Package Ravages (RAre Variant Analysis and GEnetic Simulation)

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Introduction

Ravages was developed to simulate genetic data and to perform rare variant association tests (burden tests and the variance-component test SKAT) on different types of phenotypes (Bocher et al., 2019, doi: 10.1002/gepi.22210, Bocher et al., 2020, doi:10.1038/s41431-020-00792-8) at a genome-wide scale. Ravages relies on the package Gaston developed by Herve Perdry and Claire Dandine-Roulland. Most functions are written in C++ thanks to the packages Rcpp, RcppParallel and RcppEigen.

Functions of Ravages use bed.matrix to manipulate genetic data as in the package Gaston (see documentation of this package for more details).

In this vignette, we illustrate how to perform rare variant association tests on real data on different phenotypes at a genome-wide scale. A second vignette is available showing how to simulate genetic data and how to use them for power calculation. To learn more about all options of the functions, the reader is advised to look at the manual pages.

Global parameters of Ravages

Functions in Ravages are parallelised either using RcppParallel when functions are implemented in C++ or using parallel otherwise. When the argument *cores* is present in the function, it can be directly changed to fix the number of cores to use. Otherwise, the number of threads used by multithreaded functions can be modified through RcppParallel function **setThreadOptions()**. It is advised to try several values for the number of threads, as using too many threads might be counterproductive due to an important overhead. The default value set by RcppParallel is generally too high.

Information on progression is provided if verbose = T for multiple functions in Ravages.

Example of analysis using LCT data

Below is an example of an association analysis and previous steps of data filtering using the dataset LCT.matrix available with the package Ravages. This dataset containts data from the 1000Genome project in the locus containing the Lactase gene. In this example, we look for an association between rare variants and the european populations of 1000Genomes. The population of each individual is available in the dataframe LCT.matrix.pop1000G. A classical analysis by gene is performed, and an analysis using the strategy "RAVA-FIRST".

Details about the "RAVA-FIRST" strategy and each function is given right after this example.

```
# Import data in a bed matrix
x <- as.bed.matrix(x=LCT.matrix.bed, fam=LCT.matrix.fam, bim=LCT.snps)
# Add population
x@ped[,c("pop", "superpop")] <- LCT.matrix.pop1000G[,c("population", "super.population")]
# Select EUR superpopulation
x <- select.inds(x, superpop=="EUR")</pre>
```

```
x@ped$pop <- droplevels(x@ped$pop)</pre>
# Group variants within know genes by extending their positions
# 500bp upstream and downstream
# (the function uses build 37 unless told otherwise)
x <- set.genomic.region(x, flank.width=500)
# a quick look at the result
table(x@snps$genomic.region, useNA = "ifany")
##
## R3HDM1 UBXN4
                    LCT
                          MCM6
                                 DARS
                                        <NA>
     2047
            1207
                   1454
                          1149
                                  924
                                        1295
# Filter variants with maf in the entire sample lower than 1%
# And keep only genomic region with at least 10 SNPs
x1 <- filter.rare.variants(x, filter = "whole", maf.threshold = 0.01, min.nb.snps = 10)
table(x1@snps$genomic.region, useNA="ifany")
##
## R3HDM1 UBXN4
                    LCT
                          MCM6
                                 DARS
                                  136
##
      268
             172
                    208
                           163
# run burden test CAST, using the 1000Genome population as "outcome"
# Null model for CAST
x1.H0.burden <- NullObject.parameters(x1@ped$pop, ref.level = "CEU",
                                      RVAT = "burden", pheno.type = "categorical")
burden(x1, NullObject = x1.H0.burden, burden = "CAST", cores = 1)
## Categorical phenotype
##
               p.value is.err
## R3HDM1 1.300274e-04
## UBXN4 1.993354e-05
## LCT
         4.489474e-08
## MCM6
       3.142808e-08
## DARS
        2.064661e-03
                            0
# run SKAT, using the 1000Genome population as "outcome"
# COnstruct null model for SKAT, then run test with only a few permutations
x1.H0.SKAT <- NullObject.parameters(x1@ped$pop, RVAT = "SKAT", pheno.type = "categorical")
SKAT(x1, x1.H0.SKAT, params.sampling=list(perm.target = 10, perm.max = 500))
## Categorical phenotype
## permutations
##
                        p.perm
                                     p.chi2
## R3HDM1 6.445786 0.001996008 4.186851e-07 4.186851e-07
## UBXN4 2.136390 0.011976048 1.539297e-02 1.539297e-02
         3.623174 0.001996008 3.776557e-04 3.776557e-04
## I.CT
        2.923228 0.007984032 4.636028e-03 4.636028e-03
## MCM6
        2.132967 0.041198502 5.909831e-02 4.119850e-02
# run a similar analysis but using the RAVA-FIRST approach with WSS
# RAVA-FIRST(x1, filter = "whole", maf.threshold = 0.01, min.nb.snps = 10,
           burden = TRUE, x1.H0.burden, SKAT = F)
```

