENSG00000115267
                     rs1990760
                                    4e-08
                                                coding ENSG00000115267.5
                                                coding ENSG00000120280.5
## ENSG00000120280
                                    5e-10
                     rs887369
                                                coding ENSG00000185507.19
## ENSG0000185507
                     rs1061502
                                    9e-11
## ENSG00000204366
                     rs558702
                                    8e-21
                                              promoter ENSG00000204366.3
## ENSG00000275106
                    rs10488631
                                    2e-11
                                              promoter ENSG00000275106.1
                                            intergenic ENSG00000130513.6
## 1
                     rs8105429
                                    5e-06
## 2
                                    6e-15
                                                intron ENSG00000140497.16
                     rs2289583
## 3
                     rs3794060
                                    1e-20
                                                intron ENSG00000172890.11
                                                intron ENSG00000214894.6
## 4
                     rs9267531
                                    8e-08
## 5
                   rs114090659
                                    6e-92
                                            intergenic ENSG00000214894.6
## 6
                                                intron ENSG00000214894.6
                     rs3131379
                                    2e-52
## 11
                                    4e-09
                                            intergenic ENSG00000204252.13
                     rs3129716
                    rs11185603
## 21
                                    4e-07
                                             intergenic ENSG00000185811.16
##
                   gene_symbol gene_pvalue gene_log2foldchange
## ENSG00000096968
                          JAK2 2.068794e-02
                                                      0.4854343
## ENSG00000099834
                         CDHR5 1.732902e-02
                                                      0.8539586
## ENSG0000115267
                         IFIH1 1.120363e-03
                                                       1.1494945
## ENSG00000120280
                       CXorf21 6.047898e-05
                                                      0.7819504
## ENSG0000185507
                          IRF7 2.298336e-04
                                                      1.4062704
## ENSG00000204366
                        ZBTB12 3.584479e-02
                                                      -0.3892298
## ENSG00000275106
                            NA 1.797861e-02
                                                      0.7344844
## 1
                         GDF15 4.481544e-02
                                                      0.7883703
## 2
                        SCAMP2 1.192679e-02
                                                      -0.2959934
## 3
                       NADSYN1 3.338101e-02
                                                      0.2619770
## 4
                     LINC00243 2.442643e-04
                                                      1.2684089
## 5
                     LINC00243 2.442643e-04
                                                      1.2684089
                     LINC00243 2.442643e-04
## 6
                                                      1 2684089
## 11
                       HLA-DOA 4.431304e-02
                                                     -0.4424595
## 21
                         IKZF1 1.162554e-02
                                                      -0.2575717
```

Promoter Capture Hi-C data

More recently, chromatin interaction data was generated across 17 human primary blood cell types ²⁵. More than 30,000 promoter baits were used to capture promoter-interacting regions genome-wide. These regions were then mapped to enhancers based on the Ensembl Regulatory Build ⁵⁷ and can be accessed in the supplementary data of the paper:

```
# uncomment the following line to download file
#download.file("http://www.cell.com/cms/attachment/2086554122/2074217047/mmc4.zip", destfile = "mmc4"
# uncomment the following lines to extract zipped files
#unzip("mmc4.zip")
#unzip("DATA_S1.zip")
pchic <- read.delim("ActivePromoterEnhancerLinks.tsv", stringsAsFactors = FALSE)
head(pchic)</pre>
```

