Using haploR, an R package for querying HaploReg and RegulomeDB

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Overview

HaploReg (http://archive.broadinstitute.org/mammals/haploreg/haploreg.php) and RegulomeDB (http://www.regulomedb.org) are web-based tools that extract biological information such as eQTL, LD, motifs, etc. from large genomic projects such as ENCODE, the 1000 Genomes Project, Roadmap Epigenomics Project and others. This is sometimes called “post stage GWAS” analysis.

The R-package **haploR** was developed to query those tools (HaploReg and RegulomeDB) directly from R in order to facilitate high-throughput genomic data analysis. Below we provide several examples that show how to work with this package.

Note: you must have a stable Internet connection to use this package.

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Motivation and typical analysis workflow

This package was inspired by the fact that many web-based post stage GWAS databases do not have Application Programing Interface (API) and, therefore, do not allow users to query them remotely from R environment. In our research we used HaploReg and RegulomeDB web databases. These very useful web databases show information about linkage disequilibrium of query variants and variants which are in LD with them, expression quantitative trait loci (eQTL), motifs changed and other useful information. However, it is not easy to include this information into streamlined workflow since those tools also not offer API.

We developed a custom analysis pipeline which prepares data, performs genome-wide association (GWA) analysis and presents results in a user-friendly format. Results include a list of genetic variants (also known as ‘SNP’ or single nucleotide polymorphism), their corresponding p-values, phenotypes (traits) tested and other meta-information such as LD, alternative allele, minor allele frequency, motifs changed, etc. Of course, we could go through the list of SNPs having genome-wide significant p-values (1e-8) and submit each of those SNPs to HaploReg or RegulomeDB manually, one-by-one, but it is time-consuming process and will not be fully automatic (which ruins one of the pipeline’s paradigms). This is especially difficult if the website does not offer downloading results.

Therefore, we developed **haploR**, a user-friendly R package that connects to HaploReg or RegulomeDB from R environment with methods POST and GET and downloads results in a suitable R format. This package significantly saved our time in developing reporting system for our internal genomic analysis pipeline and now we would like to present **haploR** to the research community.

Example of typical analysis workflow is shown below.

- Data preprocessing stage usually consists of basic cleaning operations (sex inconsistencies check, filtering by MAF, missing genotype rate), format conversion, population stratification, creating temporary files, etc.
- Genome-wide association study (GWAS). This includes testing hypotheses on correlation between phenotype and genotype. Usually in form of linear or logistic regression but can be quite sophisticated especially for rare variants.
Typical workflow example

Figure 1: Typical analysis workflow
• Postprocessing usually includes summarizing results in tables, creating graphics (Manhattan plot, QQ-plot), sometimes filtering by significance threshold (usually 1E-8), removing temporary files, etc.
• Post stage GWAS analysis: connect GWAS findings with existing studies and gather information about linked SNPs, chromatin state, protein binding annotation, sequence conservation across mammals, effects on regulatory motifs and on gene expression, etc. This helps researchers to more understand functions and test additional hypotheses of found genome-wide significant SNPs. At this stage haploR is especially useful because it provides a convenient R interface to mining web databases. This, in turn, streamlines analysis workflow and therefore significantly reduces analysis time. Previously researchers had to do it manually after analysis by exploring available web resources, checking each SNP of interest and downloading results (which is especially painful if website does not have a download option). With haploR such post-GWAS information are easily retrieved, shared and appeared in the final output tables at the end of analysis. This will save researchers’ time.

**Installation of haploR package**

In order to install the haploR package, the user must first install R (https://www.r-project.org). After that, haploR can be installed either:

- From CRAN (stable version):
  ```r
  install.packages("haploR", dependencies = TRUE)
  ```
- Or from the package web page (developing version):
  ```r
  devtools::install_github("izhbannikov/haploR")
  ```

haploR depends on the following packages:

- `httr`, version 1.2.1 or later.
- `XML`, version version 3.98-1.6 or later.
- `tibble`, version 1.3.0 or later.
- `RUnit`, version 0.4.31 or later.

**Examples of usage**

**Querying HaploReg**

Function

```r
queryHaploreg(query = NULL, file = NULL, study = NULL, ldThresh = 0.8, 
ldPop = "EUR", epi = "vanilla", cons = "siphy", genotypes = "gencode", 
timeout = 10, encoding = "UTF-8", verbose = FALSE)
```

queries HaploReg web-based tool and returns results.