Defining genomic regions

For rare variant association tests, the unit of analysis is not a single variant but a genomic region or 'testing unit', typically a gene. The first step of the analysis is therefore to group variants into genomic regions. This can be done using the function **set.genomic.region()** and known regions positions. It works on a bed.matrix (see Gaston) and simply adds a column "genomic.region" to the slot **x@snps** containing the genomic region (a factor) assigned to each variant. The positions of regions should be given to regions as a dataframe in a bed format containing the following columns: Chr. Start (0-based), End (1-based), Name. The dataframe should absolutely be ordered in the genome order, as well as the levels of regions\$Name.

By default, any variant being outside the gene positions won't be annotated. Regions boundaries can be extended to include more variants using the argument *flank.width* corresponding to the number of base pairs upstream and downstream the region to which expand the positions. If *flank.width=Inf*, each variant will be assigned to the nearest region.

If two regions overlap, variants in the overlapping zone will be attributed to both regions, separated by a comma. By default, split=TRUE in **set.genomic.region()**, which means that variants attributed to multiple regions will be duplicated in the bed matrix. This is done by calling the function **bed.matrix.split.genomic.region()** which takes a bed matrix as argument (x) and duplicates variants being assigned to multiple regions separated by split.pattern. If changeID=TRUE, the id from the bedmatrix will be changed to the format chr:pos:A1:A2:genomic.region to distinguish the duplicated variants.

The files **genes.b37** and **genes.b38** available in Ravages which contain gene positions from ENSEMBL versions GRCH37 and GRCH38 can be used as *regions*.

Running the following toy example will show you the behavior of these functions.

Rare variant selection

###Frequency filter To perform rare variant analysis, it is also important to define what is a rare variant in order to leave out common ones that are expected to have a weaker functional impact. The function filter.rare.variants() enables to keep only variants with a MAF (Minor Allele Frequency) below a given threshold while leaving out monomorphic variants. This function uses and returns a bed.matrix which can be filtered in three different ways:

- If filter="whole", all the variants with a MAF lower than the threshold in the entire sample will be kept.
- If filter="controls", all the variants with a MAF lower than the threshold in the controls group will be kept. In this situation, the controls group needs to be specified to the argument ref.level.

• If filter="any", all the variants with a MAF lower than the threshold in any of the groups will be kept.

It is also possible to specify the minimum number of variants needed in a genomic region to keep it using the parameter *min.nb.snps*, as well as the minimum cumulative MAF using *min.cumulative.maf*.

For the any and controls filters, the group of each individual should be given as a factor to group.

Rare variant association tests

We have implemented two burden tests extensions (CAST and WSS, doi:10.1002/gepi.22210) and an extension of the variance-component test SKAT (doi:10.1038/s41431-020-00792-8) to perform the association tests between a region and more than two groups of individuals.