```
baitChr baitSt baitEnd baitID oeChr
##
                                              oeSt
                                                     oeEnd oeID
## 1
        chr1 1206873 1212438
                                254 chr1
                                           943676
                                                    957199
                                                            228
## 2
        chr1 1206873 1212438
                                 254
                                      chr1 1034268 1040208
                                                            235
## 3
        chr1 1206873 1212438
                                 254
                                      chr1 1040208 1043143
                                                            236
## 4
        chr1 1206873 1212438
                                 254
                                      chr1 1069045 1083958
                                                            242
## 5
        chr1 1206873 1212438
                                 254
                                      chr1 1083958 1091234
                                                            243
## 6
        chr1 1206873 1212438
                                 254
                                      chr1 1585571 1619752
                                                            304
##
                         cellType.s.
## 1
## 2 nCD4,nCD8,Mac0,Mac1,Mac2,MK,Mon
## 3
         nCD4, nCD8, Mac0, Mac1, Mac2, MK
```

In this case, we will have to map the promoter baits to genes first. We can do this by converting the baits to a GRanges object and then using the TxDb object we previously built to extract positions of transcription start sites (TSSs):

```
baits <- GRanges(seqnames = pchic$baitChr, ranges = IRanges(start = pchic$baitSt,
tsss <- promoters(txdb, upstream = 0, downstream = 1, columns = "gene_id")
hits <- nearest(baits, tsss)
baits$gene_id <- unlist(tsss[hits]$gene_id)
baits</pre>
```

```
## GRanges object with 51142 ranges and 1 metadata column:
##
             seqnames
                                     ranges strand
                                                                 gene_id
##
                <Rle>
                                  <IRanges> <Rle> |
                                                             <character>
##
         [1]
                 chr1
                         [1206873, 1212438]
                                                 * | ENSG00000186827.10
##
         [2]
                 chr1
                        [1206873, 1212438]
                                                 * | ENSG00000186827.10
                        [1206873, 1212438]
[1206873, 1212438]
##
         [3]
                 chr1
                                                 * | ENSG00000186827.10
##
         [4]
                 chr1
                                                  * | ENSG00000186827.10
##
         [5]
                 chr1
                         [1206873, 1212438]
                                                  * | ENSG00000186827.10
##
##
     [51138]
                 chrY [22732049, 22743996]
                                                 *
                                                      ENSG00000230727.1
##
     [51139]
                 chrY [22732049, 22743996]
                                                  *
                                                      ENSG00000230727.1
##
     [51140]
                 chrY [22732049, 22743996]
                                                  *
                                                      ENSG00000230727.1
##
     [51141]
                 chrY [22732049, 22743996]
                                                  *
                                                       ENSG00000230727.1
##
     [51142]
                 chrY [22732049, 22743996]
                                                  *
                                                       ENSG00000230727.1
##
##
     seqinfo: 24 sequences from an unspecified genome; no seqlengths
```

Now we can create a GRanges object of the enhancers in the promoter capture Hi-C data with the bait annotation attached:

```
pchic <- GRanges(seqnames = pchic$oeChr, ranges = IRanges(start = pchic$oeSt, end = pchic$oeEnd), ge
pchic <- unique(pchic)
pchic</pre>
```

```
## GRanges object with 25232 ranges and 1 metadata column:
##
             seqnames
                                     ranges strand
                                                                  gene_id
##
                <R.1e>
                                  <IRanges> <Rle> |
                                                              <character>
##
         [1]
                         [ 943676, 957199]
                                                  * | ENSG00000186827.10
                 chr1
                         [1034268, 1040208]
##
         [2]
                                                  * | ENSG00000186827.10
                 chr1
         [3]
                         [1040208, 1043143]
##
                 chr1
                                                  * | ENSG00000186827.10
##
         [4]
                 chr1
                         [1069045, 1083958]
                                                  * | ENSG00000186827.10
##
         [5]
                 chr1
                         [1083958, 1091234]
                                                  * | ENSG00000186827.10
##
     [25228]
##
                 chrY [23401616, 23404873]
                                                  *
                                                       ENSG00000230727.1
                 chrY [23404938, 23407193]
                                                       ENSG00000230727.1
##
     [25229]
                                                  *
                                                  *
                                                       ENSG00000230727.1
##
                 chrY [23409014, 23410287]
     [25230]
                 \mathtt{chrY} \ \texttt{[23410287, 23411837]}
##
     [25231]
                                                  *
                                                        ENSG00000230727.1
                 chrY [23411837, 23412539]
##
     [25232]
                                                       ENSG00000230727.1
##
##
     seqinfo: 24 sequences from an unspecified genome; no seqlengths
```

Next, we basically repeat the steps we have taken when working with the FANTOM5 data to find SLE GWAS SNPs overlapping with these enhancers:

