Phenome Wide Association Studies (PheWAS) in R

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Packge **PheWAS** provides methods for the creation of PheWAS phenotypes, analysis, and plotting. While these methods are designed primarily for genetics based PheWAS analysis, they can perform GWAS or even phenotype only studies.

1 Data Input

There are many potential data sources and types; this necessitates that users handle the basic data i/o and formatting. Below are outlined some methods for importing common data into R.

1.1 Preparing plink data

Genome wide data is stored commonly stored in plink formats¹. The simplest method to import data from plink is the --recodeA parameter in plink. Running the following in a terminal will get one started:

```
\label{lem:plink} \begin{array}{lll} -\text{-recodeA} & -\text{-bfile example\_data --extract interesting\_snps} \\ -\text{-out } r\_\text{genotypes} \end{array}
```

This will recode the binary plink data "example_data", extracting the SNPs under investigation to the file "r_genotypes.raw". This raw data can be loaded into R with a single command²: genotypes=read.table("r_genotypes.raw",header=TRUE)

Alternatively, assuming FIDs are unique, the following will load the data ready to be put into phewas.

```
> genotypes=read.table("r_genotypes.raw",header=TRUE)[,c(-2:-6)]
> names(genotypes)[1]="id"
```

1.2 Data from file

R has robust methods for loading data from files³. For this section we will consider two examples. The first is loading a csv file containg id, icd9, and count data as appropriate for a classic

¹See http://pngu.mgh.harvard.edu/~purcell/plink/data.shtml for plink data format details.

²See http://pngu.mgh.harvard.edu/~purcell/plink/dataman.shtml#recode for details

³See ?read.table in R for the read methods discussed here.

PheWAS. id.icd9.count.csv: id,icd9,count 1,410,2 1,410.1,1 1,414.0,6 2,250.02,13

This can be loaded using csv.phenotypes=

read.csv("id.icd9.count.csv",colClasses=c("integer","character","integer"))

Pay special attention to the colClasses parameter: we need to ensure that the ICD9 codes are read as character strings so they do not lose trailing or leading zeros. This table is appropriate for use in createPhewasTable.

Another example is that the user may have exported their chart review data into a csv from a spreadsheet software.

 $example_phenotype.csv:$

```
id,T2D,max.a1c
1,T,10
2,F,NA
3,F,6
```

This can be loaded using csv.phenotypes=read.csv("example_phenotype.csv"). This table loaded into R is ready to be used in phewas-either as covariates or phenotypes (outcomes).

1.3 Data from database

The **RODBC** library contains great tools for importing data directly from electronic data warehouses. If one desired to use PheWAS codes in their analysis from an ICD9 billing code table, it might look like the following.

```
> library(RODBC)
> connection=odbcConnect("MyDSN")
> icd9.codes=sqlQuery(connection,"select id, icd9, count(distinct date)
    from icd9_codes group by id, icd9;")
> odbcClose(connection)
```

The icd9.codes data frame is ready to be used with the createPhewasTable function.

2 Data Transformation

The primary data transformation for this package is to convert and aggregate ICD9 codes into PheWAS codes. The function createPhewasTable allows for this conversion. Given the database data loaded from the above section, one can use the following code to create PheWAS phenotypes for use in phewas:

> phenotypes=createPhewasTable(icd9.codes)

There are some additional options for PheWAS code translation. Users can opt to forgo exclusions using add.exclusions=F; this increases the size of the control population, but at the cost of including potentially similar diagnoses in the control sets. The min.code.count parameter allows users to alter the specificity of case selection. It can also be set to NA to allow for continuous outcomes, the code count sum by default.

3 Phenome Wide Association Studies

The phewas function performs the PheWAS itself. Using the examples from above, one can directly pass the parameters.

> results=phewas(phenotypes=phenotypes,genotypes=genotypes)

If one wishes to speed up the analysis, a multi-threaded approach is available using snowfall.