**Arguments**

- `query`: Query (a vector of rsIDs).
- `file`: A text file (one refSNP ID per line).
- `study`: A particular study. See function getHaploRegStudyList(...). Default: NULL.
- `ldThresh`: LD threshold, r2 (select NA to show only query variants). Default: 0.8.
epi: Source for epigenomes. Possible values: vanilla for ChromHMM (Core 15-state model); imputed for ChromHMM (25-state model using 12 imputed marks); methyl for H3K4me1/H3K4me3 peaks; acetyl for H3K27ac/H3K9ac peaks. Default: vanilla.


genetypes: Show position relative to. Possible values: gencode for Gencode genes; refseq for RefSeq genes; both for both. Default: gencode.


timeout: A timeout parameter for curl. Default: 10

encoding: Set the encoding for correct retrieval web-page content. Default: UTF-8

verbose: Verbosing output. Default: FALSE.

Output

A data frame (table) wrapped into a tibble object contains data extracted from HaploReg site. The columns (33 in total at the time of writing this vignette) are specific to HaploReg output. Below we describe the columns:

- chr: Chromosome, type: numeric
- pos_hg38: Position on the human genome, type: numeric.
- r2: Linkage disequilibrium. Type: numeric.
- D': Linkage disequilibrium, alternative definition. Type: numeric.
- is_query_snp: Indicator shows query SNP, 0 - not query SNP, 1 - query SNP. Type: numeric.
- rsID: refSNP ID. Type: character.
- ref: Reference allele. Type: character.
- alt: Alternative allele. Type: character.
- AFR: r2 calculated for Africa. Type: numeric.
- AMR: r2 calculated for America. Type: numeric.
- ASN: r2 calculated for Asia. Type: numeric.
- EUR: r2 calculated for Europe. Type: numeric.
- GERP_cons: GERP scores. Type: numeric.
- SiPhy_cons: SiPhy scores. Type: numeric.
- Chromatin_States: Chromatin states: reference epigenome identifiers (EID) of chromatin-associated proteins and histone modifications in that region. Type: character.
- Chromatin_States_Imputed: Chromatin states based on imputed data. Type: character.
- Chromatin_Marks: Chromatin marks Type: character.
- DNAse: DNAse. Type: character.
- Proteins: A list of protein names. Type: character.
- eQTL: Expression Quantitative Trait Loci. Type: character.
- gwas: GWAS study name. Type: character.
- grasp: GRASP study name. Type: character.
- Motifs: Motif names. Type: character.
- GENCODE_id: GENCODE transcript ID. Type: character.
- GENCODE_name: GENCODE gene name. Type: character.
- GENCODE_direction: GENCODE direction (transcription toward 3' or 5' end of the DNA sequence). Type: numeric.
- GENCODE_distance: GENCODE distance. Type: numeric.
- RefSeq_id: NCBI Reference Sequence Accession number. Type: character.
- RefSeq_name: NCBI Reference Sequence name. Type: character.
- RefSeq_direction: NCBI Reference Sequence direction (transcription toward 3' or 5' end of the DNA sequence). Type: numeric.
- dbSNP_functional_annotation: Annotated proteins associated with the SNP. Type: numeric.
- query.snp.rsid: Query SNP rs ID. Type: character.
Number of rows is not constant, at least equal or more than the number of query SNPs, and depends on \( r^2 \) parameter chosen in a query (default 0.8). This means that the program outputs not only query SNPs, but also those SNPs that have \( r^2 \geq 0.8 \) with the query SNPs.

One or several genetic variants

```r
library(haplR)
x <- queryHaploreg(query=c("rs10048158","rs4791078"))
x
```

Here `query` is a vector with names of genetic variants.

We then can create a subset from the results, for example, to choose only SNPs with \( r^2 > 0.9 \):

```r
subset.high.LD <- x[x$r2 > 0.9, c("rsID", "r2", "chr", "pos_hg38", "is_query_snp", "ref", "alt")]
subset.high.LD
```

Here `query` is a vector with names of genetic variants.