The general idea of burden tests is to compute a genetic score per individual and per genomic region and to test if its distribution differs between the different groups of individuals. To extend these tests to more than two groups of individuals, a non-ordinal multinomial regression is used. The independant variable in this regression is the genetic effect of the region represented by the genetic score. Covariates can be added in the model. In addition to the genetic scores CAST and WSS directly implemented in the package, the user can specify another genetic score for the regression.

The variance-component test SKAT looks at the dispersion of genetics effects of rare variants. A geometrical interpretation of the test was used for its extension to more than two groups of individuals. Covariates can also be included in this model.

Genetic score for burden tests

We have implemented two functions to compute CAST and WSS scores respectively. These functions return a matrix with one row per individual and one column by genomic region. They are directly called in the function burden() if these scores are used to perform the association tests. It is also possible to compute genetic scores in a genomic region based on a vector of weights for each variant using the function burden.weighted.matrix(). It is important to note that all burden functions compute a score for the rare alleles. Therefore, if the reference allele is rare, the alleles will be flipped and the reference allele will be counted in the score instead of the alternative allele. This can be avoided only in the CAST() score if flip.rare.alleles = FALSE (it is set at TRUE by default).

CAST

CAST is based on a binary score which has a value of one if an individual carries at least one variant in the considered genomic region, and 0 otherwise. A MAF threshold for the definition of a rare variant is therefore needed (argument *maf.threshold*). This score can be computed using the function **CAST()** as shown here on the LCT data:

```
# Calculation of the genetic score with a maf threshold of 1%
CAST.score <- CAST(x = x1, genomic.region = x1@snps$genomic.region, maf.threshold = 0.01)
head(CAST.score)</pre>
```

##		R3HDM1	UBXN4	LCT	MCM6	DARS
##	HG00096	0	0	1	1	1
##	HG00097	1	0	0	0	0
##	HG00099	1	0	0	0	0
##	HG00100	0	1	0	0	0
##	HG00101	0	1	0	0	0
##	HG00102	1	0	0	0	1

WSS

WSS (Weighted Sum Statistic) is based on a continuous score giving the highest weights to the rarest variants:

$$WSS_j = \sum_{i=1}^R I_{ij} \times w_i$$

with

$$w_i = \frac{1}{\sqrt{t_i \times q_i \times (1 - q_i)}}$$

and

$$q_i = \frac{n_i + 1}{2t_i + 1}$$

Where n_i is the total number of minor alleles genotyped for variant i, t_i is the total number of alleles genotyped for variant i and I_{ij} is the number of minor alleles of variant i for the invidual j. In the original method, each variant is weighted according to its frequency in the controls group. In our version of WSS, the weights depend on allele frequencies calculated on the entire sample; this avoids using permutations. The function **WSS()** can be used to compute the WSS score as shown on the LCT data:

```
WSS.score <- WSS(x = x1, genomic.region = x1@snps$genomic.region)
head(WSS.score)</pre>
```

```
## HG00096 0.0000000 0.000000 0.8185268 1.26932 1.418436
## HG00097 0.8185268 0.000000 0.0000000 0.00000 0.000000
## HG00099 1.0019881 0.000000 0.000000 0.00000 0.000000
## HG00101 0.0000000 1.001988 0.0000000 0.00000 0.000000
## HG00102 1.0019881 0.0000000 0.000000 0.000000 0.000000
```

Other genetic scores

It is also possible to compute other genetic scores based on variants weights using the function **burden.weighted.matrix()**. The weights should be given as a vector to *weights* (with the length as the number of variants). The genetic score will be compute as:

$$Score_j = \sum_{i=1}^R I_{ij} \times w_i$$

with w_i the weight of each variant in weights, and I_{ij} the number of minor alleles for individual j in variant i. Here is an example corresponding to a genetic score with all the weights at 1, i.e. counting the number of minor alleles:

```
Sum.score <- burden.weighted.matrix(x = x1, weights = rep(1, ncol(x1)))
head(Sum.score)</pre>
```

##		R3HDM1	UBXN4	LCT	MCM6	DARS
##	HG00096	0	0	1	2	2
##	HG00097	1	0	0	0	0
##	HG00099	1	0	0	0	0
##	HG00100	0	1	0	0	0
##	HG00101	0	1	0	0	0
##	HG00102	1	0	0	0	1

