

# 🔒 Chromatin loops and expression QTL colocalization reveal novel gene targets for T1D-associated GWAS variants in immune cells

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
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This protocol was made as part of the DChallenge and gives a high level explanation of our steps.

Type 1 diabetes (T1D) is a disease characterized by the destruction of  $\beta$  cell populations in the pancreas. Immune cells, and specifically T cells, have been implicated to play a key role in destroying insulin producing  $\beta$  cells by infiltrating the pancreas. To better understand the role of immune regulation in T1D, we colocalized the gene expression quantitative trait loci (eQTL) signals from 18 different immune cell populations (15 from DICE, 3 from BLUEPRINT) with T1D GWAS signals to gather non-coding variants that are likely causal for both the gene expression and the disease association. We further overlapped these variants with chromatin loops mapped in a subset of these immune cell populations to identify potential target genes of the significant non-coding SNPs. Aside from well-studied genes such as *BACH2*, *UBASH3A*, *PTPN22* and *SIRPG*, we identified *AP003774.1*, a long non-coding RNA, that is looping to a ~15kb away regulatory element overlapping a colocalized SNP (rs479777) in various T cell subsets. The looped region overlaps the promoter of another gene (promoter-promoter loop), *CCDC88B*, however, the eQTL association for this SNP is specific to *AP003774.1* and is remarkably strong for resting T cell subsets, NK cells and naïve B cells. The same SNP creates strong binding sites for multiple important transcription factors in donors with the non-reference allele leading to higher expression of *AP003774.1*. We hypothesize that the overexpression of *AP003774.1* lncRNA mediated through specific non-coding variants in different immune cell populations play a role in immune-related aspects of T1D.

<https://github.com/joreynajr/dchallenge>

Joaquin Reyna, Sourya Bhattacharyya, Nikhil Rao, Abhijit Chakraborty, Ferhat Ay .  
Chromatin loops and expression QTL colocalization reveal novel gene targets for T1D-  
associated GWAS variants in immune cells. **protocols.io**  
<https://protocols.io/view/chromatin-loops-and-expression-qt1-colocalization-bzy7p7zn>



Type 1 Diabetes, HiChIP, HiC, Colocalization, eQTL, GWAS

<https://thenounproject.com/term/weaving/>

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## Main Colocalization Pipeline

### 1 Download the GWAS summary statistics (uses GRCh38 coordinates)

Download GWAS summary statistics from the GWAS catalogue.

Download link: [http://ftp.ebi.ac.uk/pub/databases/gwas/summary\\_statistics/GCST90014001-GCST90015000/GCST90014023/GCST90014023\\_buildGRCh38.tsv](http://ftp.ebi.ac.uk/pub/databases/gwas/summary_statistics/GCST90014001-GCST90015000/GCST90014023/GCST90014023_buildGRCh38.tsv)

Chiou, J., Geusz, R.J., Okino, ML. et al (2021). Interpreting type 1 diabetes risk with genetics and single-cell epigenomics. Nature.

<https://doi.org/10.1038/s41586-021-03552-w>

#### 1.1 Remap the GWAS SNP coordinate to GRCh37

liftOver

```
liftOver -bedPlus=3 -tab <in bed> <chain file> <out lift file>  
<unmapped>
```

UCSC tool to convert from one coordinate system to another.

## 1.2 Filter for SNPs with genome-wide significance ( $5e-8$ )

## 1.3 Reformat the file into the required format for colocalization

Estimate standard error of the SNP using beta and MAF values. Standard error is beta/z-score

## 2 Download the eQTL summary statistics (uses GRCh37 coordinates)

Many eQTL studies have already been completed with summary statistics. For our project we downloaded preprocessed data from the Mu et al., 2021 which contains Blueprint and DICE eQTL results.

Download link: <https://zenodo.org/record/4480206#.YXcophrMJyw>

Mu, Z., Wei, W., Fair, B. et al (2021). The impact of cell type and context-dependent regulatory variants on human immune traits. Genome Biol.

<https://doi.org/10.1186/s13059-021-02334-x>

## 3 Run colocalization between the T1D GWAS and all eQTL studies

The coloc package that we used can be referenced here: <https://cran.r-project.org/web/packages/coloc/index.html>

Colocalization

```
Rscript Colocalization_Analysis_GWAS.R <in GWAS> <output directory> <in  
eQTL>
```

Colocalization scripts generated by Sourya Bhattacharyya.

## 4 Intersect colocated SNPs with HiChIP data

Obtain HiChIP-seq data for several cell types. In our case, we obtained data from CD4+ T-cells, CD8+ T-cells, classical monocytes, non-classical monocytes, naive B cells, natural killer, T follicular helper, Th1, Th17, Th2 and TH1/17, TREGMEM, and TREGNAIVE cells.

## 5 Generate a Master table of SNP-gene pairs with loops

For this master table we focused on colocated SNPs, their colocated eGenes and other genes as well.

### 5.1 Extract all SNP-gene pairs that are +/- 500kb from a colocated SNP

```
bedtools intersect  
  
bedtools intersect <bed1> <bed2> > <output bed>  
Intersect two lists of genetic loci.
```

### 5.2 Intersect loops with SNP-gene pairs

```
bedtools pairtopair  
  
bedtools pairtopair <bedpe1> <bedpe2> > <output>  
Intersect two lists of genetic loci pairs.
```

### 5.3 Label each SNP-Gene pair

For each SNP-Gene pair label whether it is an eQTL, colocated pair, contains a loop and all metadata.

Gene Candidate Prioritization

## 6 Investigate Candidate Genes using the WashU Epigenome Browser

### 6.1 Generate BED + Index files for 1D tracks

- single-cell ATAC-seq data
- SNP and gene locations

bgzip

**bgzip <input> <output>**

Compress bed file with bgzip.

tabix

**tabix <bed.gz>**

Create an index for compressed bed file.

## 6.2 **Generate BEDPE + Index files for 2D tracks**

- loop data for all cell lines

bgzip

**bgzip <input> <output>**

Compress bed file with bgzip.

tabix

**tabix <bed.gz>**

Create an index for compressed bed file.

## 6.3 **Add ChromHMM tracks from the Public Hubs**

## 6.4

**Look for genes with several SNPs which overlap important loops**

## **7 Investigate TF binding sites using the Genome Browser + others**

### **7.1 Load JASPAR tracks, query SNP locations, investigate motifs**

- JASPAR tracks can be added from the Public Hub
- <https://jaspar.genereg.net/>

### **7.2 Query SNPs using ADAstra website**

<https://adastra.autosome.ru/zanthar>

## **8 Confirm expression of candidate genes using the DICE database**

<https://dice-database.org/>