```
hits <- findOverlaps(snps_hard, pchic)
snps_hard_in_pchic = snps_hard[queryHits(hits)]
pchic_with_snps_hard = pchic[subjectHits(hits)]
mcols(snps_hard_in_pchic) <- cbind(mcols(snps_hard_in_pchic), mcols(pchic_with_snps_hard))</pre>
snps_hard_in_pchic <- as.data.frame(snps_hard_in_pchic)</pre>
snps_hard_in_pchic
##
                               end width strand
                                                        SNPS P.VALUE
      segnames
                   start
## 1
          chr6
                31753256
                          31753256
                                       1
                                                   rs3131379
                                                               2e-52
## 2
          chr6
                32696681
                          32696681
                                       1
                                                   rs2647012
                                                               8e-06
                30631546
## 3
                          30631546
                                                               3e-08
         chr16
                                                   rs7197475
## 4
         chr20
                 4762059
                           4762059
                                       1
                                                   rs6084875
                                                               2e-06
## 5
                                                               4e-09
          chr6
                32689659
                          32689659
                                       1
                                                   rs3129716
## 6
                31668965
                         31668965
                                                               8e-08
          chr6
                                       1
                                                   rs9267531
## 7
          chr6 31951083 31951083
                                       1
                                                   rs1270942
                                                              2e-165
          chr6 106140931 106140931
## 8
                                       1
                                                   rs6568431
                                                               5e-14
## 9
          chr7 28146272 28146272
                                       1
                                                   rs849142
                                                               9e-11
          chr2 65381229 65381229
## 10
                                                    rs268134
                                                               1e-10
                39850937 39850937
                                              - rs143123127
## 11
         chr17
                                       1
                                                               6e-09
## 12
          chr9
                86916761 86916761
                                               * rs190029011
                                                               3e-06
## 13
         chr11
                65637829 65637829
                                                    rs931127
                                                               7e-06
         chr19 18370523 18370523
                                                               5e-06
## 14
                                       1
                                                   rs8105429
         chr16 85977731 85977731
## 15
                                       1
                                                               4e-06
                                                  rs10521318
          chr5 39406395 39406395
## 16
                                       1
                                                   rs3914167
                                                               86-06
## 17
         chr16 31315385 31315385
                                               + rs11860650
                                                               2e-20
                                       1
                            gene_id
##
        LOCATION
## 1
          intron ENSG00000219797.2
## 2
      intergenic ENSG00000204290.10
      intergenic ENSG00000180096.11
## 4
      intergenic ENSG00000212536.1
## 5
      intergenic ENSG00000204290.10
## 6
          intron ENSG00000219797.2
          intron ENSG00000225851.1
## 7
          intron ENSG00000112297.14
## 8
## 9
          intron ENSG00000106052.13
## 10
          intron ENSG00000198369.9
## 11
          intron ENSG00000277363.4
## 12 intergenic ENSG00000223012.1
## 13 intergenic ENSG00000245532.6
## 14 intergenic ENSG00000099308.10
## 15 intergenic ENSG00000279907.1
          intron ENSG00000212296.1
## 16
          intron ENSG00000260060.1
## 17
We check if any of these enhancer containing SLE variants are known to putatively regulate genes differentially
expressed in SLE:
snps_hard_in_pchic_in_degs <- merge(snps_hard_in_pchic, degs, by = "gene_id", all = FALSE)</pre>
snps_hard_in_pchic_in_degs
## DataFrame with 4 rows and 18 columns
##
                gene_id seqnames
                                                          width
                                                                   strand
                                      start
                                                  end
##
            <character> <factor> <integer> <integer> <integer> <factor>
## 1 ENSG00000106052.13
                            chr7
                                  28146272
                                             28146272
                                                              1
## 2 ENSG00000219797.2
                            chr6
                                  31753256
                                             31753256
                                                              1
## 3 ENSG00000219797.2
                            chr6
                                  31668965
                                            31668965
                                                              1
                                                                        +
## 4 ENSG00000245532.6
                           chr11 65637829 65637829
                                                              1
##
                  P.VALUE
                             LOCATION bp_length
                                                       symbol
                                                                 baseMean
##
     <character> <numeric>
                             <factor> <integer>
                                                       t>
                                                                <numeric>
## 1
       rs849142
                     9e-11
                                            9165
                                                      TAX1BP1
                                                               2406.26093
                               intron
## 2
       rs3131379
                     2e-52
                               intron
                                             498
                                                           ΝA
                                                                 74.58175
## 3
       rs9267531
                     8e-08
                                             498
                                                                 74.58175
                               intron
                                                           NΑ
```

22767 NEAT1, MIR612 17580.27601

4

rs931127

7e-06 intergenic