Regressions

We have extended burden tests using a non-ordinal multinomial regression model. Let consider C groups of individuals including a group of controls (c = 0) and C - 1 groups of cases with different sub-phenotypes of

the disease. We can compute C-1 probability ratios, one for each group of cases:

$$ln\frac{P(Y_{j}=c)}{P(Y_{j}=0)} = \beta_{0,c} + \beta_{G,c}X_{G} + \beta_{k1,c}K_{1} + \ldots + \beta_{kl,c}K_{l}$$

where Y_j corresponds to the phenotype of the individual j and K_l is a vector for the lth covariate with the corresponding coefficient β_{kl} . The genetic effect is represented by X_G and correspond to the genetic score (for example CAST or WSS) with $\beta_{G,c}$ the log-odds ratio associated to this burden score.

The p-value associated to the genetic effect is calculated using a likelihood ratio test comparing this model to the same model without the genetic effect (null hypothesis). If only two groups are compared, a classical logistic regression is performed.

This regression can be performed on a bed.matrix using the function **burden()** which relies on the package mlogit. Parameters under the null model can be obtained using the function **NullObject.parameters()**. To generate this null model, the phenotype (argument *pheno*) of each individual should be given as a factor, and the potential covariates to include in the model should be given as a matrix to the argument *data* (one row per individual and one column per covariate). If only a subset of covariates from *data* are to be included in the model, a R formula should be given to *formula* with these covariates, otherwise all the covariates will be included. In addition, the reference group should be given to the argument *ref.level*, i.e. all odds ratios will be computed in comparison to this group of individuals. The choice of the reference group won't affect the p-value. As the parameters needed to run the association tests depend on the type of test performed (burden tests or SKAT), and the type of phenotype (continuous or categorical), both arguments shouls be given to **NullObject.parameters()** (*RVAT* and *pheno.type* respectively).

The function NullObject.parameters() will return a list with the parameters to use in burden() to run the burden test, including the Log-Likelihood computed under the null model, the argument data with the covariates to include determined from the argument formula.

Once the null model has been created, the function **burden()** can be used to perform burden tests. To do so, the user needs to give the results from **NullObject.parameters()** to the argument *NullObject*, and needs to specify the genomic region associated to each variant (argument *genomic.region*).

The CAST or WSS genetic scores can be directly calculated in the regression (burden="CAST" or burden="WSS"). The user can also use another genetic score in the regression, which has to be specified as a matrix with one row per individual and one column per genomic region to burden. In this situation, no bed matrix is needed, and the result from **burden.weighted.matrix()** can be used directly.

To shorten the computation time, calculations are parallelised; the argument *cores*, set at 10 by default, controls the parallelisation.

The function **burden()** will return the p-value associated to the regression for each genomic region. If there is a convergence problem with the regression, the function will return 1 in the column is.err. The effect size (odds ratio for categorical phenotypes and beta value for continuous phenotypes) associated to each group of cases compared to the reference group (NullObject\$ref.level) with its confidence interval at a given alpha threshold (argument alpha) can also be obtained with qet.effect.size=TRUE.

An example of the p-value and OR calculation with its 95% confidence interval using WSS on the LCT data is shown below with or without the inclusion of covariates. The outcome here corresponds to the population from 1000Genome.

Finally, using Ravages, it is also possible to perform burden tests with a continuous phenotype by specifying *pheno.type* = "continuous", and by giving a numeric vector to *pheno* as showed below:

###Functionally-informed burden tests Finally, we offer to perform functionally-informed burden tests to take into account functional information. To do so, we propose to define genomic regions (for example a gene of interest) and to integrate sub-scores in the regression corresponding to different functional categories (for example distinguish between coding and regulatory variants). A burden test will then be applied for each genomic region but will take into account the different categories of variants. The number of freedom corresponds to: (number of groups of individuals - 1) * number of sub-scores.