> results=phewas(phenotypes=phenotypes,genotypes=genotypes,cores=4)

One can additionally provide covariates. In this case, we will consider an analysis adjusted by max.a1c.

```
> results=phewas(phenotypes=phenotypes,genotypes=genotypes,
+ covariates=csv.phenotypes[,c("id","max.a1c")])
```

An alternate method is to use the data parameter with name vectors in the phenotype, genotype, and covariates parameters.

```
> mydata=merge(phenotypes,genotypes)
> results=phewas(phenotypes=names(phenotypes)[-1],genotypes=c("rs1234","rs5678"),
+ data=mydata)
```

The phewas function can be used for more than just generic PheWAS. In the following example, outcomes and predictors are used for a phenotype only analysis. Note that these parameters are simply alternate names for phenotypes and genotypes, respectively.

```
> max.a1c.results=phewas(outcomes=phenotypes,
+ predictors=csv.phenotypes[,c("id","max.a1c")])
```

The phewasMeta method can assist in meta-analysis of multiple PheWAS, e.g., if one has multiple genotype platforms of data to analyze. It wraps the metagen method of the meta package.

```
> results.omni1=phewas(phenotypes=phenotypes.omni1,genotypes=genotypes.omni1)
> results.omni1$study="Omni 1"
> results.omni.express=phewas(phenotypes=phenotypes.omni.express,
+ genotypes=genotypes.omni.express)
> results.omni.express$study="Omni Express"
> results.merged=rbind(results.omni1,results.omni.express)
> results.meta=phewasMeta(results.merged)
```

4 Plotting

Three methods for plotting data are included, phewasManhattan, phenotypeManhattan, and phenotypePlot, which wrap each other. phewasManhattan is the highest level method, and can plot PheWAS results directly from phewas.

> phewasManhattan(results)

This method returns a **ggplot2** object, which can be further manipulated using methods from that package⁴. The ... parameter will pass further options into **phenotypeManhattan** and **phenotypePlot**. These lower level plot functions can be used in a stand-alone fashion for different types of data. For example, **phenotypePlot** can display information about the count for every individual of each ICD9 code.

```
> id.phenotype.value=icd9.codes
> names(id.phenotype.value)=c("id","phenotype","value")
> phenotypePlot(id.phenotype.value,use.color=F,x.group.labels=F)
```

5 Package Example

The following is the complete example from the **PheWAS** package.

```
> library(PheWAS)
> example(PheWAS)
PheWAS> ## No test:
PheWAS> #Install the recommended packages, if necessary
PheWAS> #install.packages(c("snowfall","shiny","MASS","meta"))
PheWAS> #Load the PheWAS package
PheWAS> library(PheWAS)
PheWAS> #Set the random seed so it is replicable
PheWAS> set.seed(1)
PheWAS> #Generate some example data
PheWAS> ex=generateExample()
PheWAS> #Extract the two parts from the returned list
PheWAS> id.icd9.count=ex$id.icd9.count
PheWAS> genotypes=ex$genotypes
PheWAS> #Create the PheWAS code table- translates the icd9s, adds
PheWAS> #exclusions, and reshapes to a wide format
PheWAS> phenotypes=createPhewasTable(id.icd9.count)
PheWAS> #Run the PheWAS
PheWAS> results=phewas(phenotypes,genotypes,cores=1,
          significance.threshold=c("bonferroni"))
PheWAS> #Plot the results
PheWAS> phewasManhattan(results, annotate.angle=0,
          title="My Example PheWAS Manhattan Plot")
```