We then can create a subset from the results, for example, to choose only SNPs with \( r^2 > 0.9 \):

```r
subset.high.LD <- x[x$r2 > 0.9, c("rsID", "r2", "chr", "pos_hg38", "is_query_snp", "ref", "alt")]
subset.high.LD
```
We can then save the `subset.high.LD` into an Excel workbook:

```r
require(openxlsx)
write.xlsx(x=subset.high.LD, file="subset.high.LD.xlsx")
```

This was an example of gathering post-gwas information directly from the online tool. `haploR` has an additional advantage because it downloads the full information for query retrieved by HaploReg. For example, if you go online and submit these two SNPs to HaploReg (http://archive.broadinstitute.org/mammals/haploreg/haploreg.php), you will see that some cells of columns “Motifs changed” and “Selected eQTL hits” are hidden (only number of hits are given). `haploR` retrieves this information in a form of a data frame which can be saved into Excel file.

```
x[, c("Motifs", "rsID"),]
```

```
## # A tibble: 33 x 2
##       Motifs     rsID
## 1  AP-4_3;Asc12;E2A_5;Fbecca_disc3;HEN1_2;LBP-1_2;NRSF_disc4;NRSF_disc8;NRSF_kn7
## 2 Pou5f1_disc1;RFX5_known1;Sox_4;TATA_disc7
## 3  RFX5_known5;Zbb3
## 4  AP-1_disc1;HEN1_1;Maf_disc2;NR4A_known1;RAR;RXRA_known3;T3R
## 5  Pdx1_2
## 6  AP-1_disc1;HEY1_disc1;TATA_disc2
## 7
## 8  ATF3_disc1;LXR_2;SREBP_known4
## 9  Evnf_3;Irf_disc3;STAT_disc3
##10  GR_disc4;SZF1-1
## # ... with 23 more rows, and 1 more variables: rsID <chr>
```

```
x[, c("eQTL", "rsID"),]
```

```
## # A tibble: 33 x 2
##       eQTL     rsID
## 1 GTEx2015_v6,Heart_Left_Ventricle,PRKCA,3.25964766049438e-10
## 2 GTEx2015_v6,Heart_Left_Ventricle,PRKCA,2.87827072933431e-10;Koopman2014,Hea
## 3 GTEx2015_v6,Heart_Left_Ventricle,PRKCA,2.87827072933431e-10
## 4 GTEx2015_v6,Heart_Left_Ventricle,PRKCA,5.94596439339797e-11
## 5 GTEx2015_v6,Heart_Left_Ventricle,PRKCA,6.83955923561212e-11
## 6 GTEx2015_v6,Heart_Left_Ventricle,PRKCA,6.80544182605399e-11
## 7 GTEx2015_v6,Heart_Left_Ventricle,PRKCA,6.080544182605399e-11;Koopman2014,Hea
## 8 GTEx2015_v6,Heart_Left_Ventricle,PRKCA,2.44658806733437e-10
## 9 GTEx2015_v6,Heart_Left_Ventricle,PRKCA,2.09660151875432e-10
##10 GTEx2015_v6,Heart_Left_Ventricle,PRKCA,6.18988623085424e-12
## # ... with 23 more rows, and 1 more variables: rsID <chr>
```

### Uploading file with variants

If you have a file with your SNPs you would like to analyze, you can supply it as an input as follows:

```r
library(haploR)
x <- queryHaploreg(file="extdata/snps.txt", package = "haploR")
x
```

```
## # A tibble: 33 x 33
##     chr pos_hg38    r2 `D``` is_query_snp rsID ref alt AFR
##  <dbl>  <dbl>   <dbl> <chr>     <chr> <chr> <chr> <chr>
```
Using existing studies

Sometimes one would like to explore results from already performed study. In this case you should first explore the existing studies from HaploReg web site (http://archive.broadinstitute.org/mammals/haploreg/haploreg.php) and then use one of them as an input parameter. See example below:

```r
library(haploR)
# Getting a list of existing studies:
studies <- getStudyList()
# Let us look at the first element:
studies[[1]]

# $name
# [1] "<ce><b2>2-Glycoprotein I (<ce><b2>2-GPI) plasma levels (Athanasiadis G, 2013, 9 SNPs)"
#
# $id
# [1] "1756"
# Let us look at the second element:
studies[[2]]

# $name
# [1] "5-HTT brain serotonin transporter levels (Liu X, 2011, 1 SNP)"
#
# $id
# [1] "2362"
# Query Haploreg to explore results from this study:
x <- queryHaploreg(study=studies[[1]])
x
```