To attribute variants to genomic region and subregions, the function **set.genomic.region.subregion()** needs to be used which is very similar to **set.genomic.region()**. Two dataframes (in bed format) should be included: one for the large genomic regions (regions), and one for the subregions (subregions). Two columns will be added to x@snps: genomic.region and SubRegion.

The function **burden.subscores()** can then be applied on this bed matrix, which works similarly as **burden()**, with the extra argument *SubRegion* corresponding to the vector with the subregions on which subscores should be computed. In addition, the argument *burden.function* replaces the argument *burden* in **burden()** and requires to give a function which determines how the genetic score is computed (for example *CAST* or *WSS*).

Below is an example of the two functions where subscores correspond to coding and regulatory categories in the LCT locus.

```
# *** Functionally-informed WSS analysis ***
# Attribution of variants to regions and subregions
x2 <- set.genomic.region.subregion(x, regions = genes.b37,</pre>
                                   subregions = subregions.LCT)
# Burden test
burden.subscores(x2, x1.H0.burden, cores = 1)
## Categorical phenotype
##
               p.value n_subscores is.err
## R3HDM1
                    NA
                                  3
## UBXN4 2.626565e-21
                                  2
                                         0
                                  3
## LCT
                    NA
                                         1
```

```
## MCM6 3.488717e-31 2 0
## DARS NA 3 1
```

SKAT

We also extended the variance-component test SKAT using a geometric interpretation. Unlike the burden tests, the is no burden calculated in this test: the distribution of the genetic effects in the genomic region is compared to a null distribution. SKAT is based on a linear mixed model where the random effects correspond to the genetic effects.

Before running SKAT, the function **NullObject.parameters()** first needs to be called as for the burden tests, but by specifying RVAT = "SKAT". As before, potential covariates could be included as a matrix to data. If only some of them are to be included, they should be given as a R formula to formula. This function will compute parameters to use the **SKAT()** function, and should be given to this function using the argument NullObject.

To compute the p-values, a chi-square approximation is used based on the statistics' moments. The moments can either be estimated using a sampling procedure, or be analytically computed using the method from Liu et al. 2008. The chi-square approximation can be based on the first three moments (estimation.pvalue = "skewness"), or on moments 1, 2 and 4 to have a more precise estimation of the tail distribution (estimation.pvalue = "kurtosis"). If get.moments = "theoretical" and estimation.pvalue = "skewness", it is equivalent to the "liu" method in the SKAT package, and if estimation.pvalue = "kurtosis", it corresponds to the "liu.mod" method in the SKAT package. If debug = TRUE, the statistics' moments will be returned in addition to the p-values.

If the sample size is lower than 2000, we recommand to use the sampling procedure. If no covariates are present, a simple permutation procedure can be used (get.moments = "permutations"), otherwise, a boostrap sampling should be used (get.moments = "bootstrap"). For those two situations, a sequential procedure is used to compute the p-values: permutated statistics are computed and each one is compared to the observed statistics. The sampling procedure stops when either perm.target (the number of times a permutated statistics should be greater than the observed statistics) or perm.max (the maximum number of permutations to perform) is reached. P-values are then computed in two different ways: if perm.target is reached, the p-value is computed as perm.target divided by the number of permutations performed to reach this value; if perm.max is reached before perm.target (that is, for pretty small p-values), p-values are computed using the chi-square approximation based on moments estimated from the permutated statistics. perm.target and perm.max should be given as a list to the argument params.sampling of SKAT.

If the sample size is bigger than 2000, the analytical calculation from Liu et al. (2009) can be used to compute the theoretical moments. In this situation, it is possible to parallelise the calculations using the argument *cores*, set at 10 by default. Nevertheless, this method can take more computation time than the permutation procedures when the sample size is large. We therefore recommend to use one of the permutation procedures to run a first analysis, and then to use the theoretical moments only for the most associated regions.

It is possible to clearly ask for a specific method to compute the moments using get.moments = "permutations", "bootstrap" or "theoretical". By default, get.moments = "size.based", and the method will depend on the sample size.