 $^{^4\}mathrm{See}\ \mathrm{http://docs.ggplot2.org/current/}\ \mathrm{for\ the\ web\ documentation\ of\ ggplot2}$

```
PheWAS> #Add PheWAS descriptions
PheWAS> results_d=addPhewasDescription(results)
PheWAS> #List the significant results
PheWAS> results_d[results_d$bonferroni&!is.na(results_d$p),]
    phewas_code phewas_description
                                          snp adjustment
495
            335 Multiple sclerosis rsEXAMPLE
                                                    <NA> 0.4942269 0.06611966
                             type n_total n_cases n_controls HWE_p allele_freq
495 1.63923 7.73601e-14 logistic
                                     4416
                                             1777
                                                         2639
                                                                      0.4987545
    n_no_snp note bonferroni
495
           Ω
PheWAS> #List the top 10 results
PheWAS> results_d[order(results_d$p)[1:10],]
                                phewas_description
     phewas_code
                                                          snp adjustment
                                Multiple sclerosis rsEXAMPLE
495
             335
                                                                    <NA>
414
             293 Symptoms involving head and neck rsEXAMPLE
                                                                    <NA>
456
           313.2
                               Tics and stuttering rsEXAMPLE
                                                                    <NA>
1301
           694.1
                                          Vitiligo rsEXAMPLE
                                                                    <NA>
924
           527.2
                                     Sialoadenitis rsEXAMPLE
                                                                    <NA>
1698
             994
                                   Sepsis and SIRS rsEXAMPLE
                                                                    <NA>
1700
                                                                    <NA>
           994.2
                                            Sepsis rsEXAMPLE
1441
           736.5
                     Acquired deformities of knee rsEXAMPLE
                                                                    <NA>
486
           333.1
                                  Essential tremor rsEXAMPLE
                                                                    <NA>
548
          362.26
                      Macular puckering of retina rsEXAMPLE
                                                                    <NA>
           heta
                        SE
                                   OR
                                                        type n_total n_cases
                                                 р
495
      0.4942269 0.06611966 1.6392305 7.736010e-14 logistic
                                                                4416
                                                                        1777
414
      1.3523545 0.36437457 3.8665187 2.060831e-04 logistic
                                                                4426
                                                                          29
456 -0.9695479 0.26934689 0.3792545 3.186761e-04 logistic
                                                                4781
                                                                          56
1301 -0.9887376 0.29622485 0.3720461 8.444620e-04 logistic
                                                                4300
                                                                          46
      0.9507643 0.30809623 2.5876867 2.029145e-03 logistic
                                                                4559
                                                                          43
1698 -0.7829288 0.25514548 0.4570654 2.150942e-03 logistic
                                                                5000
                                                                          64
1700 -0.9036474 0.29708980 0.4050894 2.352742e-03 logistic
                                                                4982
                                                                          46
1441 0.8309801 0.29174434 2.2955675 4.395125e-03 logistic
                                                                4502
                                                                          49
      0.7225387 0.25386960 2.0596554 4.425805e-03 logistic
486
                                                                2709
                                                                          70
    -0.6966698 0.24841660 0.4982418 5.040388e-03 logistic
                                                                          71
                                                                4113
     n_controls HWE_p allele_freq n_no_snp note bonferroni
495
           2639
                    1
                        0.4987545
                                          0
                                                        TRUE
414
           4397
                        0.4957072
                                          0
                                                       FALSE
                    1
456
           4725
                        0.4953985
                                                       FALSE
1301
           4254
                        0.4945349
                                          0
                                                      FALSE
                    1
924
           4516
                    1
                        0.4950647
                                                      FALSE
                                          0
1698
           4936
                    1
                        0.4957000
                                                      FALSE
1700
           4936
                        0.4958852
                                                      FALSE
1441
           4453
                        0.4971124
                                          0
                                                      FALSE
                    1
486
           2639
                    1
                        0.4785899
                                                      FALSE
```

0

FALSE

0.4989059

4042

1

548

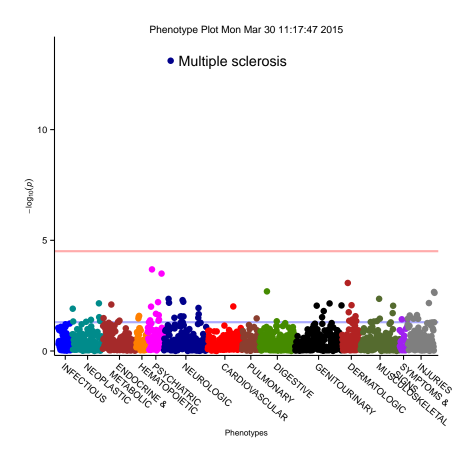


Figure 1: Example PheWAS Manhattan plot

PheWAS> ## End(No test)
PheWAS>
PheWAS>
PheWAS>

> phewasManhattan(results, annotate.angle=0)