File “snps.txt” is a text file which contains one rs-ID per line:

rs10048158
rs4791078
## <dbl>  <dbl>  <dbl>  <dbl>  <dbl>  <chr>  <chr>  <chr>  <dbl>
## 1 11 34524785 0.97 1.00 0 rs836138 C A 0.34
## 2 11 34524788 0.87 0.97 0 rs11032744 C T 0.04
## 3 11 34526877 1.00 1.00 0 rs836137 A G 0.37
## 4 11 34527359 1.00 1.00 0 rs836135 G A 0.36
## 5 11 34527815 1.00 1.00 0 rs704727 T A 0.16
## 6 11 34530979 0.96 0.99 0 rs836133 C T 0.16
## 7 11 34531545 0.90 1.00 0 rs77003093 C T 0.01
## 8 11 34533644 1.00 1.00 1 rs836132 G A 0.16
## 9 11 34534390 1.00 1.00 0 rs836131 C T 0.16
## 10 11 34535548 1.00 1.00 0 rs836130 G T 0.36
## ... with 107 more rows, and 24 more variables: AMR <dbl>, ASN <dbl>,
## EUR <dbl>, GERP_cons <dbl>, SiPhy_cons <dbl>, Chromatin_States <chr>,
## Chromatin_States_Imputed <chr>, Chromatin_Marks <chr>, DNase <chr>,
## Proteins <chr>, eQTL <chr>, gwas <chr>, grasp <chr>, Motifs <chr>,
## GENCODE_id <chr>, GENCODE_name <chr>, GENCODE_direction <dbl>,
## GENCODE_distance <dbl>, RefSeq_id <chr>, RefSeq_name <chr>,
## RefSeq_direction <dbl>, RefSeq_distance <dbl>,
## dbSNP_functional_annotation <chr>, query_snp_rsid <chr>

### Querying RegulomeDB

To query RegulomeDB use this function:

```r
queryRegulome(query = NULL,
               format = "full",
               url = "http://www.regulomedb.org/results",
               timeout = 10,
               check_bad_snps = TRUE,
               verbose = FALSE)
```

This function queries RegulomeDB http://www.regulomedb.org web-based tool and returns results in a named list.

### Arguments

- **query**: Query (a vector of rsIDs).
- **format**: An output format. Only ‘full’ is currently supported. See http://www.regulomedb.org/results.
- **check_bad_snps**: Checks if all query SNPs are annotated (i.e. presented in the Regulome Database). Default: ‘TRUE’
- **verbose**: Verbosing output. Default: FALSE.

### Output

A list of two: (1) a data frame (res.table) wrapped to a tibble object and (2) a list of bad SNP IDs (bad.snp.id). Bad SNP ID are those IDs that were not found in 1000 Genomes Phase 1 data and, therefore, in RegulomeDB.

Columns in a data frame (res.table):

- `#chromosome`: Chromosome. Type: character.
- `coordinate`: Position. Type: numeric.
- `rsid`: RefSeq SNP ID. Type: character.
- `hits`: Contains information about chromatin structure: method and cell type. Type: character.
• **score**: Internal RegulomeDB score. See http://www.regulomedb.org/help#score. Type: numeric.

Number of rows is equal or less (in case if not all SNPs were found in RegulomeDB database) to number of query SNPs.

All the information retrieved from RegulomeDB, except *hits*, are also presented in HaploReg output (queryHaploReg(...)).

**Example**

```r
library(haploR)
x <- queryRegulome(c("rs4791078","rs10048158"))
x$res.table
```

```
## # A tibble: 2 x 5
## # `chr` coordinate rsid
## 1 chr17 64236317 rs10048158
## 2 chr17 64210013 rs4791078
## # ... with 2 more variables: hits <chr>, score <dbl>
```

```r
x$bad.snp.id
```

```
## # A tibble: 0 x 1
## # ... with 1 variables: rsID <chr>
```

**Session information**

```r
sessionInfo()
```

```
## # R Under development (unstable) (2017-03-04 r72303)
## # Platform: x86_64-apple-darwin13.4.0 (64-bit)
## # Running under: macOS Sierra 10.12.4
## # Matrix products: default
## # BLAS: /Library/Frameworks/R.framework/Versions/3.4/Resources/lib/libRblas.0.dylib
## # LAPACK: /Library/Frameworks/R.framework/Versions/3.4/Resources/lib/libRlapack.dylib
## # locale:
## # [1] C
## # attached base packages:
## # [1] stats graphics grDevices utils datasets methods base
## # other attached packages:
## # [1] haploR_1.4.4
## # loaded via a namespace (and not attached):
## # [1] Rcpp_0.12.10 XML_3.98-1.6 digest_0.6.12 rprojroot_1.2
## # [5] mime_0.5 R6_2.2.0 backports_1.0.5 magrittr_1.5
## # [9] evaluate_0.10 httr_1.2.1 stringi_1.1.5 curl_2.4
## # [13] RUnit_0.4.31 rmarkdown_1.4 tools_3.4.0 stringr_1.2.0
## # [17] yaml_2.1.14 compiler_3.4.0 htmltools_0.3.5 knitr_1.15.1
## # [21] tibble_1.3.0
```