An example of the SKAT function by specifying the "permutations" or "theoretical" method is shown.

It is also possible to perform the SKAT test on a continuous phenotype by using the argument *pheno.type* = "continuous" in NullObject.parameters() before the SKAT() function:

RAVA-FIRST (RAre Variant Analysis using Functionally-InfoRmed STeps)

Ravages also offers the possibility to analyse rare variants using the 'RAVA-FIRST' strategy, composed of three main steps: defining testing units, filtering rare variants, and running functionnaly informed burden tests.

###Definition of testing units: CADD regions CADD regions are non-overlapping regions defined genome-wide using the variant pathogenicity score CADD that can be used as testing units: regions are defined between SNVs observed at least two times in GnomAD with a high CADD score. This CADD score is not the original CADD score v1.4 oublished by Rentzch et al., but an adjusted score that take into account three types of functional categories: coding, regulatory and intergenic categories. This adjusted score enables to find the most important functional variants within each of those three categories.

To attribute adjusted CADD scores to variants, the function adjustedCADD.annotation() can be used. It will call the function adjustedCADD.annotation.sNVs() and adjustedCADD.annotation.indels() to annotate SNVs and indels respectively. As CADD scores are available for every SNVs in the genome, SNVs in the bed matrix will be annotated using a provided file with adjusted CADD scores to SNVs.scores (with columns 'chr', 'pos', 'A1', 'A2', 'adjCADD'), or by downloading the adjusted scores from https://lysine.univ-brest.fr/RAVA-FIRST/ in a repository indicated to path.data. Oppositely, pre-computed PHRED CADD scores are not available for every indel in the genome and they should be annotated online in https://cadd.gs.washington.edu/. To computed adjusted CADD scores for indels, we used a set of 48M indels available at https://cadd.gs.washington.edu/download. Indels in the bed matrix will then be annotated using the file AdjustedCADD_v1.4_202204_indels.tsv.gz downloaded from https://lysine.univ-brest.fr/RAVA-FIRST/ and if new indels not present in the set of 48M indels are to be annotated, they will be given the same adjusted score as the indel with the nearest PHRED v1.4 CADD score. Therefore, if indels are present in the bed matrix, a file with the CADD PHRED v1.4 score should be given to indels.scores in adjustedCADD.annotation() (with columns 'chr', 'pos', 'A1', 'A2', 'CADD').

To assign variants to CADD regions and to the functional categories, the function **set.CADDregions()** can be used. It will add genomic.region to the bed matrix with the corresponding CADD regions present in the file "CADDRegions.2021.hg19.bed.gz" and SubRegion with the functional area present in the file "FunctionalAreas.hg19.bed.gz", both directly downloaded from https://lysine.univ-brest.fr/RAVA-FIRST/ in the repository deifined in path.data. In addition, adjCADD.Median will be added to x@snps which correspond to the median adjusted CADD score of variants (SNVs + indels) observed at least two times in GnomAD and can be used for variant filtering as explained later.

```
# Attribution of CADD regions
x.CADDregions <- set.CADDregions(x)
# Annotation of variants with adjusted CADD scores
x <- adjustedCADD.annotation(x)</pre>
```

###Filtering of rare variants: region-dependent thresholds In RAVA-FIRST, we propose a new approach to select rare variants to include in the RVAT that keeps within each CADD region only the variants with an adjusted CADD score greater than the median observed in GnomAD. To this end, the function filter.adjustedCADD() can be used which relies on the same arguments than filter.rare.variants() for the frequency parameters. Within this function, SNVs and indels of the bed matrix will be directly

annotated using adjustedCADD.annotation() described earlier. If the bed matrix has previously been annotated by adjustedCADD.annotation(), filter.adjustedCADD() will filter rare variants based on x@snps\$adjCADD without annotating the variants to gain in computation time. Only the variants with a score greater than the median will be kept. It is necessary that the bed matrix is first annotated with set.CADDregion() to get the median of each variant from its corresponding CADD region.

```
# Keep only CADD regions with 2 variants and variants with a MAF greater than 1%
# and with an adjusted CADD score greater than the median
x.median <- filter.adjustedCADD(x.CADDregions, maf.threshold = 0.01, min.nb.snps = 2)</pre>
```