```
##
     log2FoldChange
                        lfcSE
                                   stat
                                               pvalue
                                                              padj
##
          <numeric> <numeric> <numeric>
                                            <numeric>
                                                         <numeric>
## 1
          0.3438396 0.1205716 2.851746 4.347982e-03 0.0386695506
## 2
          0.5586633 0.1116884 5.001982 5.674388e-07 0.0000798169
## 3
          0.5586633 0.1116884 5.001982 5.674388e-07 0.0000798169
## 4
          0.5259525 0.1366133 3.849935 1.181492e-04 0.0032213554
##
          ensembl id
##
         <character>
## 1 ENSG00000106052
## 2 ENSG00000219797
## 3 ENSG00000219797
## 4 ENSG00000245532
```

And finally we add these 3 genes to our list. These are our final results:

```
prioritised_hits <- unique(rbind(prioritised_hits, data.frame(
    snp_id = snps_hard_in_pchic_in_degs$SNPS,
    snp_pvalue = snps_hard_in_pchic_in_degs$P.VALUE,
    snp_location = snps_hard_in_pchic_in_degs$LOCATION,
    gene_id = snps_hard_in_pchic_in_degs$gene_id,
    gene_symbol = snps_hard_in_pchic_in_degs$symbol,
    gene_pvalue = snps_hard_in_pchic_in_degs$padj,
    gene_log2foldchange = snps_hard_in_pchic_in_degs$log2FoldChange)))
prioritised_hits</pre>
```