###Functionally-informed burden tests CADD regions can overlap different types of functional categories (coding, regulatory or intergenic regions). To take them into account in RAVA-FIRST, we propose to use the functionally-informed burden test **burden.subscores()** with subscores in the regression, one for each functional category, within each CADD region. There will be at most three sub-scores in the regression. **burden.subscores()** can be directly applied on the bed matrix obtained from **set.CADDregions()** or filter.adjustedCADD().

SKAT() can also be applied on the bed matrix with variants grouped and filtered according to RAVA-FIRST but it will correspond to the classical SKAT analysis without the integration of functional information.

```
# Functionally-informed WSS analysis
x.burden <- burden.subscores(x.median, x1.H0.burden, burden.function = WSS)
# SKAT analysis
x.SKAT <- SKAT(x.median, x1.H0.SKAT)</pre>
```

###Complete RAVA-FIRST approach and whole-genome analysis RAVA-FIRST enables to perform RVAT in the whole genome. To facilitate the analysis, we propose the function **RAVA.FIRST()** which gathers all the previous functions and enables to directly performs the whole RAVA-FIRST strategy on a bed matrix. It simply needs the bed matrix and the NullObject (that can be obtained from **NullObject.parameters()** and should be given to H0.burden and/or H0.SKAT) and optional filtering arguments and parameters for the RVAT previously mentionned. burden.parameters should be a list containing burden.function and get.effect.size if burden = TRUE and SKAT.parameters should be a list containing weights, get.moments estimation.pvalue, param.sampling and debug if SKAT = TRUE. If no such list is provided, the default parameters of **burden.subscores()** and **SKAT** will be used. Due to time and memory management, we advise the user to import the data and apply the different functions of annotation, filtering and association (or directly **RAVA.FIRST()**) chromosome by chromosome for large datasets as shown in the example below.

When using RAVA-FIRST(), results of association analysis will be provided for burden and/or SKAT depending on the corresponding arguments. In addition of the results of the test, information about the CADD region will be provided: the positions of the regions, the type of genomic category overlapped by each CADD region, and finally the median of adjusted CADD scores used for the filtering of rare variants.

To get more information about a specific CADD region of interest, especially about the positions of the genomic categories in this region, the user is advised to directly use the file "FunctionalAreas.hg19.bed.gz" and the corresponding tabix file in the Ravages repository to look at the positions of the corresponding CADD region.

Data management

Data in plink format or in vcf format can be loaded in R using the functions read.bed.matrix() and read.vcf() respectively from the package gaston.

If the data for the controls and the different groups of cases are in different files, they can be loaded separately and then combined using the function **gaston::rbind()** as long as the same variants are present between the different groups of individuals.

An example is given below where the simulated data have been split according the the group of each individual, and then combined in a bed.matrix:

```
# Selection of each group of individuals
CEU <- select.inds(x1, pop=="CEU")
CEU
## A bed.matrix with 99 individuals and 947 markers.
## snps stats are set
   There are 748 monomorphic SNPs
## ped stats are set
FIN <- select.inds(x1, pop=="FIN")
FIN
## A bed.matrix with 99 individuals and 947 markers.
## snps stats are set
   There are 771 monomorphic SNPs
## ped stats are set
GBR <- select.inds(x1, pop=="GBR")
GBR
## A bed.matrix with 91 individuals and 947 markers.
## snps stats are set
    There are 792 monomorphic SNPs
## ped stats are set
# Combine in one file:
CEU.FIN.GBR <- rbind(CEU, FIN, GBR)
CEU.FIN.GBR
## A bed.matrix with 289 individuals and 947 markers.
## snps stats are set
    There are 515 monomorphic SNPs
## ped stats are set
```