```
##
                        snp_id snp_pvalue snp_location
                                                                   gene_id
## ENSG00000096968
                     rs1887428
                                    1e-06
                                               fiveUTR ENSG00000096968.13
## ENSG00000099834
                    rs58688157
                                    5e-13
                                               promoter ENSG00000099834.18
## ENSG00000115267
                     rs1990760
                                    4e-08
                                                 coding ENSG00000115267.5
                                                 coding ENSG00000120280.5
## ENSG00000120280
                     rs887369
                                    5e-10
## ENSG0000185507
                     rs1061502
                                    9e-11
                                                 coding ENSG00000185507.19
## ENSG00000204366
                      rs558702
                                    8e-21
                                               promoter ENSG00000204366.3
## ENSG00000275106
                    rs10488631
                                    2e-11
                                                        ENSG00000275106.1
                                              promoter
## 1
                     rs8105429
                                    5e-06
                                             intergenic ENSG00000130513.6
## 2
                                    6e-15
                                                 intron ENSG00000140497.16
                     rs2289583
## 3
                     rs3794060
                                    1e-20
                                                 intron ENSG00000172890.11
                                                 intron ENSG00000214894.6
## 4
                     rs9267531
                                    8e-08
                                             intergenic ENSG00000214894.6
## 5
                   rs114090659
                                    6e-92
## 6
                                    2e-52
                                                 intron ENSG00000214894.6
                     rs3131379
## 11
                     rs3129716
                                    4e-09
                                            intergenic ENSG00000204252.13
                                    4e-07
## 21
                    rs11185603
                                             intergenic ENSG00000185811.16
## 12
                      rs849142
                                    9e-11
                                                 intron ENSG00000106052.13
## 22
                     rs3131379
                                    2e-52
                                                 intron ENSG00000219797.2
## 31
                                    8e-08
                                                 intron ENSG00000219797.2
                     rs9267531
                                    7e-06
## 41
                      rs931127
                                             intergenic ENSG00000245532.6
##
                    gene_symbol gene_pvalue gene_log2foldchange
## ENSG00000096968
                           JAK2 2.068794e-02
                                                       0.4854343
## ENSG00000099834
                          CDHR5 1.732902e-02
                                                        0.8539586
## ENSG00000115267
                          IFIH1 1.120363e-03
                                                        1.1494945
## ENSG00000120280
                        CXorf21 6.047898e-05
                                                        0.7819504
## ENSG0000185507
                           IRF7 2.298336e-04
                                                        1.4062704
## ENSG00000204366
                         ZBTB12 3.584479e-02
                                                       -0.3892298
## ENSG00000275106
                             NA 1.797861e-02
                                                        0.7344844
## 1
                          GDF15 4.481544e-02
                                                        0.7883703
## 2
                         SCAMP2 1.192679e-02
                                                       -0.2959934
## 3
                        NADSYN1 3.338101e-02
                                                        0.2619770
## 4
                      LINC00243 2.442643e-04
                                                        1.2684089
## 5
                      LINC00243 2.442643e-04
                                                        1.2684089
## 6
                      LINC00243 2.442643e-04
                                                       1.2684089
## 11
                        HLA-DOA 4.431304e-02
                                                       -0.4424595
## 21
                          IKZF1 1.162554e-02
                                                       -0.2575717
## 12
                        TAX1BP1 3.866955e-02
                                                        0.3438396
## 22
                             NA 7.981690e-05
                                                        0.5586633
## 31
                             NA 7.981690e-05
                                                        0.5586633
## 41
                   NEAT1, M.... 3.221355e-03
                                                        0.5259525
```

Conclusions

In this Bioconductor workflow we have used several packages and datasets to demonstrate how regulatory genomic data can be used to annotate significant hits from GWASs and provide an intermediate layer connecting genetics and transcriptomics. Overall, we identified 17 SLE-associated SNPs that we mapped to 16 genes differentially expressed in SLE, using eQTL data 14 and enhancer - promoter relationships from CAGE 19 and promoter capture Hi-C experiments 25 .

While simplified, the workflow also demonstrates some real-world challenges encountered when working with genomic data from different sources, such as the use of different genome references and gene annotation conventions, the parsing of files with custom formats into Bioconductor-compatible objects and the mapping of genomic locations to genes.

As the sample size and power of GWASs and gene expression studies continue to increase, it will become more and more challenging to identify truly significant hits and interpret them. The use of regulatory genomics data as presented here can be an important skill and tool to gain insights into large biomedical datasets and help in the identification of biomarkers and therapeutic targets.

Abbreviations

CAGE: cap analysis of gene expression DHS: DNase I hypersensitive site eQTL: expression quantitative trait locus GWAS: genome-wide association study PheWAS: phenome-wide association study SLE: systemic lupus erythematosus SNP: single nucleotide polymorphism

TSS: trancription start site

Data and software availability

Download links for all datasets are part of the workflow. Software packages required to reproduce the analysis can be installed as part of the workflow. Code is available at https://github.com/enricoferrero/bioconductor-regulatory-genomics-workflow.

Competing interests

EF is a full time employee of GSK.

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